Methods

Tn7-Based Genome-Wide Random Insertional Mutagenesis of Candida glabrata

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We describe and characterize a method for insertional mutagenesis of the yeast pathogen Candida glabrata using the bacterial transposon Tn7. Tn7 was used to mutagenize a C. glabrata genomic fosmid library. Pools of random Tn7 insertions in individual fosmids were recovered by transformation into Escherichia coli. Subsequently, these were introduced by recombination into the C. glabrata genome. We found that C. glabrata genomic fragments carrying a Tn7 insertion could integrate into the genome by nonhomologous recombination, by single crossover (generating a duplication of the insertional mutagenized locus), and by double crossover, yielding an allele replacement. We were able to generate a highly representative set of ∼104 allele replacements in C. glabrata, and an initial characterization of these shows that a wide diversity of genes were targeted in the mutagenesis. Because the identity of disrupted genes for any mutant of interest can be rapidly identified, this method should be of general utility in functional genomic characterization of this important yeast pathogen. In addition, the method might be broadly applicable to mutational analysis of other organisms.

Candida species, primarily Candida albicans and Candida glabrata, are important human pathogens, responsible for 7% of all hospital-acquired bloodstream infections (Schaberg et al. 1991). Even with available antifungal therapies, the associated mortality for Candida bloodstream infections is high (up to 30% in cancer patients). We are using C. glabrata as a model to explore the molecular details of the host–pathogen interaction. In frequency of isolation, C. glabrata is second only to C. albicans and is responsible for 15%–20% of both mucosal (Schuman et al. 1998; Vazquez et al. 1999) and systemic (Pfaller et al. 1999, 2001) candidiasis; in spite of this epidemiological similarity, C. glabrata is phylogenetically distant from C. albicans, more highly related, for example, to Saccharomyces cerevisiae than to C. albicans (Barns et al. 1991). What strategies for host colonization are shared by C. glabrata and C. albicans remain to be determined; no genes essential for virulence have yet been described in C. glabrata. Studies primarily in C. albicans have identified multiple factors important in the pathogenesis of Candida species (for review, see Calderone and Fonzi 2001), including the ability to adhere to host tissue, the ability to grow in hyphal and yeast form (for C. albicans), the capacity to switch between different cellular phenotypes, and the ability to acquire iron in vivo. Like C. albicans, C. glabrata is able to adhere specifically to host tissue, recognizing host carbohydrate (Cormack et al. 1999). On the other hand, C. glabrata does not make hyphae, a feature of prime importance in the pathogenesis of C. albicans; rather, it grows solely in the yeast form, making pseudohyphae under conditions of nutrient starvation (Csank and Haynes 2000).

Because C. glabrata is haploid, the tools of classical genetics can be applied, and mutants defective in various aspects of virulence can be isolated and characterized. An efficient genetic analysis depends on a method of random insertional mutagenesis, and we considered various available options. In other species, numerous approaches have been taken, including the use of bacterial transposons such as Tn3 (Seifert et al. 1986; Ross-Macdonald et al. 1999), Tn7 (Biery et al. 2000), and the Drosophila melanogaster transposon Mariner (Gueiros-Filho and Beverley 1997). Tn3, in particular, has been used to advantage in mutagenesis of S. cerevisiae (Ross-Macdonald et al. 1999). In that efficient and highly random method, fragments of the S. cerevisiae genome are first mutated in S. cerevisiae to make insertion mutants and to analyze these for effects on adherence to epithelial cells (Cormack and Falkow 1999; Cormack et al. 1999). We found that the insertions were distributed more or less randomly in many different genes; however, a close analysis of the sites of insertion for 50 mutants showed that the majority (48/50) were in noncoding regions of the genome. If one were able to analyze a very large number of mutants, this bias might not be an important factor. However, for screens in which only modest numbers of mutants (20,000–30,000) are analyzed, the bias against insertions in coding regions would result in a mutational sampling of only a fraction of the genome.

As an alternative to the problematic nonhomologous-recombination-based method, we describe in this paper a novel mutagenesis approach similar in principle to the Tn3 method described above (Ross-Macdonald et al. 1999), but which exploits recent studies of in vitro transposition by the

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bacterial transposon Tn7. This method is of some general interest because the generation of mutants requires only two steps: in vitro mutagenesis by Tn7 followed by homologous recombination into the target genome (here C. glabrata). In theory, therefore, our method can easily be applied to any organism with efficient homologous recombination. We describe modifications to Tn7 to allow its use in C. glabrata and to facilitate the recovery of DNA flanking insertion sites for mutants of interest. We demonstrate that this method can be used in the efficient generation of thousands of randomly distributed insertion mutants, possessing an array of phenotypes.

**RESULTS**

**Principle of Mutagenesis and Construction of a Minitransposon With Two Yeast Selectable Markers and a Conditional Origin of Replication for E. coli**

We devised a two-step strategy for random insertional mutagenesis in C. glabrata. In this method, a collection of fosmids containing large (~40-kb) C. glabrata genomic inserts (in essence, a representation of the genome) is first insertionally mutagenized by Tn7 in vitro to create pools of mutants in E. coli. This is followed by introduction of the pool of mutants (at this point carried on fosmids in E. coli) into C. glabrata by homologous recombination. In theory, this should result in an allelic replacement, in which individual genes have been replaced by insertionally mutagenized alleles.

