RESEARCH ARTICLE SUMMARY

YEAST GENETICS

Exploring genetic suppression interactions on a global scale

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INTRODUCTION: Genetic suppression occurs when the phenotypic defects caused by a mutated gene are rescued by a mutation in another gene. These genetic interactions can connect genes that work within the same pathway or biological process, providing new mechanistic insights into cellular function, or they can correct defects in gene expression or protein production. More generally, suppression interactions may play an important role in the genetics underlying human diseases, such as the diverse penetrance of Mendelian disease variants. Our ability to interpret personal genome sequences remains limited, in part, because we lack an understanding of how sequence variants interact in nonadditive ways to generate profound phenotypes, including genetic suppression.

RATIONALE: Genetic interactions, in which mutations in two different genes combine to generate an unexpected phenotype, may underlie a significant component of trait heritability. Although genetic interactions that compromise fitness, such as synthetic lethality, have been mapped extensively, suppression interactions have not been explored systematically. To understand the general principles of genetic suppression and to examine the extent to which these interactions reflect cellular function, we harnessed the powerful genetics of the budding yeast Saccharomyces cerevisiae to assemble a global network of genetic suppression interactions.

RESULTS: By analyzing hundreds of published papers, we assembled a network of genetic suppression interactions involving ~1300 different yeast genes and ~1800 unique interactions. Through automated genetic mapping and whole-genome sequencing, we also isolated an unbiased, experimental set of ~200 spontaneous suppressor mutations that correct the fitness defects of deletion or hypomorphic mutant alleles. Integrating these results yielded a global suppression network.

The majority of suppression interactions identified novel gene-gene connections, thus providing new information about the functional wiring diagram of a cell. Most suppression pairs connected functionally related genes, including genes encoding members of the same pathway or complex. The functional enrichments observed for suppression gene pairs were several times as high as those found for other types of genetic interactions; this highlighted their discovery potential for assigning gene function. Our systematic suppression analysis also identified a prevalent allele-specific mechanism of suppression, whereby growth defects of hypomorphic alleles can be overcome by mutations that compromise either protein or mRNA degradation machineries.

From whole-genome sequencing of suppressor strains, we also identified additional secondary mutations, the vast majority of which appeared to be random passenger mutations. However, a small subset of genes was enriched for secondary mutations, several of which did not affect growth rate but rather appeared to delay the onset of the stationary phase. This delay suggests that they are selected for under laboratory growth conditions because they increase cell abundance within a propagating population.

CONCLUSION: A global network of genetic suppression interactions highlights the major potential for systematic studies of suppression to map cellular function. Our findings allowed us to formulate and quantify the general mechanisms of genetic suppression, which has the potential to guide the identification of modifier genes affecting the penetrance of genetic traits, including human disease.

Global analysis of genetic suppression. Genetic suppression interactions occur when the detrimental effects of a primary mutation can be overcome by a secondary mutation. Both literature-curated and experimentally derived suppression interactions were collected and yielded a genetic suppression network. This global network was enriched for functional relationships and defined distinct mechanistic classes of genetic suppression.

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Mapping a global suppression network

Enrichment of suppression gene pairs

Mechanistic suppression classes

Global analysis of genetic suppression. Genetic suppression interactions occur when the detrimental effects of a primary mutation can be overcome by a secondary mutation. Both literature-curated and experimentally derived suppression interactions were collected and yielded a genetic suppression network. This global network was enriched for functional relationships and defined distinct mechanistic classes of genetic suppression.

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Exploring genetic suppression interactions on a global scale

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Genetic suppression occurs when the phenotypic defects caused by a mutation in a particular gene are rescued by a mutation in a second gene. To explore the principles of genetic suppression, we examined both literature-curated and unbiased experimental data, involving systematic genetic mapping and whole-genome sequencing, to generate a large-scale suppression network among yeast genes. Most suppression pairs identified novel relationships among functionally related genes, providing new insights into the functional wiring diagram of the cell. In addition to suppressor mutations, we identified frequent secondary mutations, in a subset of genes, that likely cause a delay in the onset of stationary phase, which appears to promote their enrichment within a propagating population. These findings allow us to formulate and quantify general mechanisms of genetic suppression.

