Higher-fitness yeast genotypes are less robust to deleterious mutations

Milo S. Johnson¹², Alena Martsul⁴, Sergey Kryazhimskiy⁴⁻⁵, Michael M. Desai¹²,³,⁵⁻-six

Natural selection drives populations toward higher fitness, but second-order selection for adaptability and mutational robustness can also influence evolution. In many microbial systems, diminishing-returns epistasis contributes to a tendency for more-fit genotypes to be less adaptable, but no analogous patterns for robustness are known. To understand how robustness varies across genotypes, we measure the fitness effects of hundreds of individual insertion mutations in a panel of yeast strains. We find that more-fit strains are less robust: They have distributions of fitness effects with lower mean and higher variance. These differences arise because many mutations have more strongly deleterious effects in faster-growing strains. This negative correlation between fitness and robustness implies that second-order selection for robustness will tend to conflict with first-order selection for fitness.

The dynamics and outcomes of adaptive evolution depend on the genetic variation available to a population. Because mutations interact epistatically, the availability and strength of beneficial and deleterious mutations can vary across genotypes (1). As “first-order” natural selection drives populations toward higher fitness, “second-order” selection can favor genotypes with better prospects for future evolution, steering populations into regions of the fitness landscape with more robust population 

Many laboratory microbial populations display a consistent pattern of declining adaptability, such that less-fit genotypes adapt more rapidly than more-fit genotypes (9–13). This is at least partially explained by “diminishing-returns” epistasis, in which individual beneficial mutations become less beneficial in more-fit genetic backgrounds (12–15). In contrast to adaptability, genetic variation in mutational robustness has not been systematically characterized, and no analogs to diminishing epistasis (16–19), but little is known about how the entire distribution of fitness effects (DFE) of deleterious mutations changes across different genetic backgrounds and whether robustness, like adaptability, depends systematically on fitness.

There are several ways to define and measure mutational robustness (2–4). In this work, we consider only the single-step mutational neighborhood of a genotype. We define robustness on the basis of the DFE of single mutations and refer to strains in which these mutations are more deleterious on average as less robust (though the appropriate precise definition of mutational robustness can depend on the full DFE and the specific population genetic question). To understand genetic variation in single-step mutational robustness, we aim to measure how the DFE varies across genotypes and how these differences arise from epistasis at the level of individual mutations. To do so, we would ideally like to measure the fitness effects of identical large sets of random mutations in multiple genotypes. This would allow us to compare the DFE of these mutations across genotypes and to identify the genetic basis of differences in robustness.

To this end, we developed a pipeline to measure the effects of sets of specific insertion mutations in a panel of Saccharomyces cerevisiae genotypes (Fig. 1). Briefly, we transform yeast strains with transposon mutagenesis libraries derived from (20), in which each plasmid is tagged by multiple specific DNA barcodes. Homology-directed repair then creates the same set of transposon insertion mutations

Fig. 1. Schematic of mutagenesis and fitness-assay pipeline. Plasmids with different colors indicate different regions of homology from the yeast genome. BC, barcode; Seg., segregant; Tn7, transposon 7.
in each strain. We propagate the resulting mutant pools in batch culture and measure barcode-frequency trajectories to estimate the fitness effect of each mutation in each strain (2F) (figs. S1 and S2). Using this approach, we conducted two experiments to measure mutational robustness in F1 segregants derived from a yeast cross between a laboratory and a wine strain (BY and RM). These segregants differ at more than 35,000 loci and have previously been sequenced and phenotyped (22). They vary in fitness (across a 22.5% range) in our focal environment, and those with lower fitness are more adaptable (9).

In the first, “large library” experiment, we transformed 18 randomly chosen segregants from (9) with libraries consisting of 1147 mutations (fig. S3) (2F). Although these are not a random sample of all naturally occurring mutations in yeast, they represent an unbiased set of genomic disruptions. Because of gene essentiality, differences in cloning or transformation efficiency, or other complications, we were unable to measure the fitness effect of every mutation in every segregant (analysis of the effects of missing data is in (2F) and fig. S4). We successfully measured the effects of 710 mutations in at least one segregant (on average, 441 per segregant). Of these 710, 457 (64%) had no detectable fitness effects in any segregant. Most remaining mutations were deleterious, and their DFE varied systematically across strains. Specifically, the mean fitness effect of these mutations decreases with the background fitness of the segregant (i.e., more-fit segregants tend to be less robust with respect to random insertion mutations, $P = 0.03$, two-sided $t$ test) (Fig. 2, A and B, and fig. S5).