We used a Tn7-based in vitro mutagenesis to mutagenize a set of 100 fosmids (~0.25x genome coverage). This in vitro mutagenesis system allowed efficient transposition with minimal target-site specificity (Biery et al. 2000). The mini-Tn7 transposon previously described (mini-Tn7 SpeI-KmR-NotI; Biery et al. 2000), carries a kanamycin-resistance cassette flanked by the Tn7 left- and right-end sequences, respectively. As shown in Figure 1, we constructed two different derivatives of this transposon. To generate the first one, Tn7 URA3·KmR·R6-Ky (Tn7 UKR), we introduced into the mini-Tn7 a yeast selectable marker, the URA3 gene from S. cerevisiae, as well as the conditional origin of replication R6-Ky, which requires the II protein (the product of the pir gene) for replication (Kolter et al. 1978). The presence of this origin facilitates subsequent recovery of DNA flanking the insertion site in mutants of interest. The resulting transposon can be maintained only in an E. coli strain (BW23473) expressing the pir gene (Metcalf et al. 1996). The second transposon, Tn7 URA3·hph·KmR·R6-Ky (Tn7 UKR-H), contains, in addition, the hph gene from Klebsiella pneumoniae, which confers resistance to hygromycin B (Gritz and Davies 1983).

We tested the ability of these minitransposons to undergo transposition in vitro using different-sized target DNAs and found no difference with the parent mini-Tn7 (data not shown). We used each transposon for mutagenesis of individual fosmids and cloned and characterized 40 transposition products by digestion with restriction enzymes (data not shown). In no transposition products did we detect a gross rearrangement of fosmid DNA. In addition, the majority (32/40) of transposition products had a single insertion of the transposon in the fosmid with no other rearrangement of the fosmid DNA. In 8 out of the 40 products, there were 2 Tn7 insertions, but these were invariably in two different restriction fragments. This result is not unexpected, because over short distances the phenomenon of transposon immunity prevents the insertion of a second copy of the transposon next to the site of a transposon already present. This effect drops off over distance, so that one might expect two insertions in the same fosmid but separated by >10 kb (DeBoy and Craig 1996).

Mutagenesis of our fosmids was efficient: In a mutagenesis of 200 fosmids, we recovered a minimum of 200 and up to tens of thousands of transformants per fosmid, where each transformant represents an independent insertion in the fosmid. For each fosmid, the transformants were pooled, generating a library of insertions throughout a fosmid-borne 40-kb genomic fragment.

**Transformation of Linear Genomic Fragments Into C. glabrata Results in Three Classes of Transformants**

Before introducing a pool of Tn7-mutagenized fosmid fragments into the C. glabrata genome, we wanted to characterize in detail the in vivo fate of individual mutagenized genomic fragments, derived from one mutagenized fosmid, and to optimize conditions for recovery of homologous recombinants after transformation of mutagenized fosmids into C. glabrata. Transposition Tn7-UKR (from pIC6) was used to mutagenize one fosmid (fosmid 1). Mutagenized fosmid DNA was linearized with EcoRI, transformed into C. glabrata strain BG14 (ura2Δ), and selection was made for Ura+ transformants. Ura+ colonies were then patched on a plate lacking uracil and printed onto plates containing 5-fluoroorotic acid (5-FOA). Upon examination of Ura+ transformants obtained, it was clear that there were three different classes of transfor-

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**Figure 1** Maps of mini-Tn7 derivatives used. Tn7 UKR (URA3 Kan·R6-Ky ori) is a 3.108-kb derivative of mini-Tn7 SpeI-KmR-NotI containing the conditional origin of replication R6-Ky and the Saccharomyces cerevisiae URA3 gene. Relevant restriction sites are shown. Tn7 UKR-H is a 5.118-kb element derived from Tn7 UKR that contains the Klebsiella pneumoniae hph gene (which confers resistance to hygromycin B) driven by the PGK1 promoter from S. cerevisiae. The arrows underneath each map indicate the direction of transcription.
mants based on the phenotype displayed on 5-FOA plates: Class 1 transformants were unstable Ura+ 5-FOAR colonies; Class 2 transformants were partially stable, Ura+ 5-FOA\(^5\) transformants in which a patch on 5-FOA plates gave rise to resistant papillae after incubation at 30°C for 48 h; and Class 3 transformants were the expected Ura+ 5-FOA\(^3\) stable colonies. Each class is the result of a different fate of the transforming linear DNA fragment, as described below.

**Three Classes of Transformants Represent Three Different Fates of the Transforming DNA**

We hypothesized that Class 1 or unstable transformants (U) corresponded to linear genomic fragments (containing a Tn\(^7\) insertion), which happened to contain an autonomous replication sequence (ARS), and which were able to recircularize after transformation and remain as episomes or plasmids. These plasmids would be able to replicate, but because they would generally not contain a \(CEN\) sequence, they would be lost at a high frequency, resulting in a mixed colony containing Ura+ 5-FOA\(^3\) cells (containing the unstable plasmid), as well as Ura- 5-FOA\(^5\) cells, which have lost the plasmid. If this hypothesis were true, then it should be possible to isolate this freely replicating plasmid from genomic DNA preps from unstable transformants, and functionally identify it by its ability to transform the \(E. coli\) strain expressing the pir gene. We prepared genomic DNA from four individual Class 1 transformants, and used it directly to transform the \(E. coli\) strain BW23473. As expected, undigested genomic DNA from each unstable transformant produced thousands of transformants in this \(E. coli\) host, but not in an \(E. coli\) strain not expressing the pir gene (data not shown). Furthermore, plasmid DNA isolated from \(E. coli\) strain BW23473 could be transformed back into \(C. glabrata\) at the high frequency (\(>10^5\)/\(\mu g\) of DNA) typical of plasmid transformations in \(C. glabrata\) (Cormack and Falkow 1999).

