

# Genotype to Phenotype: A Complex Problem

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Rapid genome-sequencing methods coupled with whole-genome transcription profiling suggests that it may be possible to predict phenotype from a genotype. Human genetic association studies of common single-nucleotide polymorphisms (SNPs) explain only a fraction of phenotypic variation among individuals (1). This may be due to rare SNPs (1), structural (2) and epigenetic variants, or multiple alleles with additive effects or synergistic genetic interactions associated with complex combinations of genetic variation (3).

To address the genotype-to-phenotype problem, we developed a simple comparative model for the budding yeast *Saccharomyces cerevisiae* that enables a comprehensive assessment of the genetic mechanisms leading to different phenotypes for the same mutation in two different genetic backgrounds. The strain  $\Sigma$ 1278b mates and forms viable meiotic progeny with the reference strain, S288c, and the divergence between the two strains is roughly equal to the divergence between the genomes of two humans (4, 5).

We sequenced and assembled the 12-Mb  $\Sigma$ 1278b genome, annotating 6923 open reading frames (ORFs) and RNAs, of which 6848 have orthologs within S288c (5). The order of genes between the strains was the same (except in the highly variable subtelomeric regions), and the sequence of 46% of the  $\Sigma$ 1278b ORFs was identical to those in S288c. Differences between the strains were largely due to small insertions and deletions or SNPs, with an average SNP density of 3.2 per kilobase.

We deleted ~5100 genes within  $\Sigma$ 1278b (5) to systematically compare identical deletion

mutants (6). In particular, we identified “conditional essentials,” those genes required uniquely for viability in either strain (fig. S3 and table S1). We scored colonies as dead or alive and surveyed all vital pathways for individual-specific genetic interactions. We expected such conditional essential genes to be rare because the genomes of  $\Sigma$ 1278b and S288c are nearly identical.

Although 894 genes were essential in both S288c and  $\Sigma$ 1278b, 44 genes were essential only in  $\Sigma$ 1278b and 13 genes were essential only in S288c (Fig. 1A). The conditional subsets included genes of various functions; however, the  $\Sigma$ 1278b subset was enriched for genes involved in mRNA metabolic process, whereas the S288c set was enriched for genes annotated to SRP-dependent co-translational targeting. These biological biases suggest that these phenotypes result from genetic interactions associated with an individual genotype.

Hybrid strain crosses and tetrad analysis focusing on 18 mutants that were lethal in  $\Sigma$ 1278b with wild-type levels of fitness when deleted in S288c were used to investigate conditional essentiality. We mated viable haploid S288c deletion mutants to wild-type  $\Sigma$ 1278b and analyzed the hybrid diploid progeny by tetrad analysis. The number of viable meiotic progeny carrying the deletion allele is related to the number of unlinked background-specific modifiers (5) that contribute to the genetic interaction. In all 18 cases, the conditional phenotype was associated with numerous modifier genes that differ between strains. The simplest cases, *SKI7* and *BEM1*, are likely due to a genetic interaction with at least two or more modifiers, but all other cases were more complex (Fig. 1B). Thus,

our analysis showed that conditional essentiality is almost always a consequence of complex genetic interactions involving multiple modifiers associated with strain-specific genetic variation rather than classic digenic synthetic lethality (6, 7).

Our genome-wide survey of conditionally essential genes demonstrates that in most cases a complex set of background-specific modifiers influence a mutation whose phenotype differs between individuals. These results raise the possibility that similar complex modifiers may largely explain the difficulty in identifying the genetic basis for individual phenotypes. The potential for genetic interactions to control individual phenotypes becomes even more important if different combinations of alleles can lead to the same physiological state. The ability to identify these conditional essential phenotypes in yeast provides a framework to unravel the fundamental principles of genetic networks resulting from natural variation, including those that underlie human disease.

## References and Notes

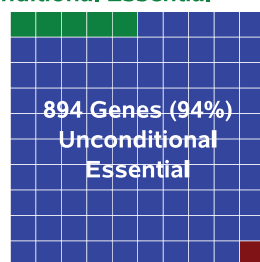
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## Supporting Online Material

www.sciencemag.org/cgi/content/full/328/5977/469/DC1  
Materials and Methods  
Figs. S1 and S2  
Tables S1 to S3  
References

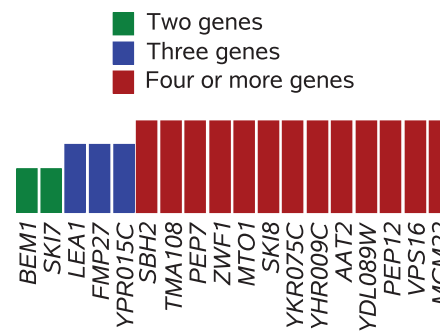
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## A $\Sigma$ 1278b: 44 Genes (5%) Conditional Essential



S288c: 13 Genes (1%)  
Conditional Essential

## B Number of Modifiers



**Fig. 1. (A)** Most S288c essential genes are also essential in  $\Sigma$ 1278b (94%); however, ~5% are essential only in the  $\Sigma$ 1278b genetic background, whereas ~1% are essential only in S288c. **(B)** Conditional essential genes in  $\Sigma$ 1278b are the consequence of complex genetics.  $\chi^2$  tests indicated the number of modifiers associated with conditional essentiality (5).

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