

Evolution in *Candida albicans* Populations During a Single Passage Through a Mouse Host

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ABSTRACT

The mechanisms and rates by which genotypic and phenotypic variation is generated in opportunistic, eukaryotic pathogens during growth in hosts are not well understood. We evaluated genomewide genetic and phenotypic evolution in *Candida albicans*, an opportunistic fungal pathogen of humans, during passage through a mouse host (*in vivo*) and during propagation in liquid culture (*in vitro*). We found slower population growth and higher rates of chromosome-level genetic variation in populations passaged *in vivo* relative to those grown *in vitro*. Interestingly, the distribution of long-range loss of heterozygosity (LOH) and chromosome rearrangement events across the genome differed for the two growth environments, while rates of short-range LOH were comparable for *in vivo* and *in vitro* populations. Further, for the *in vivo* populations, there was a positive correlation of cells demonstrating genetic alterations and variation in colony growth and morphology. For *in vitro* populations, no variation in growth phenotypes was detected. Together, our results demonstrate that passage through a living host leads to slower growth and higher rates of genomic and phenotypic variation compared to *in vitro* populations. Results suggest that the dynamics of population growth and genomewide rearrangement contribute to the maintenance of a commensal and opportunistic life history of *C. albicans*.

OPPORTUNISTIC pathogens such as *Candida albicans* often reside in the host as benign, commensal organisms until the immune system is weakened. In patients undergoing organ transplants or chemotherapy, or when indigenous competitors are eliminated upon antibiotic treatment, opportunistic pathogens may gain access to vulnerable tissues, causing death in $\leq 50\%$ of infected patients (WILSON *et al.* 2002). Consequently, it is important to understand the genetic mechanisms underlying the survival and adaptation of opportunistic pathogens to growth in host environments (MARGOLIS and LEVIN 2007). Here, we used a genomewide array of single nucleotide polymorphisms (SNPs) to characterize the rates of genetic and phenotypic evolution accompanying the growth of *C. albicans* in contact with a mammalian host and compared these to rates of evolution during *in vitro* growth.

Genome evolution during interactions with hosts varies considerably across different microbial pathogens. The specific genome rearrangements leading to phase change and antigenic switching that allow patho-

gens to evade host immune responses are well described for only a few pathogens such as trypanosomes (BORST and RUDENKO 1994) and Plasmodium (KYES *et al.* 2001). Obligate intracellular symbiotic microbes, such as Buchnera (MORAN 1996) and Pneumocystis (STROBEL and ARNOLD 2004), propagate asexually and often carry a minimal but stable genome, making them wholly dependent on life within their hosts (WREN 2000). Although both opportunistic and obligate pathogens commonly propagate by asexual means, these organisms often maintain large genomes and generate substantial genomic and phenotypic variation via genome rearrangements (VICTOIR and DUJARDIN 2002; KLINE *et al.* 2003) and heritable silencing at telomeres (CROSS *et al.* 1998; BORST 2002; GUPTA 2005). Given that many commensal and apparently harmless symbionts may become invasive pathogens in immunocompromised hosts, the mechanisms underlying the maintenance of genetic variation and of the commensal state bear investigation (LEVIN *et al.* 2000).

As the most common commensal fungus of the human microbial flora, *C. albicans* provides a model for the study of opportunistic pathogens because it reproduces primarily asexually and demonstrates a high degree of genetic and genomic variability among isolates (COWEN *et al.* 1999; IWAGUCHI *et al.* 2000; JOLY *et al.* 2002; PUJOL *et al.* 2002; LEGRAND *et al.* 2004). The complete genome sequence revealed high levels of

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heterozygosity (~4%) across the 16-Mb diploid genome (JONES *et al.* 2004; VAN HET HOOG *et al.* 2007), and population-level variation has been demonstrated in clinical populations from different continents, regions, hospitals, and families (FORCHE *et al.* 1999; PUJOL *et al.* 2002; BOUGNOUX *et al.* 2006). However, the genome and population processes underlying observed variation in host populations is not well understood. Appreciable rates of mitotic recombination estimated at specific genome regions (LEPHART *et al.* 2005; LEPHART and MAGEE 2006) and in repetitive regions (ZHANG *et al.* 2003) have been evaluated primarily from *in vitro* cultures. Chromosomal variation as well as point mutations accumulate rapidly in populations evolving resistance to azole antifungal drugs (SELMECKI *et al.* 2006; COSTE *et al.* 2007), and in a few cases, the evolution of a pathogen within the same individual has been studied over the time course of antifungal drug treatment (LOPEZ-RIBOT *et al.* 1998; MARR *et al.* 1998; COSTE *et al.* 2007; SELMECKI *et al.* 2008). Together, clinical studies reveal the accumulation of variation in host-associated populations, but the evolutionary relationship among isolates is not clear, and the number of isolates obtained during the course of infection are insufficient to allow a comprehensive view of population dynamics.

With the goal of understanding mechanisms by which genetic and phenotypic variation arise as a pathogen propagates in its host, we tracked genomewide dynamics in *C. albicans* populations during passage through a susceptible host (*in vivo*) and compared results to populations propagated in liquid culture (*in vitro*). We first asked if population growth rates differ when cells are grown in a mammalian host relative to when they are grown in liquid culture. We then compared the rates and types of short- and long-range loss of heterozygosity (LOH) events that arose during *in vivo* relative to *in vitro* propagation. Finally, we determined the rates and types of phenotypic variation in colony growth that arose during *in vivo* and *in vitro* propagation. To conduct the analyses, we exploited the counterselectable marker *GALI*, measured recombination as LOH using genome-wide SNPs, and evaluated changes in chromosome copy number using competitive genome hybridization (CGH) (FORCHE *et al.* 2005; SELMECKI *et al.* 2005). We found fivefold lower population growth rates and distinctly different genome dynamics arising in response to growth *in vivo* compared to growth *in vitro*. Furthermore, we found that variation in *C. albicans* colony size and morphology arose during *in vivo* propagation only and was positively associated with short-range and chromosome-level recombination events. Taken together, our results suggest that passage through a mammalian host is accompanied by slow population growth and elevated levels of genetic and phenotypic variation relative to the rates of variation observed with propagation in the laboratory.

TABLE 1

The number and chromosomal location of heterozygous SNP loci analyzed in reference strains SC5314 and AF7

Chromosome	Strain SC5314	Strain AF7
R	19	19
1	23	13
2	19	19
3	11	10
4	21	20
5	12	12
6	11	11
7	7	6
Total	123	110

SNPs heterozygous in AF7 were analyzed for all experimental strains.

MATERIALS AND METHODS

Strains and media used in this study: To study recombination events across the entire *C. albicans* genome, we used a system with two components: a heterozygous counterselectable *GALI* marker that permitted selection of isolates in which LOH at *GALI* had occurred and 123 SNP loci positioned ~100 kb apart across most of the genome (Table 1; supporting information, Table S1). *C. albicans* strain AF7 is a derivative of sequenced strain SC5314 in which one copy of *GALI* was replaced with *URA3* to generate the *GALI/galI* heterozygous locus (*GALI/ΔgalI::URA3*) (FORCHE *et al.* 2003). *Gal*⁺ strains are sensitive to 2-deoxygalactose (2DG^s) because metabolism of 2DG results in a toxic product (PLATT 1984) but grow on media with galactose as the sole carbon source. *Gal*⁻ cells lack a functional *GALI* locus and are 2DG resistant (2DG^r), but do not grow on media with galactose as the sole carbon source.