We hypothesized that Class 2 or partially stable (PS) transformants were events in which the incoming, linear DNA recircularized before, or coincident with, the homologous recombination event. This would lead to a partial duplication of this particular locus, in which an insertional mutagenized allele and the wild-type allele are present as tandem direct repeats (Fig. 2A). Because of the direct repeat, it is possible that at some low frequency, the duplication could resolve by homologous recombination between the direct repeats, yielding Ura- 5-FOA\(^5\) papillae on an otherwise 5-FOA\(^3\) patch. To test this hypothesis, we first generated a defined

**Figure 2** Two of the pathways of homologous recombination after transformation of linearized mutant fosmids. (A) Class 2 (partially stable, PS) transformants: A linearized genomic fragment containing a transposon insertion can recircularize before, or at the time of, homologous recombination, generating a duplication of the region as indicated. Primers 1 and 2 are oriented toward, and primers 3 and 4 oriented away from the Tn\(^7\) insertion. (B) Result of PCR amplification of genomic DNA from one Class 2 transformant. (genom. PS lane) Template DNA for PCR was genomic DNA; (plasm. lane) template DNA for PCR was pIC14. (C) Class 3 (stable, S) transformants: allele replacement. A linearized genomic fragment containing a transposon insertion recombines by double crossover and replaces the genomic locus. (D) Result of PCR amplification of genomic DNA from nine Class 3 (S) transformants. (Lane 1) Genomic DNA from a nonhomologous recombinant was used for the PCR and bands for both the wild type and Tn7-disrupted copies are amplified. (Lanes 2–9) Results for homologous recombinants. Controls: (wt) Cg14 genomic DNA was used as template for the PCR; (ins.) pIC14 was used as template.


insertion in a single *C. glabrata* sequence. DNA from the mutagenized fosmid was digested with *Eco*RI, recircularized, and used to transform *E. coli* strain BW23473 (*pir*) to kanamycin resistance. This generated a smaller, conditional replicon, designated pIC14, containing a single Tn7 insertion and 8 kb of the flanking genomic *C. glabrata* sequence, which was sequenced using primers that anneal to each end of the Tn7 and are oriented outward. We then designed primers in both directions from the flanking genomic sequence obtained, toward and away from the insertion (Fig. 2A). pIC14 was linearized with *Eco*RI, transformed into BG14 (*ura3*Δ), and selection was made for *Ura*’ transformants. If the homologous recombination resulted in a duplication of this region as illustrated in Figure 2A, then by using the pair of primers whose direction of annealing is away from the insertion (“out-PCR”), it should be possible to amplify a fragment containing the duplication, but not if there was no duplication, because the two primers are oriented away from each other. Furthermore, by using the pair of primers oriented toward the insertion (“in-PCR”), two regions should be amplified from a duplication event: a wild-type small fragment of 614 bp and a larger fragment corresponding to the wild type plus the Tn7 UKR insertion (3720 bp).

Genomic DNA was prepared from two Class 2 (PS) transformants and was used as template for PCR reactions using the two pairs of primers from the sequences flanking the insertion. Figure 2B shows the results obtained with one PS transformant. As is clear from the picture, the primer pair oriented toward the insertion amplified the two fragments of the size expected of a duplication. When the primers directed away from the insertion were used, a product of the size expected was amplified. This result indicates that, in fact, a duplication occurred after transformation of this fragment of DNA containing the Tn7 UKR insertion either before or during homologous recombination. We also gel-purified high-molecular-weight genomic DNA from this and other PS transformant and obtained the same results with “in” and “out” primer pairs, indicating that in PS class transformants, the transforming DNA underwent a recombination pathway that led to a duplication of the region in the genome. Unlike in the case of Class 1 (unstable) transformants, we were unable to transform the *E. coli* permissive host BW23473 to kanamycin resistance with genomic DNA purified from the two Class 2 transformants (data not shown). This is again consistent with our hypothesis that these transformants represent tandem duplications of the Tn7-disrupted genomic locus.

Class 3 or stable (S) transformants represent allele replacements in which the insertion of Tn7 UKR at that particular locus replaces the wild-type region, resulting in a stable *Ura*’ colony (Fig. 2C). We made genomic DNA of 30 stable transformants derived from pIC14 transformation and used the PCR primers directed toward the insertion to analyze them. Figure 2D shows nine transformants of which eight using primers 708 and 711 (corresponding to insertion 3 in Table 1) toward the insertion; instead, a 3.553-kb band was amplified, corresponding in size to the insertion of the Tn7 UKR at that site. Furthermore, only these same two genomic DNAs showed the expected 0.207-kb and 0.237-kb fragments amplified with primers from the left and right junctions, respectively, of that particular insertion. These results indicate that in these two transformants, a replacement event took place, and also that insertion 5 was not present in any of the 20 other transformants as a nonhomologous recombinant or a double insertion. The results of the complete set of PCR

**The Majority of 12 Individual Insertions in a Pool Are Represented in a Small Pool of *C. glabrata* Stable Transformants**