Because we only measured fitness effects across 18 segregants in our large library experiment, and 64% of the mutations were indistinguishable from neutral, we were unable to detect more-subtle changes in the DFE or to connect DFE-level changes to patterns of epistasis for individual mutations. To address this limitation, in our second experiment we transformed a larger group of 163 randomly chosen segregants with a “small” library, created by selecting a subset of 91 insertions from our large library that had significant fitness effects in the largest number of segregants (2F). This small library filters out insertions with undetectable or rare effects but is otherwise unbiased.

Again, we find that the mean of the DFE decreases as segregant fitness increases (Fig. 2, C to E). We also find significant correlations between background fitness and the variance and skew of the DFE: more-fit segregants have wider DFEs that are skewed toward more-deleterious mutations (Fig. 2, C, F, and G, and fig. S6). These results imply that mutational robustness is negatively correlated with background fitness. This suggests that second-order selection for robustness could be constrained by conflict with first-order selection for fitness.

The DFE is composed of individual mutations. To understand why the shape of the DFE varies between segregants, we examined how the effects of these individual mutations vary. We observe a variety of types of epistasis (Fig. 3 and fig. S7), including nearly constant effects across backgrounds (e.g., PAH1, MME1) and diminishing returns (e.g., SIR3). However, the most frequent pattern (48 cases) is “increasing-cost” epistasis such that the mutation is more deleterious in more-fit segregants (Figs. 3 and 4A). This is the deleterious-mutation analog of diminishing-returns epistasis. Notably, this effect can cross zero: Some mutations are beneficial in the least-fit segregants, neutral in intermediate-fitness segregants, and deleterious in higher-fitness segregants (e.g., RPL16A). This negative correlation is not universal; six mutations exhibit the opposite pattern (e.g., KRI1).

In addition to background fitness, specific genetic loci can influence the fitness effects of individual mutations (e.g., NOP16) (Fig. 3). We used the same procedure as in (9, 22) to identify such quantitative trait loci (QTLs) for each mutation (fig. S8). To quantify how these QTLs and background fitness explain the variation in the fitness effect of each mutation, we fit three linear models to our data (2F). The “fitness” model includes segregant fitness as the only predictor. The “QTL” model includes only
The segregant genotype at a small number of QTLs. The “full model” includes both segregant fitness and QTL effects.

We find that at least one of these three models is significantly better than the null model without epistasis for 71 of the 80 mutations for which we have measurements in at least 50 segregants (Fig. 4B). For mutations in which the fitness model has explanatory power (64 cases, 80.0%), most are more deleterious in more-fit backgrounds (58 cases), and most of these are deleterious on average (i.e., exhibit increasing-cost epistasis, 48 cases). QTLs have explanatory power in 60 cases (75.0%), but because many QTLs also affect segregant fitness, QTL and fitness contributions are confounded. To understand which of our models best explains the fitness effects of each mutation while involving as few parameters as possible, we compare the Akaike information criterion (AIC) of each model. The AIC is lowest for the background fitness model for 10 mutations (12.5%), the QTL model for 16 mutations (20%), and the full model for 45 mutations (56.25%). However, even when the background fitness model does not have the lowest AIC, in many cases it explains close to as much variation as the QTL or full model.

One potential explanation for increasing-cost epistasis is that mutations have effects during the saturation phase of our batch culture propagation, and more-fit strains are less robust because they spend longer in saturation. We measured fitness within a single growth cycle and found that this is not the case. Instead, differences in fitness arise almost exclusively during exponential growth (21) (figs. S9 to S11).