Colonies of *C. albicans* were routinely grown on the non-selective YEPD medium (1% yeast extract, 1% yeast peptone, 2% glucose; 1.5% agar for plate cultures). To distinguish between *Gal*⁻ (2DG^r) and *Gal*⁺ (2DG^s) phenotypes, strains are plated onto a synthetic 2-deoxygalactose medium (2DG; 0.67% yeast nitrogen base without amino acids, 0.1% 2-deoxygalactose, 1.5% agar) and counterselected on synthetic galactose medium (0.67% yeast nitrogen base without amino acids, 2% galactose, 1.5% agar).

***In vivo* populations:** In a previous study, 10⁶ cells of the parent strain AF7 (*GALI/ΔgalI::URA3*) were injected into the tail vein of 13 outbred ICR male mice (22–25 g; Harlan, Indianapolis) (FORCHE *et al.* 2003). Mice were observed and, when moribund at 5–7 days, anesthetized using isoflurane and euthanized, and both kidneys were removed. Kidneys were combined, homogenized with 1 ml of water, and dilutions at 1:1000 of the kidney homogenate were plated onto YEPD medium to obtain total colony counts. The same homogenate was diluted 1:10 and plated onto 2DG medium to obtain *Gal*⁻ colony counts at 3 days. At 3 days, *Gal*⁻ colonies arising by mutation do not grow to an observable size but colonies are apparent for a control *ΔgalI/ΔgalI* strain. (FORCHE *et al.* 2003).

***In vitro* populations:** To generate *in vitro* populations, strain AF7 was streaked out on synthetic galactose medium and grown 2 days at 30° to obtain single colonies, which were removed and transferred to each of 20, 5 ml YEPD liquid cultures (BECKERMAN *et al.* 2001; SPELL and JINKS-ROBERTSON 2004; MOOKERJEE and SIA 2006). These were grown for 16 hr in a roller tube incubator at 30°. The cultures were spun down,

washed once with sterile water, and resuspended in 1 ml of sterile water. The resuspended cells were then plated onto YEPD plates at 10^{-7} dilutions and grown at 30° for 2 days to obtain the total cell count and plated on 2DG plates at 10^{-3} dilution and grown at 30° to obtain a count of Gal⁻ colonies at 3 days.

Diagnostic PCR to determine GALI status: Diagnostic PCRs were carried out using primers flanking the *GALI* locus to determine if Gal⁻ phenotypes (2DG^R) obtained in the above experiments were due to loss of the remaining *GALI* copy or due to mutation in the *GALI* ORF. Primers gal1-detF, ura3-detR, and 2020 were used for upstream and primers 2278, 2279, and 2280 were used for downstream diagnostic PCR (Table S1). Total genomic DNA extractions were carried out as described previously (BECKERMAN *et al.* 2001). PCRs were carried out in a total volume of 25 μ l with 10 mM Tris-HCl (pH 8.0); 50 mM KCl; 3 mM MgCl₂; 100 μ M each dATP, dCTP, dGTP, and dTTP; 2.5 units Taq polymerase (rTaq, TAKARA); either 5 or 10 μ mol of each primer (see Table S1); and 30 ng of genomic DNA under the following conditions: initial denaturation for 3 min at 94° , 30 cycles of a denaturation step for 1 min at 94° , a primer annealing step for 30 sec at 54° , and an extension step for 1 min at 72° . The final extension step was 5 min at 72° . PCR fragments were size fractionated on agarose gels [1% in $1\times$ TBE (0.89 M Tris; 0.02 M EDTA-NA₂H₂O; 0.89 M boric acid)] and compared to the size of products from positive control strains [SC5314 (*GALI/GALI*) and AF7 (*GALI/gal1Δ*)] and the negative control strain [AF27 (Δ gal1/ Δ gal1)] (FORCHE *et al.* 2003).

Population growth rate: For *in vivo* experimental populations, we used two methods to obtain estimates of net population growth rate and the number of cell divisions. First, we obtained and analyzed published results from a careful time-course experiment tracking *C. albicans* population growth *in vivo* (MACCALLUM and ODDS 2004). Our study used strain genotypes and inoculation cell numbers comparable to those used in that study (MACCALLUM and ODDS: SC5314 at 5×10^4 cells/g body weight of BALB/c mice; our study: AF7 at 4×10^4 cells/g body weight of ICR mice), both with tail-vein infection. Strain AF7 is derived from SC5314, is prototrophic, shows no growth rate differences, and is not attenuated in virulence in comparison to SC5314 (FORCHE *et al.* 2003). Numbers of colony-forming units (CFUs) in kidney tissues over time were log transformed and regressed against time (minutes) in Microsoft Excel 2004, version 11.3.7. Obtaining an increasing function, the slope of the regression of ln(cell number) *vs.* time for the period of 2–72 hr after inoculation provided an estimate of the intrinsic population growth rate, *r*. The number of cells generated per parent cell during time, *t*, of increasing growth was calculated as e^{rt} , and the number of doublings (cell divisions) was calculated by solving for $x = \log_2(e^{rt})$. A second estimate of population growth *in vivo* was made using total colony counts from homogenized kidney tissue and by estimating the number of doublings (cell divisions) required to produce the observed increases in cell numbers: $x = \log_2(e^{rt})$. We assumed a bottleneck population size as a fraction of the final population size (MACCALLUM and ODDS (2004) to estimate starting population sizes in the kidney.

For *in vitro* experimental populations, cell numbers at the start and end of population growth were counted as CFUs on YEPD plates and population growth rates were estimated as described above for *in vivo* experiments.

Rates of LOH at GALI: For both the *in vivo* and *in vitro* experiments, we used LEA and COULSON's (1949) method of the median to estimate the rate at which the heterozygous *GALI*/ Δ gal1 (2DG^S) locus is converted to the homozygous state, Δ gal1/ Δ gal1 (2DG^R). The method uses the variance

among independent cultures for the proportion of 2DG^R cells generated relative to the total number of cells generated over the same time to estimate the rates of the event per generation (Table S2). The method accounts for differences of *in vitro* and *in vivo* growth rates and permits direct comparison of LOH rates at *GALI*.

SNP microarray analysis: We expanded a previously described SNP microarray (FORCHE *et al.* 2005), adding 98 SNP loci to cover a total of 123 SNP loci (Table 1). The SNP loci are positioned ~100 kb apart across the eight *C. albicans* chromosomes except for regions of low polymorphism in the fully sequenced genome of strain SC5314 (Chr3 right arm, Chr7 left arm, and telomere distal regions of ChrR (VAN HET HOOG *et al.* 2007). Design of allele-specific oligonucleotides, probe generation, slide preparation and hybridization, and data analysis was performed as described earlier (FORCHE *et al.* 2005). Four multiplex PCR reactions with 28–34 primer pairs/reaction (Table S1) were carried out for each *C. albicans* isolate, using 60 ng of genomic DNA and a Qiagen Multiplex PCR kit following the manufacturer's instructions (Qiagen, Valencia, CA).

The SNP genotypes were determined using the allelic fraction (AF), which describes the relative intensity of the control probe (SC5314: labeled with Cy5) compared to the experimental probe (test strain: labeled with Cy3) in competitive hybridizations to the microarray according to the protocol detailed in FORCHE *et al.* (2005). The AF values are normalized to give an expected value of 0.5 for heterozygous loci. The distribution of AF values for homozygous and heterozygous loci and the cutoff values to score homozygous states were as described previously (FORCHE *et al.* 2005). For each individual SNP locus, AF values between 0.40 and 0.60, inclusive, were scored as heterozygous, AF values <0.40 were scored as allele 1 homozygous, and AF values >0.60 were scored as allele 2 homozygous.

We used SNP genotypes and information on their physical linkage to infer chromosome-level rearrangement and altered ploidy. Individual SNP loci were scored as potentially homozygous, heterozygous, or trisomic by AF values as follows: 0–0.342 (homozygous for a1); 0.343–0.422 (trisomy; two copies of a1 and one copy of a2); 0.423–0.573 (heterozygous; a1/a2); 0.574–0.657 (trisomy; 2 copies of a2 and 1 copy of a1); and 0.658–1.0 (homozygous for a2) (data not shown). Where AF values suggested altered ploidy at linked SNP loci over large segments or entire chromosomes, we confirmed ploidy levels using CGH (SELMECKI *et al.* 2005; LEGRAND *et al.* 2008) as described below.