We next wanted to know whether in a small pool of insertions in one fosmid, there would be homologous recombination and replacement of each insertion after transformation into *C. glabrata* and screening for stable transformants. This was an important parameter to understand because we worried that a single insertion might recombine at much greater than average efficiency and “poison” the pool of *C. glabrata* transformants derived from a pool of insertions in *E. coli*. We made a pool of 12 Tn7 UKR insertions in one fosmid, of which 4 were in the fosmid vector, and the remaining 8 were distributed in the five different-sized *Eco*RI fosmid fragments shown schematically in Table 1. The target site of each of these 8 insertions was sequenced using the Tn7 end primers facing outward, and pairs of primers were designed from each flanking sequence toward the insertion. We made DNA of the pool of 12 insertions from *E. coli* and digested it with *Eco*RI and used it to transform *BG14 (ura3Δ)*, selecting for *Ura*’. Genomic DNA from 20 stable Class 3 (S) FOA*’* transformants was prepared and used as template for PCR analysis. We used three pairs of primers for each of the insertions in the pool and used these to PCR genomic DNA from each of the 20 FOA*’* transformants. The first pair is oriented toward the insertion, and the other two amplify each junction of the Tn7 insertion. Figure 3 shows only one example using 11 FOA*’* DNAs and the three primer pairs specific for one of the eight insertions. In this case, 2 of the 11 genomic DNAs used (numbers 7 and 11) did not give amplification of a wild-type 0.445-kb fragment using primers 708 and 711 (corresponding to insertion 5 in Table 1) toward the insertion; instead, a 3.553-kb band was amplified, corresponding in size to the insertion of the Tn7 UKR at that site. Furthermore, only these same two genomic DNAs showed the expected 0.207-kb and 0.237-kb fragments amplified with primers from the left and right junctions, respectively, of that particular insertion. These results indicate that in these two transformants, a replacement event took place, and also that insertion 5 was not present in any of the 20 other transformants as a nonhomologous recombinant or a double insertion. The results of the complete set of PCR
reactions for the 20 genomic DNAs from PS transformants are summarized in Table 1. The first four insertions fell in the largest EcoRI fragment; none of these happened to disrupt any ORF within this fragment, and all four were found among the 20 transformants analyzed; insertion number 1 was the most commonly found (6/20 FOAS genomic DNAs). The four insertions in this fragment accounted for two-thirds (14/20) of the transformants analyzed, not unexpectedly because they accounted for one-half of the total number of insertions in the pooled DNAs, which can recombine by homologous recombination (4/8). Insertion number 5 in the second largest EcoRI fragment is at amino acid 456 of the *C. glabrata* INP53 homolog and was found three times in the 20 DNAs monitored (disruption of the ortholog of this gene in *S. cerevisiae* results in no phenotype). Insertion number 6 in the 2.8-kb EcoRI fragment was found once among the genomic DNAs; insertion number 7, in a 1.5-kb EcoRI fragment, which disrupts the YNL001w ortholog, was not found in any of the 20 genomic DNAs tested. Null mutations in this ORF result in slow growth in *S. cerevisiae*, and it is expected that mutations that confer a slow growth will be underrepresented in our method because in the initial transformation, these colonies would probably be avoided. We also did not find any transformants carrying insertion 8 in the smallest EcoRI fragment (400 bp), which disrupts YNL107w. It may be that this insertion results in a slow growth or inviability phenotype and therefore could not be recovered. Alternatively, 400 bp of homology is too low to give efficient homologous recombination.

From the data in Table 1, we can conclude that many different insertions in a pool can be recovered among a small number of recombinants in *C. glabrata*; for small fragments, there might be a correlation between the size of the fragment carrying the Tn7 and the likelihood of its integrating by homologous recombination. In all, of the 20 genomic DNAs from stable FOAS transformants, 18 of them were caused by homologous recombination involving six of the eight insertions in the genomic DNA; the remaining two transformants were caused by nonhomologous recombination; in one case, insertion 9 in the vector backbone recombined by nonhomologous recombination and in one case insertion 1 in the largest EcoRI fragment recombined by nonhomologous recombination. The initial pool of 12 *E. coli* insertions contained 4 (or one third of the pool) insertions in the vector, and these can integrate only by nonhomologous recombination.
Consistent with nonhomologous recombination being less efficient than homologous recombination, only 1 of 20 transformants was caused by recombination of a vector-derived fragment (as opposed to the 7 expected if homologous and nonhomologous recombination happened at equal frequency). It is clear from this analysis, as well as from the earlier analysis of a single fragment, that ∼10% of the stable transformants derived from this method are caused by nonhomologous recombination. The corollary is that ∼90% of the stable FOA⁺ transformants are simple allele replacements.

Most Insertions in a Large Pool Transform C. glabrata at Equal Efficiency

We wanted to further verify for a large pool of Tn7 insertions in a fosmid that individual insertions in the pool would give rise to transformants in C. glabrata at approximately the same frequency that they were represented in the E. coli pool. If all fragments in a pool recombined at equal frequency, this would be true. The exception to this would be insertions, which disrupt essential genes (which cannot be recovered at all in our mutagenesis). We mutagenized in vitro two fosmids and pooled ~150 E. coli individual transformants for each fosmid. DNA from these two pools was digested with BglI plus MfI (for neither one of which recognition sites are present in Tn7) and used to transform BG14 (ura3Δ) to Ura⁺. MfI recognition sites are present approximately every 3000 bp in the genome; BglI sites are present on average every 10 kb. The BglI sites in pBAC-BglI immediately flank the cloning site for the genomic fragment and therefore digestion with BglI releases the mutagenized genomic DNA fragments from any vector sequences, leaving genomic fragments with ends precisely complementary to the corresponding genomic locus. Genomic DNA from pools of 35 or 44 stable Class 3 (FOA⁺) transformants, as well as DNA from the E. coli pool that generated them, was analyzed by Southern blot using a labeled URA3 probe. Figure 4 shows the results of this experiment: The first two lanes correspond to the first C. glabrata pool and the E. coli pool, respectively; all of the bands from the bacterial pool are shared with the yeast pool, indicating that most of the individual insertions are indeed represented in C. glabrata transformants. Because the relative intensities of the individual bands in the E. coli pool and the C. glabrata pools are the same, this shows that the recombination of individual fragments carrying Tn7 occurs at approximately equal frequencies for most of the insertions in the pool, consistent with what we had found for the smaller pool of 12 insertions. The same result was found for the second fosmid (Fig. 4, lanes 4, 5). Stripping the nylon membrane and reprobing with vector sequences showed that there were no vector sequences in the C. glabrata pool as expected (data not shown). In a control experiment shown in lanes 6–8, fosmid vector pBAC was mutagenized with Tn7 UKR-H, and a pool of >1000 insertions in pBAC was generated. DNA from the pool was digested with BglI and MfI and used to transform BG14 (ura3Δ) to Ura⁺ by nonhomologous recombination of the vector fragments containing Tn7 UKR-H insertions. Genomic DNA from a pool of five C. glabrata stable transformants was analyzed by Southern blot. When a labeled URA3 probe was used, none of the bands in the E. coli pool (Fig. 4, lane 8) were also present in the yeast pool (Fig. 4, lane 6), showing that all of the C. glabrata transformants were nonhomologous recombinants.