Although causative variants have been identified for many Mendelian disorders, challenges remain in understanding how genetic variants combine to generate phenotypes. Great progress has been made in mapping and interpreting genetic interactions in yeast, by using growth rate as a proxy for fitness. High-throughput genetic interaction studies have identified hundreds of thousands of negative and positive interactions, in which the fitness defect of a yeast double mutant is either more or less severe, respectively, than the expected effect of combining the single mutants (Fig. 1A) (1, 2). Positive interactions indicate that the phenotypic effects associated with detrimental mutations can be masked or overcome and may explain why certain individuals are healthy despite carrying severe disease-causing mutations (3).

Positive interactions can be further classified by their relative strength, ranging from masking, in which the double mutant fitness is higher than expected but less than or equal to that of the slowest growing single mutant, to suppression, in which the double mutant is healthier than the slowest growing single mutant and possibly has a fitness that is comparable to that of wild type (Fig. 1A) (1, 4). These classes of positive interactions can represent biologically distinct functional relationships (4, 5). Most positive interactions identified by systematic genetic interaction screens in yeast, based on synthetic genetic array (SGA) analysis with loss-of-function mutations (2, 6), are relatively weak masking interactions (fig. S1A), such as the positive interactions that occur among genes within the same nonessential complex or pathway (7). By contrast, stronger suppression interactions remain largely unexplored.

Spontaneous suppressor mutations can be selected to overcome the fitness defect associated with a specific mutant allele. Extragenic suppressor mutations encompass two basic classes: (i) informational suppressors that change the protein translational or mRNA transcriptional machinery, such that the primary mutation is reinterpreted, and (ii) functional suppressors in which a mutation in a second gene functionally compensates for the defect associated with the primary mutation (8). Here, our major goal was to investigate the general principles of functional suppression by assembling a global network of these interactions, which should provide new mechanistic insights about protein function and enable the ordering of components of biological pathways.

A network of literature-curated suppression interactions

To capture existing suppression interactions in Saccharomyces cerevisiae, we examined ~6000 potential interactions in ~1700 published papers derived from the BioGRID’s “synthetic rescue” data set (9). From each interaction, we annotated the type of suppressor mutation (e.g., spontaneous mutation or deletion allele); the type of mutation that is being suppressed, which we refer to as a “query” mutation; and the use of specific conditions (e.g., a drug or specialized carbon source). Suppression interactions that were intragenic, involved a specific phenotype other than growth, or included more than two genes were excluded from the final data set. We also removed suppression interactions derived from high-throughput experiments or dosage interactions in which either the query or the suppressor was overexpressed. The resulting literature-curated network encompassed 1304 genes and 1842 unique suppression interactions (table S1). We visualized this network using a force-directed layout (10), so that query genes that share a common suppressor tend to be positioned together (Fig. 1B). Most query genes (69%) are suppressed by one or two suppressor genes, whereas a small subset of queries (5%) have numerous (10 to 27) reported interactions (fig. S1B). Despite the relatively low average network degree, genes involved in highly studied processes, such as DNA replication and repair or chromatin and transcription, tend to group together because of their shared suppression interactions (Fig. 1B).