Genomic distribution of short-range LOH events: For *in vivo* and *in vitro* populations, we counted the number of short-range LOH events occurring on each chromosome and compared the observed number of LOH events to that expected for a random distribution across the genome within each population. We tested the significance of higher or lower numbers of observed events compared to that expected using a binomial test (SOKAL and ROLF 1981) with an expected value set at the average number. Because the number of events detected is very low relative to the number of loci evaluated and because the binomial test lacks power, we also conducted a permutation test. Here, we randomly permuted loci (columns in Table S3) across all chromosomes and divided columns into "chromosomes" with the same number of loci as in the empirical data set. Repeating permutations, we generated 1000 random distribution data sets. Significance was evaluated as the number of times that the observed number of events occurred in the permuted data sets, divided by 1000, the total number of randomized data sets.

Comparative genome hybridization: Comparative genome hybridization uses a microarray format with probes for all of

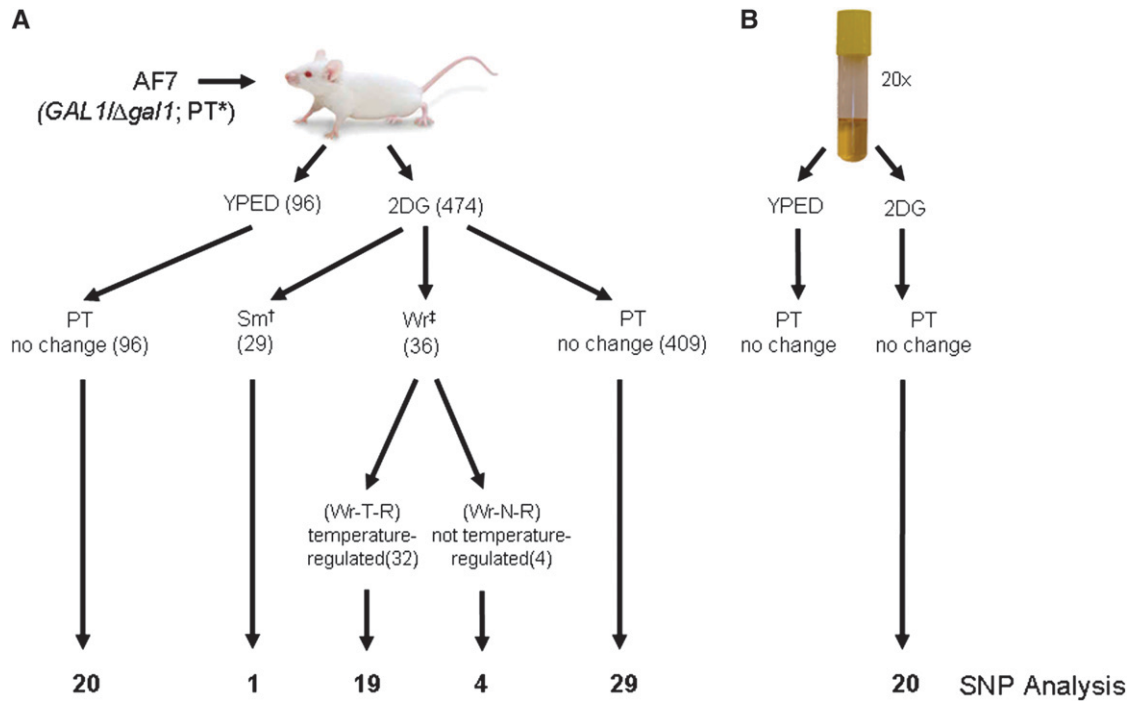


FIGURE 1.—Propagation of *C. albicans* populations for *in vivo* and *in vitro* experiments, media used for isolation (YEPD, 2DG), and occurrence of altered colony phenotypes. Numbers of strains isolated for study are shown in parentheses, and numbers of isolates analyzed using the SNP microarray are shown at the bottom. (A) *In vivo* experiments. Colony growth phenotype indicated as PT*, parental type; SM^f, small colony; and Wr⁺, wrinkly colony. Wr-T-R, “wrinkly, temperature regulated”; Wr-N-R, “wrinkly, not regulated by temperature.” (B) *In vitro* analysis. Twenty strains isolated for SNP analyses from 2DG plates. PT colony growth phenotype observed for all isolates.

the ~6800 *C. albicans* ORFs and detects changes in gene copy number using hybridization signal intensity (SELMECKI *et al.* 2005). CGH was carried out as described previously (SELMECKI *et al.* 2005) using strains SC5314 and AF7 as heterozygous, disomic reference strains. An updated version of the Ch_map program, provided by Sven Bergmann (SELMECKI *et al.* 2005), was used to visualize combined SNP and CGH results.

Colony growth phenotype: Colonies obtained from experimental populations were streaked for single colonies on YEPD plates, and colony morphology was scored as follows: the parental colony phenotype (PT) that is indistinguishable from that of AF7, the small colony phenotype (Sm), and the wrinkly colony phenotype (Wr). In addition, isolates were grown for 2 days at 23°, 32°, and 37° in duplicate YEPD plates. Strains that formed wrinkled colonies at all three temperatures were designated as “wrinkling not regulated by temperature” (Wr-N-R). Strains that formed smooth yeast colonies at 23° and formed filamentous, wrinkled colonies at 32° and 37° were designated as “wrinkling that is temperature regulated” (Wr-T-R). The frequency of each phenotype in experimental populations was recorded, and plates were photographed.

RESULTS

Our overall goal was to estimate and compare rates at which genetic and phenotypic variation is generated in *C. albicans* populations propagated *in vivo* and *in vitro*. From the *in vivo* experimental populations, 96 single-colony isolates (strains) were obtained from nonselective YEPD plates (*Gal*⁺, 2DG^S) and 474 strains were isolated from 2DG plates (*Gal*⁻, 2DG^R) for further study.

From the *in vitro* experiments, 20 2DG^R strains were isolated to serve as a control population (Figure 1). In the results reported below, *GALI* status of single colony isolates was determined by plating on selective media and by diagnostic PCR (data not shown; see MATERIALS AND METHODS).

***In vivo* populations:** We estimated the total CFUs at 8.0×10^4 – 1.7×10^6 cells/g kidney across kidneys of 13 infected mice (FORCHE *et al.* 2004). Because kidney tissues were thoroughly homogenized, we assumed that each CFU represented a single nucleate hyphal or yeast cell. We estimated the proportion of cells that had undergone LOH at the *GALI* locus (2DG^R) at 8.7×10^{-5} – 1.7×10^{-2} /cell for all mice. Among 474 2DG^R isolates, 65 (13.7%) exhibited colony phenotypes different from that of the parent AF7, and the remainder demonstrated AF7 PTs. The altered colony phenotypes included Sm colonies due to slow growth and Wr colonies due to filamentous cell growth compared to AF7 (Figure 1); these are fully characterized below. In contrast, none of the 96 colonies isolated on YEPD medium demonstrated altered colony growth phenotypes.