Thus, all (or the majority) of the linear fragments of genomic DNA derived from a mutagenized fosmid containing
We determined the DNA sequence surrounding the insertion point for many of the amino acid auxotrophic mutants. Genomic DNA for any mutant of interest was digested either with XbaI or MfeI, and DNA ligase was added. This generated a circular plasmid containing the transposon and the DNA flanking the insertion site, which was recovered by transformation into E. coli selecting for Km resistance. Of 68 sequenced mutants, we determined that 58 (86%) were derived by homologous recombination. For the auxotrophs, the insertion points and identity of the disrupted gene for one representative insert in each of 11 loci are shown in Table 2. Approximately three-fourths of the inserts recovered from the C. glabrata genome were in the coding regions of genes.

**DISCUSSION**

We have mutagenized C. glabrata using a modified Tn7 transposon, derivatized with the S. cerevisiae URA3 gene, the K. pneumoniae hph gene, and the E. coli R6Kγ conditional origin. After mutagenesis of large fosmid-borne fragments of the genome in vitro and recovery of these insertion mutants in E. coli, the insertion mutants were introduced by allele replacement into the genome of C. glabrata. We had first tried the simpler approach of mutagenizing C. glabrata chromosomal DNA with Tn7 and directly transforming this pool of DNA into the organism, bypassing the need for the fosmid library or recovery of insertions in E. coli. However, probably because C. glabrata transformation is relatively inefficient and because, as detailed in this paper, there are multiple ways in which a linear DNA can integrate into the genome, direct transformation of mutagenized chromosomal DNA yielded no simple Tn7 insertions in the genome. We therefore used the scheme described in this paper.

The distribution of insertion sites for Tn7 itself in vitro showed essentially no bias at all, and we present evidence here that the full range of mutations generated in vitro can be introduced by homologous recombination into the genome of C. glabrata. The resultant library of mutants is a highly random collection of insertion mutations. This method, al-

### Table 2. Analysis of Auxotrophic Mutants

<table>
<thead>
<tr>
<th>Mutant</th>
<th>Gene</th>
<th>Insertion site</th>
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</thead>
<tbody>
<tr>
<td>15-04</td>
<td>ILV1</td>
<td>aa 268</td>
</tr>
<tr>
<td>31-36</td>
<td>ARG2</td>
<td>aa 441</td>
</tr>
<tr>
<td>37-26</td>
<td>SUL2</td>
<td>aa 388</td>
</tr>
<tr>
<td>42-37</td>
<td>lys14</td>
<td>aa 755</td>
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<tr>
<td>47-40</td>
<td>MET3</td>
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<td>lys12</td>
<td>aa 348</td>
</tr>
<tr>
<td>53-73</td>
<td>ARG8</td>
<td>aa 113</td>
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<tr>
<td>59-09</td>
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<td>aa 116</td>
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<tr>
<td>92-53</td>
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</tr>
<tr>
<td>95-75</td>
<td>lys5</td>
<td>aa 240</td>
</tr>
</tbody>
</table>

Shown are the genes disrupted for the 11 different loci disrupted in a total of 43 mutants (derived by homologous recombination) isolated by screening 9216 mutants. The position of the Tn7 insertion for each of the insertions is given. For multiple auxotrophic mutants derived from the same fosmid (and therefore predicted to be mutants in the same gene), at least two insertions were sequenced. In all cases, as expected, the same gene was disrupted but at a different site of insertion.
though not as random as high-throughput disruption approaches, nonetheless generates, with moderate effort, more highly distributed mutations than, for example, those generated by Ty1 mutagenesis in *S. cerevisiae*. The Tn3 mutagenesis technique in *S. cerevisiae* (Ross-Macdonald et al. 1999) probably yields a similar spectrum of mutations as our Tn7 method, although no analysis (similar to that presented here) of rates of Tn3-mutagenized linear fragments has been reported for *S. cerevisiae*. However, given the low rates of nonhomologous recombination in *S. cerevisiae* compared with *C. glabrata*, it is likely that the majority of transformants in *S. cerevisiae* resulting from transformation with a linear genomic fragment containing a Tn3 insertion are, in fact, allele replacements. The complications in our mutagenesis strategy resulting from the isolation of three classes of *C. glabrata* transformants were related to recombination parameters, which were specific to *C. glabrata*. Lastly, the ratio of homologous to nonhomologous recombinants is likely to be species-specific and the utility of Tn7 as a mutagenesis method in other organisms will certainly depend in part on the efficiency of homologous recombination and equally importantly on the ratio of homologous to nonhomologous recombination.

The R6K* origin that we engineered into the transposon facilitates recovery of the transposon and associated flanking genomic DNA for any insertion of interest. The R6K* origin is silent in any *E. coli* strain lacking the o-ring proteins; thus, fosmids carrying the Tn7 insertion could be maintained stably in DH5αMCR. When cloning genomic DNA flanking the transposon, we transform the permissive host and easily recover the transposon and associated DNA.

The insertions that we isolated are targeted to a diverse subset of genes in *C. glabrata*. Among the mutations that we found among the first 9216 generated by this method were 52 amino acid auxotrophs with insertions in 11 genes known from *S. cerevisiae* to be required for amino acid prototrophy. We also found insertions in two loci that profoundly altered susceptibility to the antifungal fluconazole. The first of these was in CgCDR1, the *C. glabrata* ortholog of the *S. cerevisiae* PDR5 gene, which encodes an ABC transporter (Balzi et al. 1994). This gene has previously been shown to be required for resistance to azoles (Miyazaki et al. 1998; Sanglard et al. 1999); the second was caused by a nonhomologous recombination event and has not been characterized further. We identified 38 fluconazole-resistant mutants that will be characterized elsewhere.