To understand why faster-growing strains are less robust, we looked for functional similarities among genes disrupted by mutations with similar epistatic patterns [by mutations whose fitness effects were modified by the same QTL (fig. S12) or by mutations with strong fitness-mediated epistasis (21)]. We found enrichment in several Gene Ontology (GO) terms among such genes at P < 0.05 (though none remain significant after multiple-hypothesis correction) (table S1) (23). Most notably, the fitness-mediated epistasis set and the set associated with the most commonly observed QTL are enriched for ribosome- and translation-related functions. This QTL is also the strongest background fitness QTL and includes variants in KRE33, a gene involved in small ribosomal subunit assembly (9). Metabolic control theory provides a possible link between this functional information and increasing-cost epistasis. Specifically, it predicts that a deleterious mutation in one enzyme will have a weaker effect if other enzymes in any sequential biological pathway are already defective, given that fitness is correlated with metabolic flux through that pathway (18, 24).
This also applies to sequential pathways in transcription and translation (25), so increasing costs epistasis could arise because more-fit segregants have a better-optimized ribosome synthesis or protein synthesis pathway and therefore experience greater costs when deleterious mutations affect these pathways. Although these results are not definitive, they suggest that further generalizations could be drawn from a deeper understanding of the cell-physiological basis of fitness-mediated epistasis.

Our results are limited to the analysis of individual gene disruption mutations in a specific set of yeast strains, and it is possible that other types of mutations (e.g., regulatory changes or second-step mutations) exhibit different patterns. However, to the extent that the patterns of fitness-dependent epistasis observed in this research and in previous work on beneficial mutations (9–15) hold more broadly and over multiple mutational steps, their net effect is a predictable change in the local properties of the fitness landscape as populations adapt. This local mutational neighborhood becomes less favorable in more-fit genotypes: Uphill steps become flatter or even change to downhill, and many downhill paths become steeper. On such landscapes, first-order selection for high-fit genotypes conflicts with second-order selection, and populations evolve toward more-fit but less robust and less adaptable genotypes. Thus, even if true fitness peaks exist, populations may never attain them, instead reaching a dynamic balance between beneficial and deleterious mutations (26).

REFERENCES AND NOTES
21. Materials and methods and supplementary text are available as supplementary materials.

ACKNOWLEDGMENTS
We thank E. Jerison, L. Rast, A. Nguyen-Ba, J. Yodh, G. Wildenberg, and members of the Desai lab for experimental assistance and/or comments on the manuscript. Funding: This work was supported by an NSF Graduate Research Fellowship (to M.S.J.), a BWF Career Award at the Scientific Interface (grant 1010791.01), the Alfred P. Sloan Foundation (grant FG-2011-2017), the Hellman Foundation, the Simons Foundation (grant 375796), the NSF (DBI-1655960), and the NIH (GM104239). Computational work was performed on the Odyssey cluster supported by the Research Computing Group at Harvard University. Author contributions: Conceptualization: S.K. and M.M.D.; experimental design: M.S.J., A.M., S.K., and M.M.D.; methods development: M.S.J. and A.M.; experiments: M.S.J.; analysis: M.S.J. and S.K.; writing: M.S.J., S.K., and M.M.D. Competing interests: None declared. Data and materials availability: Data described in the paper are presented in the supplementary materials. Raw sequencing data are publicly available at the NCBI Sequence Read Archive (accession no. SRP216610), and all analysis code is available from Zenodo (27).

SUPPLEMENTARY MATERIALS
science.sciencemag.org/content/366/6464/490/suppl/DC1

Materials and Methods
Supplementary Text
Figs. S1 to S12
Tables S1 to S6
References (28–40)
Data File Descriptions
Data Files S1 to S3 (Excel format)

18. June 2019; accepted 11 September 2019
10.1126/science.aay4199
Higher-fitness yeast genotypes are less robust to deleterious mutations
Milo S. Johnson, Alena Martsul, Sergey Kryazhimskiy and Michael M. Desai

Science 366 (6464), 490-493.
DOI: 10.1126/science.aay4199

Genetic background affects variation
Robustness, or the effect of mutations on fitness, can affect the evolutionary trajectory of a species. By introducing a large number of deleterious mutations into many different genetic backgrounds of yeast, Johnson et al. found that, for many mutations, the more fit the background, the larger the deleterious effect of the mutation (see the Perspective by Miller). A more-fit lineage is thus less tolerant to deleterious mutations, whereas less-fit lineages can tolerate more mutations. This observation supports a tendency toward diminishing returns for beneficial mutations, which has been shown to influence patterns of adaptation.
Science, this issue p. 490; see also p. 418