***In vitro* populations:** The proportion of cells that had undergone LOH at the *GALI* locus during *in vitro* propagation was 3.2×10^{-5} – 1.1×10^{-4} /cell across the 20 independent YEPD cultures. No altered growth phenotypes were detected among ~7500 colonies

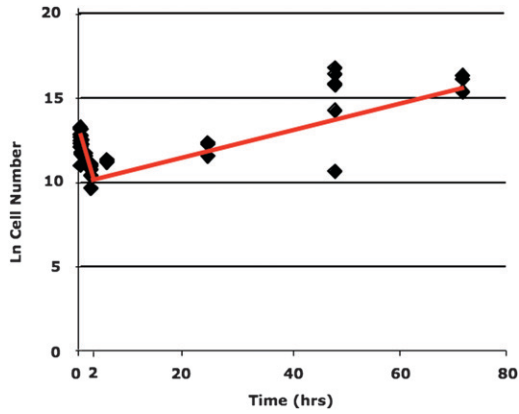


FIGURE 2.—Graph of log-phase population growth during *in vivo* propagation, ln (cell number) over time in hours. From the data of MACCALLUM and ODDS (2004).

grown on YEPD or on 2DG plates after *in vitro* propagation (Figure 1).

Population growth rates *in vivo*: We employed two different approaches to estimate growth rates for *in vivo* populations, and these yielded similar results. We report net population growth rates because rates of birth and death cannot be separately estimated. First, we estimated the population growth rate and the number of cell divisions using the published results of a careful time-course study (MACCALLUM and ODDS 2004). These data show that most *C. albicans* cells are cleared from the blood and that population numbers in the kidney greatly decline during the first 2 hr after inoculation (MACCALLUM and ODDS 2004). The estimated population bottleneck size was 0.45% of the final population size at 2 hr. After 2 hr, populations in the kidneys demonstrated log growth, and the slope of the regression of ln(cell number) *vs.* time for the period of 2–72 hr after inoculation (Figure 2) provides an

estimate of the intrinsic population growth rate, r , at 1.3×10^{-3} ($r^2 = 0.75$; SE ± 0.0002). The number of cells generated during the time $t = 70$ hr of increasing growth is $e^{rt} = 3.2 \times 10^2$ cells/cell/g kidney. On average, $\log_2(320) = 8.3$ cell divisions/parent cell in the population is required to generate 320 cells.

Second, we assumed a bottleneck population size at 0.45% of the final population size for our experimental *in vivo* populations and then calculated the average net increase in cell numbers required to obtain final population sizes of 2.2×10^2 cells/cell/g kidney. The estimated number of cell divisions is $\log_2(220) = 7.8$ cell divisions/parent cell in the population, a value similar to that obtained above.

Rates of cell division are slower *in vivo* than *in vitro*: The average rate of cell division for *in vivo* populations estimated from data of MACCALLUM and ODDS (2004) is 8.3 cell divisions over 70 hr, or 0.12 generations/hr. From our experimental *in vivo* data, we estimated a rate of 7.8 cell divisions over 5 days, or 0.065 generations/hr, which is even slower. For *in vitro* populations, we used direct counts of total CFUs before and after growth in liquid culture to estimate an average 9.9 cell divisions/parent cell and an average rate of cell division over the 16-hr experiment at 0.62 divisions/hr. Thus, net population growth rates were 5- to 10-fold slower *in vivo* than *in vitro*.

Rates of recombination at *GAL1* are higher *in vivo* than *in vitro*: For both the *in vivo* and *in vitro* populations, we used LEA and COULSON’S (1949) method of the median to estimate the rate at which the heterozygous *GAL1*/ $\Delta gal1$ locus is converted to the homozygous state, $\Delta gal1/\Delta gal1$ (Table 2; Table S2). For the *in vivo* populations, we estimated the total CFUs at 8.0×10^4 – 1.7×10^6 cells/g kidney across the 13 mouse populations. We estimated the proportion of cells that had undergone LOH at the *GAL1* locus to generate 2DG^R cells at $8.7 \times$

TABLE 2

Number, type, and rates per generation (\pm SD) of genome rearrangement in strains derived from *in vitro* and *in vivo* experiments

Type of events	<i>In vivo</i> experiment		<i>In vitro</i> experiment	
	No. of events	Rate ^a	No. of events	Rate ^a
<i>GAL1</i>		1.7×10^{-4} ($\pm 2.2 \times 10^{-4}$)		6.0×10^{-6} ($\pm 2.0 \times 10^{-6}$)
SNP LOH	16	3.3×10^{-4} ($\pm 5.8 \times 10^{-4}$)	7	3.5×10^{-4} ($\pm 2.4 \times 10^{-5}$)
BIR	1		0	0
Whole chromosomes	5		0	0
All chromosomes	6	1.2×10^{-3} ($\pm 4.2 \times 10^{-3}$)	0	0

Standard deviation was calculated across loci (short-range LOH) or across strains (chromosome-level rates).
^a *GAL1* LOH rates were calculated as events per generation, SNP LOH rates were calculated as events per locus per generation, and chromosomal rates were calculated as events per chromosome per generation.

TABLE 3

Short-range LOH and chromosome-level events observed for strains from *in vivo* and *in vitro* experiments

Event	Chromosome	Description	Host ID	Strain
<i>In vivo</i> experiments				
Chr	R	Whole-chromosome LOH	E63(3) ^a	AF617
Chr	R	Whole-chromosome LOH	A22(15)	AF3976
Chr	R	Whole-chromosome trisomy	A22(15)	AF3977
Chr	2	BIR, ^b 11 contiguous loci	B22(8)	AF21
Chr	2	Whole-chromosome LOH	A22(15)	AF3990
Chr	2	Whole-chromosome trisomy	B63(9)	AF540
LOH	R	Single LOH at 1694/2254	A22(15)	AF656
LOH	R	Single LOH at 1645/2358	A63(8)	AF121
LOH	1	Single LOH at 2347/2406	A53(4)	AF42
LOH	1	Single LOH at 1916/2198	A53(4)	AF51
LOH	1	Single LOH at 2347/2406	B63(9)	AF160
LOH	1	Single LOH at CPH1	B63(9)	AF540
LOH	1	Single LOH at 2080/2297	B63(9)	AF169
LOH	2	Single LOH at 1760/2185	C63(5)	AF194
LOH	3	Single LOH at 2095/2215	A63(8)	AF122
LOH	3	Single LOH at 1629/1863	B63(9)	AF157
LOH	4	LOH (three continuous loci)	C63(5)	AF558
LOH	4	Single LOH at 2101/2204	C63(5)	AF558
LOH	6	Single LOH at 1372/2355	B53(5)	AF72
LOH	6	Single LOH at 1372/2355	B63(9)	AF148
LOH	7	Single LOH at 6.1345	B22(8)	AF22
LOH	Unknown	Single LOH at B5B7	A22(15)	AF3990
<i>In vitro</i> experiments				
LOH	2	Single LOH at 1775/2077	—	AF7-8
LOH	2	Single LOH at 1179/2238	—	AF7-14
LOH	2	Single LOH at 2184/2321	—	AF7-16
LOH	3	Single LOH at 1629/1863	—	AF7-3
LOH	3	Single LOH at 2195/2207	—	AF7-7
LOH	3	Single LOH at 1629/1863	—	AF7-17
LOH	4	Single LOH at 2004/2043	—	AF7-12

^a Number of strains analyzed per mouse in parentheses.^b BIR, break-induced replication.

10^{-5} – 1.7×10^{-2} /cell across the 13 mouse populations and rates of LOH at *GALI* at 1.7×10^{-4} ($\pm 2.12 \times 10^{-4}$ SD) events/generation.

For the 20 *in vitro* cultures, we estimated the proportion of cells that had undergone LOH at the *GALI* locus at 3.2×10^{-5} – 1.1×10^{-4} /cell across cultures and the rates of LOH at 6.0×10^{-6} ($\pm 2.0 \times 10^{-6}$ SD) events/generation. Thus, the rate of recombination at *GALI* is ~ 28 -fold greater during *in vivo* growth than during *in vitro* growth (Table 2).