Combining data from multiple studies can reveal suppression mechanisms between pathways or protein complexes that may not be apparent from any individual study. Indeed, a subnetwork focused on DNA replication and repair pathways showed that many of the interactions appear to represent the activation of alternative DNA repair pathways (Fig. 1C). For example, mutations that perturb Rad51-dependent homologous recombination (HR) often lead to toxic chromosomal deletions or rearrangements due to increased repair of double-strand DNA breaks by nonhomologous end joining (NHEJ) (11). In this case, suppression can occur through...
Fig. 1. A global network of literature-curated suppression interactions for *S. cerevisiae*. (A) Genetic interaction classes. When two single mutants (xxxΔ and yyyΔ) have a relative fitness of 0.8 and 0.7, the expected fitness of the resultant double mutant (xxxΔ yyyΔ) based on a multiplicative model is 0.8 × 0.7 = 0.56. A negative genetic interaction occurs when the observed double mutant fitness is lower than this expected fitness. A masking positive interaction occurs when the fitness of the double mutant is greater than expected, but lower or equal to that of the slowest growing single mutant. Suppression positive interactions occur when the double mutant fitness is greater than that of the slowest growing single mutant. (B) A global network of literature-curated suppression interactions for *S. cerevisiae*. Genes are represented as nodes and interactions as edges. The nodes were distributed using a force-directed layout, such that genes that share a suppressor tend to be close together on the network. Genes involved in chromatin and transcription or DNA replication and repair are highlighted in magenta and cyan, respectively. (C and D) Regions of the global network highlighting suppression interactions between complexes and pathways involved in chromatin and transcription (C) or DNA replication and repair (D) are shown. Arrows point from the suppressor to the query. PCNA, proliferating cell nuclear antigen.
NHEJ inactivation, which favors double-strand break repair by the compromised, but more accurate, HR machinery (11). Similar trends are observed for genes involved in transcription, for which suppression interactions between pathways mainly represent activation or repression of transcription (Fig. 1D). For example, mutations in genes encoding Mediator or RNA polymerase II subunits can reduce transcription efficiency, which can suppress the toxic effects of derepressed transcription caused by loss-of-function mutations in the NC2 transcription regulator complex (12). Thus, by integrating data from hundreds of papers, we derived a suppression network that provides insight on general suppression relationships and the ordering of pathways and complexes within a biological process.

**Suppression interactions within and across cellular processes**

Consistent with other biological networks (2, 13–15), many suppression interactions occurred between functionally related genes, such that a query mutant tended to be suppressed by another gene annotated to the same biological process (Fig. 2A). Genes connected by suppression interactions also tended to be coexpressed and encode proteins that function in the same subcellular compartment and/or belong to the same pathway or protein complex (Fig. 2B). The extent of functional relatedness between suppression gene pairs did not depend on the conditions under which the interaction was identified (e.g., a specific drug or carbon source), or whether the suppressor was isolated as a spontaneous suppressor mutation as opposed to an engineered allele that was directly tested for an interaction (fig. S2A). However, the frequency of shared complex membership was significantly higher for gene pairs in which the suppressor gene carried a gain-of-function mutation compared with gene pairs involving loss-of-function suppressor mutations (P = 0.01, Fisher's exact test). Thus, when a query mutation perturbs a subunit of a complex, compensating mutations in another subunit can be gain of function—for example, by stabilizing the complex.

Notably, the functional enrichment observed in the genetic suppression network was substantially stronger than in a global network of negative and positive genetic interactions generated with SGA (6) (Fig. 2B). In fact, most positive genetic interactions identified in the global SGA network, especially among loss-of-function alleles of essential genes, do not overlap with other functional interaction data. Suppression interactions thus constitute a special class of positive genetic interaction that captures highly specific functional relationships between gene pairs (fig. S2B).

Despite their tendency to connect functionally related genes, suppression interactions also connect different biological processes. These interactions often occurred between genes involved in related processes, such as Golgi, endosome, or vacuole sorting and ER-Golgi traffic (Fig. 2A). Note that genes involved in protein degradation suppress growth defects associated with mutation of genes involved in many different biological processes.