Our analysis of small pools of insertion mutants indicated that the mutagenesis method is, indeed, random. Southern analysis of pools of ~40 insertions in *C. glabrata* derived from pools of 150 insertions in *E. coli* showed that all the bands in *E. coli* were, in fact, present at approximately the same intensity in the pool of *C. glabrata* mutants. This indicated strongly that most mutants in a pool of insertions in *E. coli* recombine at approximately equal frequency into the *C. glabrata* genome. However, because each band on the Southern could in theory correspond to any one of many different insertions in that fragment, it is possible that each band on the Southern in the *C. glabrata* lane was, in fact, caused by differential recombination (poisoning) by one or a small subset of the many insertions in that fragment present in the *E. coli* library (each of which contributes to the corresponding band in the *E. coli* lane). We view this formal possibility as unlikely given the randomness of the insertions that we have analyzed by sequencing. For 21 different loci, we isolated and sequenced multiple insertions giving the same phenotype (as many as 7 insertions for the same locus). For none of these 54 mutants were the insertion sites the same, implying that for a variety of loci, multiple insertions in the same genomic fragment (in the *E. coli* library) all recombine into the *C. glabrata* genome. Our characterization of ~100 initial insertion mutants indicates in other ways that the method is, indeed, random. First, three-fourths of the insertions generated in *C. glabrata* were in coding regions of the genome, which corresponds well with the ratio of coding to noncoding regions in the *S. cerevisiae* genome. Second, we identified 11 loci giving tight auxotrophic phenotypes in a mutagenesis that covers 20%–25% of the genome. Extrapolating from *S. cerevisiae*, in which disruptions of ~65 genes (taken from the SGD database) show tight auxotrophic phenotypes on media lacking amino acids, we might have expected to identify ~14 mutants.

The presence of the R6K* origin in the Tn7 transposon permits the rapid rescue of a plasmid containing the Tn7 transposon and DNA flanking the insertion site. This plasmid can be used to obtain sequence information for the locus disrupted. Equally importantly, this rescued plasmid can be used (via homologous recombination and allele replacement) to introduce the mutation into a clean background and formally verify that the phenotype in question is the result of the Tn7 insertion. Lastly, although they are not used for analysis here, the *C. glabrata* mutants are all barcoded with short oligonucleotide tags (see Cormack et al. 1999 and Methods). This will permit parallel analysis of the mutants in future screens.

This method of mutagenesis is potentially broadly applicable to other organisms, requiring only that the organism be transformable and that transforming DNA integrate via homologous recombination. Its advantages over chemical methods of mutagenesis are obviously in the rapidity of the subsequent analysis of mutants with phenotypes of interest. We would also suggest that it has potential advantages over in vivo methods of transposon mutagenesis depending on any in vivo bias for transposon insertion sites (e.g., the bias to insertion sites upstream of RNA polymerase III promoters or in silent chromatin displayed by the *S. cerevisiae* Ty family of transposons; Ji et al. 1993; Zou et al. 1996; Kim et al. 1998). In summary, we present the characterization of a novel approach to insertional mutagenesis in the pathogenic yeast *C. glabrata*. This method yields a randomly distributed set of insertion mutations throughout the genome, which can be screened for phenotypes affecting diverse aspects of metabolism, physiology, or virulence.

**METHODS**

**Strains and Growth Conditions**

*E. coli* strain BW23473 (Δlac-169 rhaA1 creC510 hisdR514 ΔalA-$\gamma$ pir endA recA1; Metcalf et al. 1996) was used for maintenance of conditional replicons carrying an R6K* origin of replication that require expression of the protein II (the product of the *pir* gene) for replication. Otherwise, strain DH10 (GIBCO BRL) was used routinely for electroporation of plasmids or fosmids. Barcoded derivatives of *C. glabrata* strain BG14 (ura3Δ::Tn903Neo2; Cormack and Falkow 1999) were used for all the experiments described. These 96 barcoded strains are identical except that each has a unique oligonucleotide integrated in the genome, which permits tracking of the strain in a mixed pool of barcoded strains (Cormack et al. 1999).
Media

Electrocompetent cells were prepared according to Dower et al. (1988); plasmid and fosmid DNA preps were made using Qiaspin miniprep columns (QIAGEN) according to the manufacturer’s instructions. To obtain reasonable yields for fosmid DNA preps that are maintained at a single copy in *E. coli*, 10-ml cultures were grown in 2 × YT at 37°C overnight. The cell pellet was resuspended and lysed in twice the volume recommended for regular minipreps, and then passed over two Qiaspin columns. For growth of fosmid DNAs, media was supplemented with Cm to 10 mg/L and for mutagenized fosmids with Km to 30 mg/L.

Yeast cells were routinely grown on standard *S. cerevisiae* media (Sherman et al. 1986): YPD (YP supplemented with 2% final glucose concentration at 30°C). For *Ura* selection of BG14, SD plates supplemented with 0.6% of casamino acids (GIBCO) were used. To score for resistance or sensitivity to 5-fluorotic acid (5-FOA), YNB plates were supplemented with a 500 µg/mL final concentration of hygromycin B. The phenotype of the insertion mutants was scored by growth on YPD plates supplemented with a 2.0-kb hygromycin-resistance cassette containing (sequence deposited at GenBank accession number) after addition of EDTA (10 mM) to stop the digestion, fractions were analyzed by gel electrophoresis to find the sample in which the average molecular weight of the DNA was well above 40 kb. This sample was treated with shrimp alkaline phosphatase (Episcience), and the phosphatase was inactivated by heat. To generate the fosmid arms, pBAC-Bcg3 was digested with HindIII and phosphatased with calf intestinal phosphatase (Boehringer Mannheim). The phosphatase was removed by phenol extraction, and the sample was redigested with BamHI to generate one of the fosmid arms (a HindIII-BamHI fragment). The second fosmid arm (EcoRI-BamHI) was generated by digesting the fosmid with EcoRI, followed by treatment with calf intestinal phosphatase. The phosphatase was removed by phenol extraction, and the product was digested with BamHI. These two fosmid arms were ligated to the genomic DNA; the resultant ligation was packaged using XL packaging extract (Stratagene) and used for infection of DH5αMCR cells. Individual Cm® transformants were picked and arrayed in 96-well plates. For 700 of these, fosmid DNA was prepared from 10-ml cultures using the REAL prep kit in a 96-well format (Stratagene). The library is highly representative. We picked 5500 fosmids (~15 × genome coverage) and analyzed by PCR how many times particular genes were present. The majority of genes were present between 10 and 15 times (TRP1, 16 times; HIS3, 15 times; YERO19w homolog, 12 times; EPA1, 7 times).