Rates of LOH at individual SNP loci are similar *in vivo* and *in vitro*: For *in vivo*-propagated populations, we determined whole-genome SNP genotypes for 73 strains that were chosen to represent the range of observed 2DG and colony morphology phenotypes. We analyzed SNP genotypes for 20 2DG^S and 53 2DG^R strains, and among these 2DG^R strains, we analyzed SNPs for 24 strains with altered colony phenotypes (Sm and Wr) and 29 strains with PT phenotypes. For the *in vitro* populations, we analyzed 20 2DG^R isolates

(Figure 1). In the course of these genotypic analyses, we found that 13 of the 123 SNP loci were homozygous in the AF7 parent strain; 10 on Chr1, and one each on Chr3, Chr4, and Chr7. These were excluded from further analyses, giving a total of 110 SNP loci analyzed for the 93 strains from both experiments (Table 1). Diagnostic PCR confirmed that all 2DG^R strains lacked both copies of *GALI* and that all 2DG^S strains retained at least one copy of *GALI* (data not shown).

After first determining that the LOH events located in genomes of different isolates obtained from the same mouse represented independent events, we counted 16 short-range LOH events over the 7442 individual SNP loci for which unambiguous data were obtained (Table S3); Of these, 15 altered only a single SNP locus and one longer recombination event spanned three contiguous SNP loci on Chr4 (Table 3). Assuming that each cell represents an average of 7.8–8.3 mitotic cell divisions, the estimated rate is 2.5×10^{-4} short-range LOH events/locus/generation ($\pm 5.2 \times 10^{-4}$ SD). To deter-

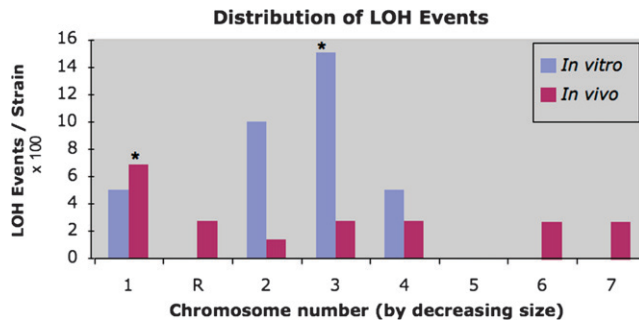


FIGURE 3.—Chromosomal distribution of short-range LOH events involving one to three SNP loci, analyzed separately for *in vivo* and *in vitro* data sets. Shown is the number of rearrangement events per strain for each chromosome ($\times 100$) ordered from largest (Chr1) to smallest (Chr7) chromosome. An asterisk indicates the chromosomes for which greater than expected (average) numbers of LOH events were observed ($P < 0.05$) within the *in vivo* or *in vitro* data sets.

mine the sensitivity of the estimate to the population evaluated, we analyzed results for only the 2DG^R strains where 15 events were observed. The calculated rate is slightly higher at 3.3×10^{-4} events/locus/generation ($\pm 5.8 \times 10^{-4}$ SD). Both estimates are comparable to the rate of LOH at *GALI* during *in vivo* growth that we obtained above (1.7×10^{-4} events/generation).

For the *in vitro*-propagated populations, we detected seven LOH events at single SNP loci and no events composed of multiple SNP loci over the 2012 individual SNP loci for which unambiguous data were obtained. Assuming that each cell represents an average of 9.9 mitotic cell divisions, we obtained a rate of 3.5×10^{-4} short-range LOH events/locus/generation ($\pm 2.4 \times 10^{-5}$ SD) (Table 2). We conclude that the average rates of short-range LOH due to recombination and independent of those occurring at *GALI* are not significantly different for *in vivo* and *in vitro* populations.

Genomewide distribution of short-range LOH events: We next evaluated separately the distribution of short-range LOH events across chromosomes for *in vivo* and *in vitro* propagated populations (Figure 3). Among the *in vivo*-propagated strains, LOH events were distributed across all chromosomes except for Chr5 and Chr5. Comparing the number of LOH events/strain for each chromosome to that expected for each chromosome if LOH events were randomly distributed, the number of events on Chr1 was significantly greater than expected for a random distribution ($P < 0.05$; binomial test or permutation test; see MATERIALS AND METHODS). The larger number of LOH events on Chr1 parallels the higher rate of LOH at *GALI*, also located on Chr1.

For the *in vitro*-propagated populations, short-range LOH events were less evenly distributed across chromosomes than for *in vivo* strains. LOH occurred primarily on the larger chromosomes, Chr1, Chr2, Chr3, and Chr4, although no events were detected on the second largest chromosome (ChrR) or on the smaller chromo-

somes (Chr5, Chr6, and Chr7) (Figure 3). The number of events on Chr3 was significantly greater than expected for a random distribution (binomial test or permutation test; see MATERIALS AND METHODS).

Comparing the results for *in vivo* and *in vitro* populations, it is apparent that the distribution of short-range LOH events differs (Figure 3). For either the *in vitro* or the *in vivo* populations, results show no significant correlation between the number of LOH events and either chromosome size or number of SNP loci analyzed per chromosome (data not shown). For both populations, permutation tests revealed that low numbers of events on some chromosomes could have occurred by chance. These results suggest that the observed differences in the distribution of short-range LOH events across the genome are driven more by the growth environment or the chromosome structure than by the size of the chromosome.

Chromosome-level changes were observed for *in vivo*, but for not *in vitro*, populations: We inferred long-range chromosome-level rearrangement events when AF values for SNP loci and CGH results demonstrated changes spanning all or most of a chromosome (SELMECKI *et al.* 2005). Importantly, we detected six long-range events in isolates from *in vivo* populations (AF617, AF3976, AF3977 for ChrR; AF21, AF540, AF3990 for Chr2) but no long-range events in isolates from *in vitro* populations. The six long-range events that we detected involved two chromosomes, ChrR and Chr2 (Tables 2 and 3).

For ChrR, we recovered two strains with disomic, homozygous ChrR and one strain carrying a trisomic ChrR. Homozygous disomy for ChrR was inferred for strains AF617 and AF3976 because the AF values at most individual SNP loci were close to either 0 or 1, indicating that the entire ChrR was homozygous (Table S3). CGH analysis for these same strains demonstrated that ChrR was present in two copies (Figure 4), suggesting that one ChrR homolog was lost and the second homolog reduplicated. Interestingly, AF values were ~ 0 for strain AF617 and ~ 1 for AF3976 at some SNPs (*e.g.*, locus1381/2345); they were the opposite at other SNPs: ratios were ~ 1 for strain AF617 and ~ 0 for AF3976 (*e.g.*, locus 1694/2254) (Table S3). We infer that strains AF617 and AF3976 are homozygous for different ChrR haplotypes (phases of alleles). We defined ChrR haplotype 1 as the set of SNP alleles found in strain AF3976 and ChrR haplotype 2 as the set of alleles found in strain AF617. Because strains AF617 and AF3976 were obtained from different mice, they must represent independent chromosome loss and reduplication, or nondisjunction, events.

Trisomy was inferred for ChrR of strain AF3977 because most SNP loci demonstrated AF values at either ~ 0.3 or ~ 0.7 , which is intermediate between that expected for heterozygous loci (~ 0.5) and that expected for homozygous loci (~ 0 or 1) (Table S3).