**Fig. 2. Properties of the suppression network.** (A) Frequency of suppression interactions connecting genes within and across indicated biological processes. Node size reflects fold enrichment for interacting gene pairs observed for a given pair of biological processes. Significance of the enrichment was determined by Fisher's exact test, comparing the observed frequency of suppression interactions between two given functional categories with the global frequency. The total number of suppression interactions involving genes annotated to a particular process is indicated. Kinet., kinetochore. (B and C) Fold enrichment for (B) colocalization, GO coannotation, coexpression, same pathway membership, and same complex membership for gene pairs involved in different types of genetic interaction (GI); and (C) overlap of literature-curated suppression interactions with dosage suppression interactions (13), or with negative and positive genetic interactions identified by SGA analysis using either an intermediate or a stringent interaction score threshold (6). A Fisher's exact test was performed to determine statistical significance of the results. (D) An example of a gene pair showing suppression, dosage suppression, and negative genetic interactions.
processes. This central role for protein turnover in the suppression network likely reflects a more general mechanism whereby growth defects of conditional temperature-sensitive (TS) alleles of essential query genes, which are often hypomorphic (partially functional) even at a permissive temperature, can be overcome by additional mutations that weaken the protein degradation machinery and elevate protein levels.

**Overlap with other genetic networks**

The suppression network shows significant overlap with a dosage suppression network (13) \((P = 2 \times 10^{-34})\); Fisher's exact test) and with SGA-derived positive and negative genetic interaction networks (2, 6) \((P = 5 \times 10^{-47} \text{ and } P = 1 \times 10^{-35})\), respectively, Fisher's exact test). The overlap with positive genetic interactions (fivefold enrichment) (Fig. 2C) is expected, as suppression interactions are an extreme type of positive interaction. Indeed, this overlap increases (11-fold enrichment) for stronger positive genetic interactions. The overlap of the suppression network with dosage-suppression interactions associated with gene overexpression reflects that overexpression may lead to a gain-of-function phenotype (16) and suppression can involve gain-of-function alleles (Fig. 2C and fig. S2C). Gain-of-function suppressor mutations also explain the 2.5-fold enrichment for negative genetic interactions between loss-of-function alleles (Fig. 2C and fig. S2C). For example, whereas the growth defect associated with loss-of-function mutations in CDC25, which encodes the guanine nucleotide exchange factor that activates Ras2, can be suppressed by gain-of-function mutations in RAS2, loss-of-function mutations in RAS2 exacerbate the cdc25 growth defect, thereby causing a synthetic lethal negative genetic interaction (Fig. 2D). Despite the overlap with other genetic networks, most suppression interactions (78%) are specific to the suppression network and thus provide novel insights into the functional wiring diagram of a cell.

**Systematic identification of spontaneous suppressor mutations**

Literature-curated data can come from specific hypothesis-driven experiments and may thus be biased (15, 17). We therefore compared the curated suppression network to an independent experimental set of spontaneous suppressor mutations identified through the large-scale application of SGA analysis. In SGA, a specific natMX-marked query mutation is crossed to an array of ~5000 kanMX-marked deletion mutants, to systematically construct a complete set of haploid natMX- and kanMX-marked double mutants (18, 19). This also represents a genome-wide set of two-factor crosses, enabling us to scan the query strain genome for the presence of an unmarked extragenic suppressor locus, which SGA analysis reveals as a collarine set of small colonies spanning the genomic location of the suppressor mutation, which we refer to as a linkage group (20, 21) (fig. S3A). In total, we completed 7056 full-genome SGA screens, involving mutant strains carrying deletion or hypomorphic alleles of 5845 different genes (2, 6). In 251 SGA screens (~49%), we identified a linkage group that suggested the presence of a spontaneous extragenic suppressor mutation (tables S2 and S3).

The 251 candidate suppressor strains were analyzed by whole-genome sequencing, and for 216 (86%) of these, a mutation was discovered within the suppressor locus identified by SGA (fig. S3A and table S2). Almost all (98%) of these mutations were subsequently confirmed by Sanger sequencing (table S2). For 24 genes, multiple independently generated query strains carried a potential extragenic suppressor mutation (table S2). In 13 (54%) of these 24 cases, the candidate extragenic suppressor mutations were in the same gene, whereas in the remaining 11 cases, two different suppressor genes were identified. In three instances, these different suppressor genes encoded known members of the same complex.