**Plasmid Constructions**

**pBC166.2**

pBC166.2 is derived from plasmid pMCB40 ([porKΔΔΔ::miniTn7 spe1·KmR·NorI](http://www.genome.org)). This plasmid contains the conditional origin of replication R6KΔ and a mini-Tn7 element conferring resistance to kanamycin (Biery et al. 2000). We first introduced the *URA3* gene of *S. cerevisiae* into this mini-Tn7. We PCR-amplified the gene with primers 341 and 343 containing NorI sites. The PCR product was digested with NorI and ligated to pMCB40 to generate plasmid pBSC166.2.

**pIC6**

Plasmid pBC166.2 was used as template to PCR-amplify a 398-bp fragment containing the R6KΔ origin of replication, using primers S38 and S39 containing XhoI sites; the PCR product was digested with XhoI and ligated to Spel-digested pBSC166.2. This plasmid contains two R6KΔ origins of replication: the original one in the backbone and the newly cloned copy within the Tn7 element. To remove the backbone R6KΔ origin, plasmid pIC2 was digested with *AatI* and *PmlI*, blunted, ligated, and transformed into the permissive strain BW23473. The plasmid generated, pIC6, carries the element Tn7 URA3 (Tn7·URA3·KmR·Spe1). This minitransposon contains within it a yeast-selectable marker (*URA3*), a kanamycin-resistance cassette, and the conditional origin of replication R6KΔ. The element carries the only origin of replication of the plasmid, and must be maintained in strain BW23473 (Metcal et al. 1996), which expresses the *Aur* protein from a chromosomal fusion to the *uidA* gene.

**pIC31**

We then modified pIC6 by cloning the *lhp* gene conferring resistance to hygromycin B (Tn7·UKR-H) into the Tn7 element. To do this, we first destroyed one of the two NorI sites in pIC6 by partially digesting, blunting, and religiating. This generated plasmid pIC28, which contains only one NorI site (downstream from *URA3*). We then PCR-amplified the *K. pneumoniae* lhp gene conferring resistance to hygromycin B (Gritz and Davies 1983) from plasmid pAG26 (Goldstein et al. 1999); the PCR product was cloned into the PGK1 gene promoter from *S. cerevisiae* P-PRG6 in a derivative of pRS316 to generate plasmid pAP358. *C. glabrata* transformants with this plasmid grow on YPD plates supplemented with 200 µg/mL of hygromycin B. The 2.0-kb hygromycin-resistance cassette from plasmid pAP358 was excised by digesting with SacI

**Construction of Fosmid Library**

The plasmid pBAC carries the *F* origin, the *cat* gene conferring Cm resistance, and a cos site for packaging (Kim et al. 1992). We modified pBAC by introducing two *BglII* sites immediately flanking the *BamHI* site. This was accomplished by first digesting pBAC with *NorI* and then cloning a double-stranded DNA oligonucleotide obtained by allowing oligonucleotides *BglII* (**GCGCCGCGAATTATGCTGGC**GATCCGCGAATTATGCTGGC) to anneal. The final vector, pBAC-Bcg3, is similar to the original, but the *BamHI* cloning site is now immediately flanked by *BglII* sites (sequence deposited at GenBank accession number). The advantage of this is that once a fragment of the genome has been cloned into the *BamHI* site, it can be precisely released with *BglII*, which cleaves 7 bases downstream of its site (inside the cloned genomic insert). Thus, the genomic DNA ends generated by *BglII* digestion are precisely homologous to the genome, with no nucleotides derived from the pBAC vector.

Genomic DNA from strain BG14 was carefully prepared according to Goshorn et al. (1992). The DNA was digested with decreasing amounts of *SacI* and after addition of EDTA (10 mM) to stop the digestion, fractions were analyzed by gel electrophoresis to find the sample in which the average molecular weight of the DNA was well above 40 kb. This sample

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*Genome Research* 913

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and KpnI, blunted, and ligated to pUC28 cut with NotI and blunted, generating plasmid pIC31. This plasmid carries mini-
transposon Tn7 UKR (URA3·HphI·KmR·R6K), which contains the URA3 selectable marker, the kanamycin-
resistance gene, the conditional origin of replication R6K, and the hygromycin-resistance cassette.

pIC3.3::Tn7-UKR
One random insertion of the Tn7 UKR transposon in one arbitrarily chosen fosmid was used to generate a smaller plasmid. The mutagenized fosmid was digested with EcoRI and religated; the resulting plasmid is an 8.4-kb conditional repli-
con consisting of a 5.3-kb EcoRI C. glabrata genomic frag-
ment with a Tn7 UKR insertion.

In Vitro Mutagenesis
In vitro transposition reactions were performed as described by Biery and Craig (Biery et al. 2000) using purified trans-
posase proteins (TnsA, TnsB, and TnsC<sup>25V</sup> purchased from New England Biolabs). For fosmid mutagenesis, we first pre-
pared fosmid templates from 10-mL cultures using the QIA-
GEN midiprep kit according to the manufacturer’s instructions.