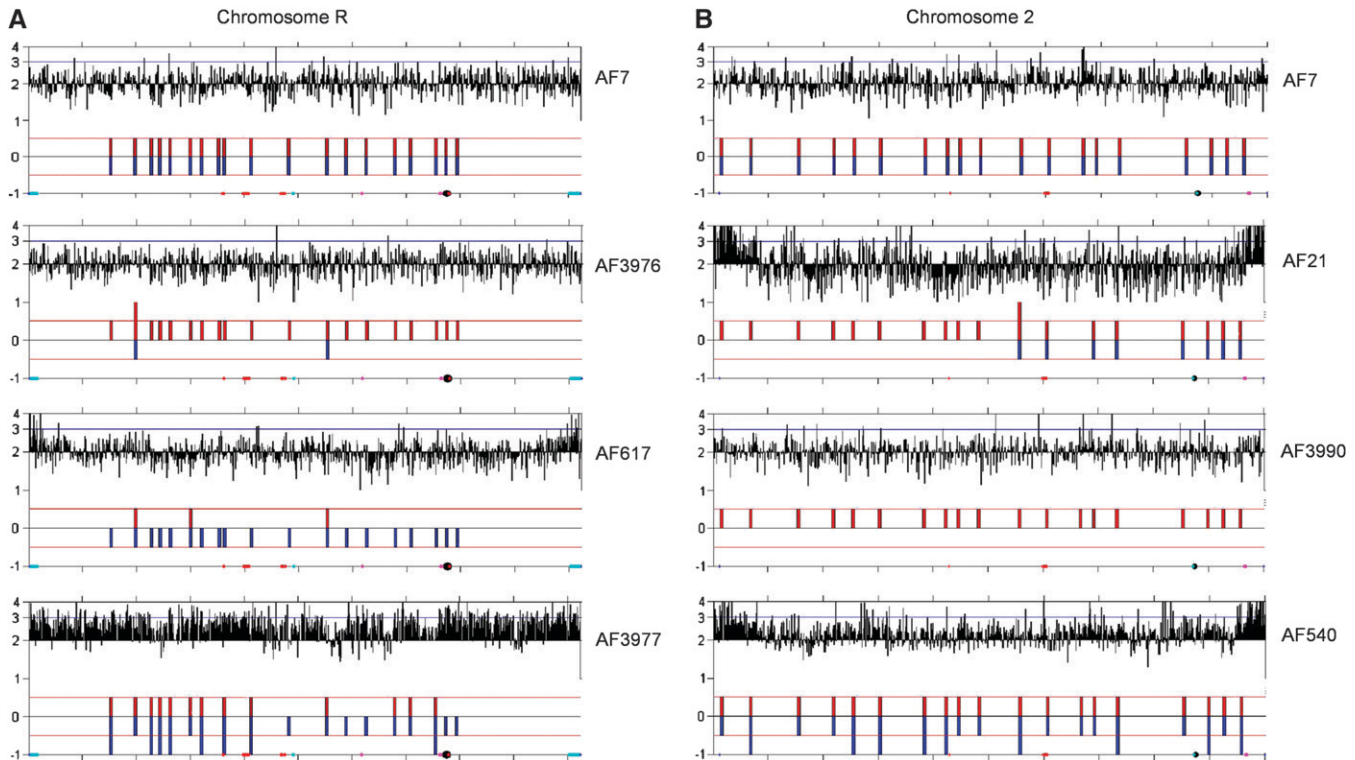


FIGURE 4.—Correspondence of CGH and SNP array results showing chromosome copy number for (A) ChrR and (B) Chr2. Results are shown for parental strain AF7 and those strains demonstrating chromosomal-level rearrangement. Data are plotted as a function of chromosome position using a modified version of Ch_map (SELMECKI *et al.* 2005). Strong signals at telomeres for CGH results are due to incomplete digestion of genomic DNA.

CGH analyses confirmed that ChrR is trisomic (Figure 4A). Following the haplotype designations above, strain AF3977 likely carries two copies of haplotype 2 and one copy of haplotype 1 at ChrR. Nondisjunction could explain the ChrR complement for strains disomic AF3976 and trisomic AF3977 because these strains were isolated from the same mouse.

For Chr2, we recovered three different strains demonstrating chromosome-level events (Figure 4B, Table 3). Because these were recovered from populations in three different mice, we assume that they represent independent events. In strain AF3990, the AF values for most SNP loci were close to either 0 or 1, revealing that Chr2 was homozygous. CGH analysis determined that Chr2 was disomic. In strain AF540, AF values were intermediate between expected homozygous and heterozygous values, suggesting that markers on Chr2 were trisomic, a result confirmed by CGH analysis. In the third strain, AF21, only SNP loci on the left arm of Chr2 were homozygous, and CGH showed that Chr2 was disomic, leading to the conclusion that break-induced replication (BIR) or a single crossover (KRAUS *et al.* 2001) generated the left arm of Chr2 (Figure 5). We designated haplotype 1 as the set of alleles found homozygous on Chr2 in strain AF3990. This same set of alleles was homozygous for most of the left arm of Chr2 in strain AF21. Haplotype 2 was defined as the set

of alleles with AF values of ~ 0.7 on trisomic Chr2 in strain AF540 (Table S3).

In summary, all six chromosome-level recombination events were recovered from *in vivo* propagated strains, and these involved only two chromosomes, ChrR and Chr2 (Tables 2 and 3). Among these six long-range LOH events, we found two cases of whole-chromosome trisomy (one for ChrR and one for Chr2), three cases of chromosome homolog loss with duplication of the remaining homolog by reduplication or nondisjunction, and one case in which most of one chromosome arm likely underwent BIR or a single crossover (KRAUS *et al.* 2001) (Tables 2 and 3). The *in vivo* rate of long-range chromosome-level events was estimated at 1.2×10^{-3} events/chromosome/cell division ($\pm 4.3 \times 10^{-3}$ SD) among the 584 chromosomes evaluated, an order of magnitude higher than the rate of short-range LOH events in this same population. Chromosome-level events were not detected among the 160 chromosomes evaluated from *in vitro*-propagated isolates (Tables 2 and 3).

LOH at *GALI* is associated with recovery of additional rearrangement events: We asked whether LOH at *GALI*, detected by 2DG selection, was associated with recovery of genome rearrangement events occurring independently of those at the *GALI* locus. Among the *in vivo*-propagated strains, we performed SNP anal-

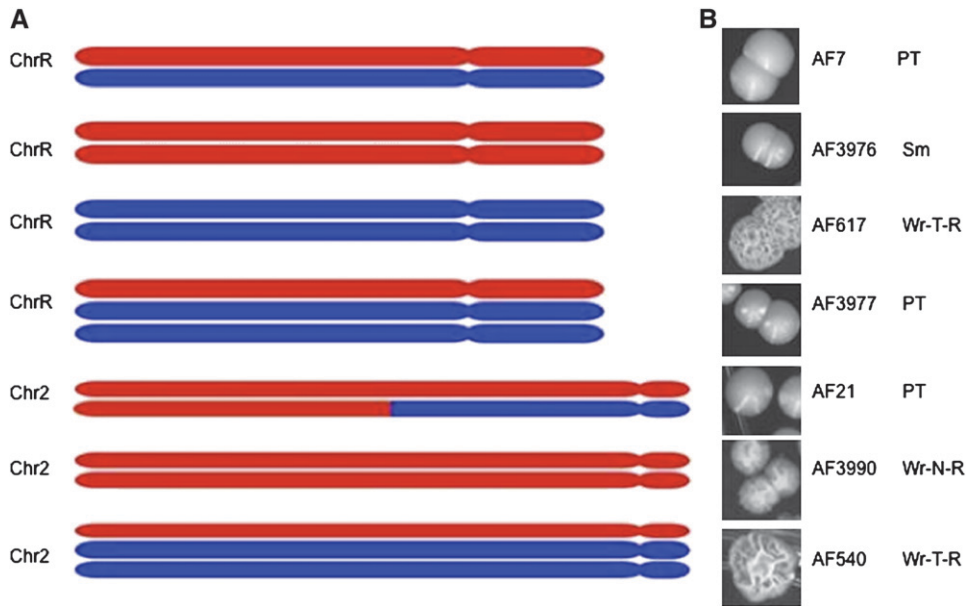


FIGURE 5.—Chromosome rearrangement is associated with altered colony phenotypes. (A) Chromosome diagrams for ChrR and Chr2. Haplotype 1 is indicated in red and haplotype 2 is indicated in blue. (B) Colony morphology, strain identification, and colony growth phenotype associated with chromosomal genotypes.

ysis on 53 2DG^R and 20 2DG^S strains. Among the 2DG^R isolates, we found 14 short-range LOH events covering a single SNP locus, one event covering three SNPs, and six chromosome-level events described above. In the 20 2DG^S strains, we detected only a single SNP LOH event, on ChrR, in strain AF656 (Table 3). From the *in vitro*-propagated populations, 20 2DG^R strains demonstrated seven different LOH events at single SNP loci and no chromosome-level events.