We next validated candidate suppressor genes using several genetic tests, including plasmid-based complementation assays and tetrat analysis of meiotic progeny derived from crossing each suppressor strain to a wild-type strain, a strain with a marked deletion that was genetically linked to the candidate suppressor, or a strain carrying a deletion or hypomorphic allele of the suppressor gene (fig. S3A) (21). Of the suppressor interactions, 88% gave a positive result in at least one assay (table S2). Based on these assays and the type of mutation, one-third (33%) of the suppressor mutations appeared to be gain-of-function, and two-thirds (67%) appeared to be loss-of-function mutations. We also randomly selected four potential loss-of-function and five potential gain-of-function suppressor alleles and introduced those into a diploid strain that was heterozygous for the corresponding query mutation. In all cases, sporulation and tetrad analysis of the resulting diploids confirmed the genetic interaction and identity of the suppressor mutation (table S2 and fig. S3A). Thus, we identified 216 unbiased mutations that arose spontaneously to suppress severe growth defects associated with 146 deletion mutants of nonessential genes and 70 hypomorphic alleles of essential genes (table S2).

Although we observed significant overlap with the literature-curated data set (15 shared interactions, \(P = 1 \times 10^{-29}\); Fisher's exact test), most of the spontaneous suppressor interactions identified through SGA (92%) have not been reported previously; this indicates that the yeast genetic suppression network has remained largely unexplored. The experimentally derived suppressor interactions showed similar significant enrichments as the literature-curated set for different types of genetic interactions, as well as for functionally related gene pairs, suggesting that suppression interactions in both networks define close functional relationships between genes and share the same basic properties (fig. S3, B and C).

**Suppression interaction magnitude correlates with functional relatedness**

Given that suppression interactions tend to connect functionally related genes, we examined whether the relative magnitude of a given suppression interaction was indicative of the extent of functional overlap. We estimated the relative magnitude of suppression for our systematic interactions (table S4) (27), ranked the suppression pairs by suppression magnitude, and calculated the fraction of functionally related pairs for the 33% strongest and weakest suppression interactions (fig. S4). Gene pairs exhibiting more severe suppression interactions showed stronger enrichments for various measures of functional relatedness (fig. S4), in line with what has been described for positive and negative genetic interactions (2). Thus, large improvements in fitness appear to be caused by mutations in genes that are functionally similar to the query, whereas weaker suppression may be achieved by more general or diverse mechanisms.

**Systematic analysis identifies suppressor hubs**

The literature-curated network is enriched for genes involved in highly studied processes, such as chromatin and transcription, as well as DNA replication and repair (Fig. 3A). In contrast, in the experimentally derived network, queries and suppressors were more evenly spread over the various biological processes. As we found for the literature network (Fig. 2), genes involved in protein degradation were underrepresented in the spontaneous suppression interactions identified through SGA (92%) have not been reported previously; this indicates that the yeast genetic suppression network has remained largely unexplored. The experimentally derived suppressor interactions showed similar significant enrichments as the literature-curated set for different types of genetic interactions, as well as for functionally related gene pairs, suggesting that suppression interactions in both networks define close functional relationships between genes and share the same basic properties (fig. S3, B and C).

It is noteworthy that suppressed queries with roles in ribosome biogenesis and translation were underrepresented in the literature but overrepresented in our systematic data set (Fig. 3A). This enrichment was driven by a set of 34 query genes, each encoding a component of the mitochondrial translation machinery. All 34 queries were suppressed by missense mutations in the \(a, \beta, \gamma\) subunits of the \(F_0\) domain of the mitochondrial adenosine triphosphate (ATP) synthase, and the majority of the substituted residues were located at the interfaces between these subunits (fig. 3B). Mutations in the same mitochondrial ATP synthase subunits also suppressed deletion alleles of mitochondrial DNA and RNA polymerase genes, as well as three relatively uncharacterized genes: IRC69, PET330, and YPR117W (table S2). All of these query mutations led to loss of the mitochondrial genome (mtDNA), which results in decreased growth due to a defect in the import of proteins into the mitochondria (27) (Fig. 3, C and D, and fig. S5A). The ATP synthase suppressor mutations could restore both fitness and mitochondrial protein import in the absence of mtDNA (Fig. 3, C and D, and fig. S5B). Note that an
activity of the ATP synthase other than ATP synthesis was required for this suppression phenotype (fig. S5, C and D). Although the mechanism by which the suppressor mutations increase protein import is unclear, one possibility is that the mutations reverse ATP synthase activity to generate ADP\(^{3-}\) instead of ATP\(^{4-}\). The charge difference between these two nucleotide phosphates could be exploited by adenine nucleotide translocators to rebuild the mitochondrial membrane potential, which is lost in the absence of mtDNA and is thought to be required for protein import into the mitochondria (Fig. 3D) (24).