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pared fosmid templates from 10-mL cultures using the QIA-
GEN midiprep kit according to the manufacturer’s instructions.

We used 400 ng of target fosmid DNA (0.12 nM), and 100 ng of donor plasmid carrying the transposon: pIC31 or pIC6 (0.19 and 0.22 nM, respectively). The reaction was stopped by phenol extraction and precipitation, and the DNA was resus-
pended in 10 µL of TE. Then 1 µL of this mutagenesis mix was used to transform DH10 electrocompetent cells, and trans-
formants were selected on media containing chloramphenicol (10 mg/ml) and kanamycin (50 mg/ml). Only those fosmid
molecules into which the mini-Tn7 transposed are able to replicate in this strain, because donor molecules contain only
the conditional R6K origin, and unmutagenized fosmid is not resistant to kanamycin. For each fosmid, a minimum of
200 and a maximum of tens of transformants were pooled and grown in a 100-ml culture to saturation at 30°C. This culture was used to prepare fosmid DNA using the QIAGEN midiprep kit according to the manufacturer’s in-
nstructions.

Classification of C. glabrata Transformants
To introduce the mutagenized fosmids into C. glabrata, we used DNA from 96 mutagenized fosmids to transform the 96 oligonucleotide-tagged (barcoded) derivatives of BG14 (ura3Δ37;Tn903Neo<sup>a</sup>). First, 1–2 µg of a mutagenized fosmid DNA was digested with either BglII or with MfeI prior to trans-
formation. The transformation mixes were plated on SD
plates lacking uracil, and Ura<sup>+</sup> isolates were scored after
growth at 30°C for 2–3 d. Ura<sup>+</sup> transformants fall in three
different classes based on the phenotype displayed on 5-FOA
plates. The first class of unstable (U) transformants is com-
posed of colonies that are able to grow both on 5-FOA plates
as well as on plates lacking uracil, therefore appearing as Ura<sup>+</sup>
5-FOA<sup>a</sup> colonies. The second class of partially stable (PS)
transformants refers to colonies that appear to be Ura<sup>+</sup> FOA<sup>a</sup>;
however, upon incubation at 30°C (48–72 h), patches of the
transformant engender a small number of 5-FOA<sup>a</sup> papillae on
the 5-FOA plate. The third class of stable (S) transformants
consists of colonies that display the expected Ura<sup>+</sup> 5-FOA<sup>a</sup>
phenotype without papillation even after prolonged incuba-
tion on 5-FOA plates. For any given fosmid tested, the percent of transformants derived from each class of transformant did
not change by altering parameters of the transformation pro-
tocol; the parameters tested were inclusion or omission of
DMSO and varying the time (15 min to 1 h) and temperature
(from 37°C to 45°C) of the heat-shock temperature (data not shown).

Killing of Untransformed Background and Abortive
Transformants of C. glabrata
C. glabrata was transformed to Ura<sup>+</sup> using linearized, in vitro
mutagenized fosmids. Transformants were picked and ar-
ranged on a plate lacking uracil, allowed to grow overnight at
30°C, and printed onto 5-FOA plates. The majority of Ura<sup>+</sup>
colonies tested this way were contaminated with untrans-
formed background (Ura<sup>+</sup> 5-FOA<sup>a</sup>), which are Ura<sup>+</sup> and thus
5-FOA<sup>a</sup>; therefore, most 5-FOA<sup>a</sup> transformants appeared to
give 5-FOA<sup>a</sup> papillae. To identify transformants that were
truly 5-FOA<sup>a</sup>, it was necessary to eliminate these contaminat-
ing cells. This could be easily accomplished by colony-
purifying the transformants on plates lacking uracil. How-
ever, we decided this was not practical for thousands of trans-
formants. Therefore, a step to screen for hygromycin-resistant
transformants was introduced. Transformants patched on
plates were replica-plated onto a hygromycin-containing YPD
plate, allowed to grow, and printed a second time onto hy-
gromycin plates; patches were subsequently printed onto
5-FOA plates and incubated at 30°C for 48–72 h. After these
two consecutive prints on hygromycin plates, all of the un-
transformed background was killed, because they are sensitive
to hygromycin. In control experiments, identical results were
obtained by streak purification and replica-printing onto hy-
gromycin plates: that is, the bona fide Ura<sup>+</sup> 5-FOA<sup>a</sup> colonies
identified by colony purification were also identified by print-
ing onto hygromycin plates (data not shown).

Consideration of Multiple Mutants Derived
From a Single Fosmid
Because the majority of our stable recombinants are derived by homologous recombination, the expectation is that if a
fosmid contains a gene that can mutate to a given phenotype,
them in a pool of 100 transformants derived from that fosmid,
there will likely be multiple mutants, representing indepen-
dent insertions in the same gene and giving the same pheno-
type. Given that we have mutagenized only 20%–25% of the
genome, most fosmids analyzed here will not physically over-
lap. Therefore, the number of fosmids from which mutants
with a phenotype were derived, rather than the number of
mutants themselves, gives a good indication of the number of
loci identified. Thus, for auxotrophs, we identified 52 mu-
tants, but these were derived from 20 fosmids (between 1 and
9 auxotroph mutants isolated per fosmid), and we therefore
estimate that our mutants identify ~20 loci. For fluconazole
sensitivity, we identified 15 insertions, but all derived from
two fosmids. We therefore estimate that our mutants identify
only two loci. That this line of reasoning is valid for the
majority of cases is indicated by the fact that we have semceived the insertions for up to three independent auxo-
trophs derived from the same fosmid; in 5/5 cases, mutants
from the same fosmid with the same phenotype represented
independent insertions in the same gene; in contrast, auxo-
trophic mutants derived from different fosmids were all in
different genes.

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914 Genome Research
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