An association test showed that genome rearrangement events were significantly associated with 2DG^R strains ($G_{stat} = 6.23$, d.f. = 1, $P = 0.01$). From the data available, we cannot determine if events at *GALI* and other SNP loci or chromosomes occurred in the same cell cycle or at different times during cell division in the host. Nonetheless, the 2DG selection for strains exhibiting LOH at *GALI* yields strains that have undergone additional LOH events at other loci and chromosomes.

Altered colony phenotypes arose during *in vivo* but not *in vitro* propagation: We next determined the rate at which altered colony growth phenotypes arose and the relationship between these phenotypes and genome-wide genetic changes. For the *in vivo* populations, when grown at 32°, 65 of the 474 2DG^R isolates (13.7%) exhibited colony growth phenotypes different from that of parent strain AF7 (“PT” in Figure 5). Of these 65 strains with altered colony phenotype, 29 formed smaller colonies (Sm) than did AF7, and 36 exhibited filamentous Wr colony morphologies with varying degrees of wrinkling relative to strain AF7. The Sm phenotype was due to slower cell growth, as determined by growth-rate measurements in liquid culture for two of these Sm strains (data not shown). Of the 36 Wr strains, 32 formed smooth colonies at 23°, but formed filamentous, wrinkled colonies at 32° and 37°, the Wr-T-R phenotypes. Four of the 36 Wr strains formed wrinkled

colonies at all three temperatures, the Wr-N-R phenotypes (Figure 5, Table 4).

In striking contrast to the *in vivo* populations, we did not observe altered colony phenotypes among >7000 colonies observed during the *in vitro* experiment. Although filamentous colony phenotypes are detected occasionally during routine maintenance of SC5314 and its derivative strains, these events are rarely observed with growth at 32°. We discount the explanation that 2DG media is mutagenic and leads to increased rates of LOH and phenotypic variation because the *in vitro*-propagated cells that were subjected to 2DG media did not show elevated rates of LOH or phenotypic variation. Thus, the higher rate of altered colony phenotypes observed in strains from *in vivo* propagation and selected on 2DG is attributable to different growth

TABLE 4
Association of genotypic and phenotypic change

Genome change	Colony growth phenotype ^a				
	Parental		Nonparental		
	PT	Sm	Wr-N-R	Wr-T-R	Total ^b
No change	42	0	0	10	52
Chromosome and BIR ^c	2	1	1	2	6
LOH	5	0	4	7	16
Total	49	1	5	19	74

Goodness-of-fit statistic: 26.45, d.f. = 10, $P < 0.005$.

^a PT, parental colony phenotype; Sm, small colony phenotype; Wr-N-R, not regulated by temperature; Wr-T-R, colony phenotype dependent on temperature.

^b One strain exhibited two LOH events, giving a total of 74 events in 73 strains.

^c BIR, break-induced replication.

conditions *in vivo* compared to those *in vitro*, and not to the 2DG selection itself.

Genome rearrangement is associated with altered colony phenotypes: We examined the relationship of altered colony phenotypes and genome rearrangement within the data set for *in vivo* populations. Using the goodness-of-fit test, there is a significant association between altered colony phenotypes (Sm, Wr) and 2DG^R phenotypes relative to 2DG^S phenotypes ($G_{\text{stat}} = 19.45$, d.f. = 3, $P = 0.01$).

We then evaluated the relationship of altered colony phenotypes (Sm, Wr) and those short-range and chromosome-level LOH events at loci other than *GALI*. We found that altered phenotypes were significantly and positively associated with LOH events while the PT smooth colony phenotype was more often associated with the absence of detected LOH events than would be expected by chance (Table 4). Examining the specific events and strains, we found that four of the five strains in which LOH occurred along an entire chromosome also exhibited an altered colony phenotype. For ChrR, homozygous haplotype 1 (strain AF3976) was associated with the Sm phenotype while homozygous haplotype 2 (strain AF617) was associated with the Wr-T-R phenotype (Figure 5). Interestingly, trisomy of ChrR with an extra copy of haplotype 2 (strain AF3977) is not associated with an altered colony phenotype. These results suggest that the loss of specific ChrR alleles leads to altered colony phenotypes, while a change in the ratios of alleles from 1:1 to 2:1 on ChrR does not have the same effect.

We also detected changes in colony phenotype associated with chromosome-level changes on Chr2. Homozygous haplotype 1 is associated with the Wr-N-R colony phenotype (strain AF3990), and trisomy of Chr2 with two copies of haplotype 2 is associated with the Wr-T-R colony phenotype (strain AF540) (Figure 5). In contrast to ChrR results, we found that both the loss of a haplotype or an imbalance of allele ratios on Chr2 gave rise to altered colony phenotypes. Finally, conversion of the left half of Chr2 to homozygous haplotype 1 was not associated with a change in colony phenotype (strain AF21), which suggests that the Wr-T-R phenotype in strain AF3990 is due to homozygous haplotype 1 loci on the right half of Chr2. Together, results show varying effects of trisomy and hemizyosity at ChrR and Chr2 on growth phenotype.

Synthesizing results for LOH at *GALI*, rearrangement events independent of *GALI*, and altered colony phenotypes, we find that altered colony phenotypes are observed more frequently in strains that have undergone at least one LOH event at *GALI* but that the underlying cause of altered colony phenotypes is additional, *GALI*-independent LOH events. A large proportion of isolates with altered colony phenotypes also had undergone chromosomal changes via short-range LOH events (11/17, or 65%) or chromosome-

level events (4/6, or 80%) (Table 4). Overall, our results show that the choice of isolates with altered colony phenotypes increased the detection of genome changes and are consistent with findings that LOH at many different loci influences filamentous growth in *C. albicans* (UHL *et al.* 2003).

DISCUSSION

Opportunistic pathogens often live as commensal associates of plants and animals, becoming dangerous infectious agents when host defenses are weakened or new virulence types arise by mutation (MARGOLIS and LEVIN 2007). Since many of these microbial pathogens propagate asexually, understanding the mechanisms by which opportunistic pathogen populations respond to host environments is essential to developing effective disease control and antifungal therapy and to understanding the fundamental evolutionary question of how adaptation occurs in asexual lineages. Here, we report differing types and distribution of recombination events across the genome for *in vivo*- vs. *in vitro*-propagated populations and that altered colony growth phenotypes arose only within the *in vivo* populations. Our results demonstrate that *C. albicans* populations generate considerably more phenotypic and genetic variation during growth in a living host than in the relatively benign environment of *in vitro* culture.

Net population growth rates are 5- to 10-fold lower *in vivo* than *in vitro*. Immediately after *C. albicans* is introduced into the mouse host, population sizes dramatically decline in the bloodstream and kidneys and subsequently expand exponentially in the kidney, causing renal failure and death (ODDS *et al.* 2000; MACCALLUM and ODDS 2004). Because log growth is density independent, the results suggest that the slower growth trajectory characteristic of *in vivo* populations is less likely a result of direct competition among *C. albicans* cells and more likely a result of higher cell mortality due to the host immune response or slower cell division due to nutrient limitations (BARELLE *et al.* 2006; BERNARDIS *et al.* 2007). The linear growth rate of pseudohyphal and hyphal cells (HAUSAUER *et al.* 2005) may also contribute to slower population growth in the *in vivo* environment. Slower growth *in vivo* is not surprising; the result is important because relative rates of genetic change can be evaluated and because of the empirical insight provided into the population dynamics of an important pathogen.