**Suppressor identification can predict novel gene function**

The functional relationship observed between a query mutant and its suppressor can be exploited to assign gene function to previously uncharacterized genes. For example, in our systematically mapped suppressor network, we found that loss-of-function mutations in an uncharacterized gene, YMR010W, suppressed the growth defect of mon2Δ mutants (Fig. 4, A and B). Ymr010w belongs to the family of PQ-loop proteins, some of which function as membrane transporters (25), and localizes to both the Golgi and late endosomes (fig. S6A). Mon2 is distantly related to the Sec7 family of guanine nucleotide exchange factors and physically interacts with Dop1, a

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**Fig. 3. The mitochondrial F\(_1\) ATPase (adenosine triphosphatase) is a suppressor hub in the systematic suppression network.** (A) The distribution of query and suppressor mutants in both the literature-curated and the systematic experimental network across different biological processes. Node size reflects fold enrichment or depletion for query and suppressor mutants observed for a given biological processes. Significat enrichment or depletion was determined by Fisher’s exact test, comparing the observed to the expected proportion of genes in each functional category. Bonferroni-corrected \(P\) values are indicated. (B) Bottom view, facing the inner membrane from the mitochondrial matrix, of the yeast mitochondrial F\(_1\) ATPase structure 2HLD. Residues that were found to suppress the growth defect of mitochondrial transcription or translation mutants are highlighted in red. Orange spheres represent the nucleotides bound to the catalytic sites. (C) Fraction of wild-type and ATP synthase-mutant cells either with intact (\(r^+\)) or (partially) deleted (\(r^-\)) mtDNA that show mitochondrial localization of GFP fused to a mitochondrial-targeting signal (MTS-GFP). Averages (\(n = 4\)) and SD are shown. (D) Model of ATP synthase–dependent suppression of mitochondrial mutants (top) and corresponding representative images of MTS-GFP import (bottom). Localization of outer mitochondrial membrane protein mCherry-Fis1 shows the presence and position of mitochondria. ETC, electron transport chain; \(\Delta\Psi_m\), inner mitochondrial membrane potential; ANT, adenine nucleotide translocator. Scale bar, 5 \(\mu\)m.
Fig. 4. Characterization of YMR010W (ANY1). (A) Predicted membrane topology of Ymr010w. Sites of suppressor mutations, ubiquitination, and phosphorylation are indicated. Single-letter abbreviations for the amino acid residues are as follows: A, Ala; C, Cys; D, Asp; E, Glu; F, Phe; G, Gly; H, His; I, Ile; K, Lys; L, Leu; M, Met; N, Asn; P, Pro; Q, Gln; R, Arg; S, Ser; T, Thr; V, Val; W, Trp; and Y, Tyr. (B) Suppression of the growth defect caused by a mon2Δ deletion allele, or TS alleles dop1-1 and neo1-2, by deletion of YMR010W. Series of 10-fold dilutions of exponentially growing cultures of the indicated strains were spotted on plates with YPD medium and incubated at either 22°C or 38°C for 2 days. (C) Deletion of YMR010W restores membrane asymmetry in neo1-2 cells. Wild-type, ymr010wΔ, neo1-2 and neo1-2 ymr010wΔ cells were grown at 34°C in the presence of the phosphatidylserine (PS) targeting peptide papuamide A, or the phosphatidylethanolamine (PE) targeting peptide duramycin. Growth relative to vehicle-treated wild-type strain is plotted. SEM is indicated by shading (n = 2 to 3). (D) Intracellular distribution of PS, visualized using GFP-LactC2 (31). Shown are representative confocal fluorescent micrographs of exponentially growing cells of the indicated strains. The fraction of cells was calculated for each of the following groups: those (i) that showed diffuse cytosolic fluorescence or (ii) localization of GFP-LactC2 to the plasma membrane, or (iii) in which GFP-LactC2 was partially localized to distinct internal structures. Measurements were performed in triplicate on at least 100 cells, and averages are shown. (E) Model of suppression of flippase mutants by loss of Ymr010w.
conserved membrane protein involved in endosome to Golgi transport, as well as Neo1, an essential member of the phospholipid flippase family (26, 27). When tested directly, we found that a ymr010wΔ deletion allele also suppressed the growth defects of neo1-2 and dopl-1 TS mutants (Fig. 4B). Moreover, a ymr010wΔ deletion allele suppressed the lethality associated with deletion alleles of the essential genes NEO1 and DOP1 (fig. S6B). Loss of YMR010W function can thus bypass the requirement for the Mon2/Dop1/Neo1 module.

The essential function of the Mon2/Dop1/Neo1 module is likely performed by Neo1, which is thought to flip phosphatidylserine (PS) and phosphatidylethanolamine (PE) from the exoplasmic to the cytoplasmic leaflet of membrane bilayers and thereby to establish an asymmetric distribution of these lipids (28). A neo1-2 TS mutant is defective in establishing membrane asymmetry. This leads to hypersensitivity to papuamide A and duramycin, bioactive peptides that disrupt membranes through the binding of exposed PS and PE, respectively (28–30), and reduced plasma membrane localization of green fluorescent protein (GFP)–LactC2, a probe for visualizing the distribution of PS over cyttoplasmic membrane leaflets (31) (Fig. 4, C and D). Overexpression of YMR010W also led to reduced levels of PS at the cytoplasmic leaflet of the plasma membrane, and accumulation of GFP-LactC2 in internal structures (Fig. 4D), thus mimicking the phenotype of a neo1-2 mutant. We found that a ymr010wΔ deletion allele suppressed both the sensitivity of a neo1-2 TS mutant to papuamide A and duramycin (Fig. 4C), and the neo1-2 GFP-LactC2 localization defect (Fig. 4D). The absence of these phenotypes suggests that the neo1-2 phospholipid distribution defects are corrected in the double mutant.

In addition to suppressing loss of Neo1 function, a ymr010wΔ deletion allele suppressed the cold sensitivity caused by loss of the flippase Dns2 (fig. S6C). Moreover, neo1Δ lethality was no longer suppressed by ymr010wΔ in the absence of Dns2 (fig. S6C). An intriguing possibility is that Ymr010w functions as a scramblase that transports PS and PE bidirectionally to at least partially collapse the membrane asymmetry established by Neo1 and other flippases (Fig. 4E).

Deletion of YMR010W would then allow Dns2, possibly with the help of other flippases, to more easily establish membrane asymmetry in the absence of Neo1. We named the YMR010W open reading frame ANY7 for antagonizes Neo1 yeast phospholipid flippase.

**Frequent secondary mutations delay the onset of stationary phase**

Whole-genome sequencing revealed that, besides the suppressor mutation, each suppressor strain carried on average eight additional secondary mutations (table S5). Unlike the suppressor mutations, none of these secondary mutations affected exponential cell growth enough to be detected by SGA mapping analysis (table S3), suggesting the majority are random mutations that arose during DNA replication. We therefore refer to these additional secondary mutations as “passenger” mutations. We identified a similar number of passenger mutations in a control set of 72 strains that did not carry a suppressor mutation that affects growth of the query mutant (table S5). Of the 304 strains that were sequenced at a coverage >10 times, only one query strain, deleted for PMS1 that encodes a mismatch repair protein, displayed a mutator phenotype, exhibiting a relatively large number (76%) of passenger mutations. In total, we identified 2024 unique passenger mutations, of which 946 were in coding regions, affecting 744 protein or RNA-encoding genes. The fraction of missense, nonsense, and frameshift mutations was substantially smaller among the passenger mutations than among the suppressor mutations (Fig. 5A). In fact, most of the passenger mutations (64%) resulted in synonymous changes or mapped to intergenic regions (Fig. 5A). Furthermore, passenger missense mutations occurred less frequently in essential genes, were predicted to be less deleterious, were less often at protein–protein interaction interfaces, and occurred more often in disordered protein regions than suppressor missense mutations (Fig. 5B). Thus, the majority of the passenger mutations, which have no effect on exponential growth of the query strain, have a lower putative functional impact than the suppressor mutations that do affect query strain cell growth.