We found that average rates of LOH at individual SNP loci across the genome are quite similar for *in vivo* and *in vitro* populations ($\sim 10^{-4}$ events/generation). Yet the distribution of events differed with greater numbers of short-range LOH events on Chr1 for *in vivo* populations and on Chr3 for *in vitro*-propagated populations than expected for a random distribution across all SNP loci

and chromosomes. Higher rates of LOH at the *GALI* locus on Chr1 were observed during *in vivo* growth than during *in vitro* growth. Loss of the *URA3* gene with LOH at the engineered *GALI* locus could lead to lower virulence (STAAB and SUNDSTROM 2003), but such an effect should decrease, rather than increase, the recovery of Gal⁻ strains. Previous results show little evidence that *URA3* copy number affected survival in our experimental system (FORCHE *et al.* 2003). The results show that the growth environment strongly affects the genomic distribution of short-range LOH events and that, in either environment, LOH events are unevenly distributed across the genome.

Both the rates and the distribution of chromosome-level events across the genome differed from the rates and distribution of short-range LOH events. Most striking, chromosome-level events were observed only for *in vivo* populations, and in those populations, only for ChrR and Chr2. Our results are supported by those of DIAGO *et al.* (2009) for a limited number of *C. albicans* strains isolated from the digestive tract of healthy individuals where chromosome-level events (trisomy, nondisjunction) were prevalent. We conclude that conditions during growth in a mammalian host affect chromosome nondisjunction more strongly than they affect mitotic crossover or other recombination processes. While the type of rearrangements differs from the mutational spectrum observed in bacterial pathogens during growth in stressful conditions (VICTOIR and DUJARDIN 2002; TENAILLON *et al.* 2004; PONDER *et al.* 2005), common mechanisms involving recombination and repair may yet be identified.

The rates of chromosome-level recombination that we observed ($\sim 10^{-3}$ events/generation) are comparable to those reported for rates in other fungi at $\sim 10^{-3}$ events/kb/generation (AWADALLA 2003). Although high rates of chromosomal rearrangement have been observed at specific chromosomal loci of other pathogens (HENDERSON *et al.* 1999; VICTOIR and DUJARDIN 2002; KLINE *et al.* 2003; UHL *et al.* 2003; ZHANG *et al.* 2003), few quantitative estimates of genomewide change are available to explain the variation apparent in clinical populations (GRAESER *et al.* 1996; FRIES and CASADEVALL 1998; IWAGUCHI *et al.* 2000; FORCHE *et al.* 2005). The rates determined in our study are certainly sufficient to generate heritable variation in *C. albicans* populations during a single passage through the host.

The variation in colony phenotypes generated during *in vivo* growth is of most direct consequence to *C. albicans* fitness in the host. We estimated the recovery of altered colony phenotypes at $\sim 10^{-6}$ /generation with a large fraction of these also regulated by temperature (Wr-T-R) and the greatest trait expression at 37°. This finding is consistent with genetic studies that found that many genes, when hemizygous, affect filamentous growth *in vitro* (UHL *et al.* 2003) and suggests that the number of genes that affect the quality of filamentous

growth is larger than the number of genes that affect the temperature regulation of filamentous growth. We found a positive association between altered colony phenotypes and short-range and chromosome-level LOH events and infer that choosing phenotypic variants for analysis increased the overall rate of the recovery of rearrangement events. Within *in vitro*-propagated strains, we did not observe altered colony growth phenotypes, which is consistent with the very low recovery of chromosomal rearrangement or altered colony growth phenotypes observed for *in vitro* growth (reviewed in RUSTCHENKO 2007). Our observation of the appreciable phenotypic variation arising during a single passage through a mouse host is striking and is of sufficient magnitude to strongly affect the evolution of *C. albicans* populations within the short lifetime of the affected host.

Recombination events as well as the altered colony phenotypes were positively associated with the recovery of 2DG^R strains. We considered the possibility that selective growth on 2DG, like growth on 5-fluoroorotic acid (BOEKE *et al.* 1987), might be mutagenic. However, no altered colony phenotypes were recovered among the many 2DG^R strains derived from *in vitro* experiments, and controls show that cells were subject to 2DG selection for too little time to generate and recover colonies arising from *GALI* mutation. Further, rates at *GALI* (10^{-6} /generation) *in vitro* are comparable to those reported for *Saccharomyces cerevisiae* at the *URA3* locus at $\sim 5 \times 10^{-6}$ events/generation (calculated from HIRAOKA *et al.* 2000). The *GALI* locus resides on Chr1, suggesting that the higher rate of LOH at *GALI* for *in vivo* populations results from a greater rate of recovering short-range LOH events there. We conclude that 2DG selection is not mutagenic; rather, it provides a portal through which genome rearrangement events and associated phenotypic variation can be efficiently recovered and studied.

Together, our results show that relatively few mitotic generations occurring during a single passage through a mouse host generate measurable genomewide and phenotypic variation within populations. Our results shed light on the role of recombination in the evolution of asexual organisms (DUNHAM *et al.* 2002), especially opportunistic pathogens. The evolution of genetic factors that increase recombination rates will be favored under environmental uncertainty (LENORMAND and OTTO 2000; OTTO 2002), and previous studies with *S. cerevisiae* demonstrate an advantage to recombining populations during infectious growth (GRIMBERG and ZEYL 2005). We propose that the commensal state of opportunistic pathogens such as *C. albicans* is maintained by the dynamic fluctuations of populations (LEVIN *et al.* 2000; LEVIN and ANITA 2001). In this model, population bottlenecks decrease the efficiency of selection in removing deleterious phenotypes that arise during infectious growth. Even though population

sizes may be quite large at later infection stages, populations will be dominated by those mutations arising early in passage through the host. In contrast, traits under selection during commensal growth will likely differ from those beneficial to invasive growth, and larger population sizes allow efficient selection. Either the retention of deleterious alleles by drift when population sizes are small, or clonal interference among genotypes carrying beneficial mutations (KAO and SHERLOCK 2008; CAMPOS and WAHL 2009) when populations are large, will slow evolution of virulence and host adaptation. If so, further experiments should find that colony growth and virulence phenotypes generated during host passage are more variable than expected under strong directional selection for increased virulence and instead constitute “sink” populations that are unable to compete in the commensal environment or survive past host death (ROMANI *et al.* 2003; SOKURENKO *et al.* 2006). As in drug resistance evolution (ANDERSON 2005), it will be important to evaluate fitness associated with differing growth conditions and varying rates of genome rearrangement. Results presented here portray the evolution of a genome strongly responsive to growth environment as well as suggest an important role of recurrent bottlenecks and population expansion during host passage and reinfection common to clinical settings.

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Supporting Information

<http://www.genetics.org/cgi/content/full/genetics.109.103325/DC1>

**Evolution in *Candida albicans* Populations During a Single
Passage Through a Mouse Host**

Anja Forche, P. T. Magee, Anna Selmecki, Judith Berman and Georgiana May

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TABLES S1 – S3

Tables S1-S3 are available as Excel files at <http://www.genetics.org/cgi/content/full/genetics.109.103325/DC1>.

Table S1: Primer sequences for SNP loci, Allele-Specific Oligonucleotides (ASO), *GALI* diagnostic primers

Table S2: Rates of LOH at the *GALI* locus for *in vivo* and *in vitro* experiments

Table S3: Analysis of SNP loci for *in vivo* and *in vitro* propagated strains