A previous study suggested that deletion of a particular query gene may select for further genetic changes, such as the occurrence of specific secondary nonsuppressor mutations (32). However, we did not observe a correlation between the number of passenger mutations and the fitness of the query strain (fig. S7A). Moreover, genes carrying passenger mutations do not tend to be coannotated or coexpressed with the corresponding query or suppressor gene (fig. S7B). In addition, we did not find any enrichment for particular GO terms among query genes that shared the same passenger mutation, or for shared passenger mutant genes among multiple, independent isolates of a particular query mutant strain. However, we found that 10 strains that all carried a suppressor mutation in ATP2 but had different query mutations involved in mitochondrial transcription or translation, also harbored a third mutation in HEM1, TPP1, or HAP1. These three genes are important for heme biosynthesis, and these mutations may thus be selected for to maintain heme homeostasis in the absence of mitochondrial transcription, translation, or ATP synthase activity. Still, in most cases, different isolates of the same query suppressor strains did not contain mutations in the same passenger genes, and most passenger genes were not functionally related to either the query or the suppressor gene, indicating that passenger mutations are not generally dependent on preexisting mutations.

We did find several genes that were mutated in a large fraction of the sequenced strains; this suggested that they may be adaptive and may not represent innocuous passenger mutations (Fig. 5C). Of all sequenced strains, including wild-type controls, 29% carried unique mutations in WHI2, IRA1, IRA2, RIM15, CUP9, and/or UBC3. Multiple experimental evolution studies have identified a similar set of frequently mutated yeast genes (33–35). Most of the mutations were frameshift or nonsense mutations, suggesting a selection for loss-of-function of these genes (Fig. 5C). Exponential growth rates of seh12a, iru2a, rim15a, and ubc3a deletion mutants were not enhanced relative to a his3A deletion mutant control. It thus appears that there was no selection for these frequent secondary mutations on the basis of an increased maximum growth rate (fig. S7C). Note that Whi2, Ira1, Ira2, and Rim15 are all negative regulators of the RAS/cyclic adenosine monophosphate (cAMP)/protein kinase A (PKA) pathway, which, in response to glucose, stimulates population expansion (36–39). When glucose levels become limited, the RAS/cAMP/PKA pathway is repressed, thereby causing cells to stop dividing and enter stationary phase. Disruptive mutations in WHI2, IRA1, IRA2, or RIM15 may cause a delayed response to low glucose levels that enables a few additional rounds of cell division before cells enter stationary phase and, thereby, lead to increased representation of these mutants after serial passaging under laboratory conditions. We constructed mixed populations consisting of a strain deleted for one of the frequently mutated genes and a wild-type strain and followed their ratio for six rounds of serial passaging under conditions with a relatively prolonged stationary phase (21). Indeed, the relative abundance of strains deleted for WHI2, IRA2, RIM15, or UBC3 increased with each round of serial passaging, whereas five control mutant strains maintained abundances similar to or lower than the wild-type reference strain (Fig. 5D and fig. S7D). Similar results were obtained for IRA1 and IRA2 in another strain background, W303 (fig. S7, E and F). Thus, our data suggest that the vast majority of passenger mutations are random and not dependent on the query or suppressor mutation and, further, that a few additional secondary mutations arise at high frequency because of a selection for mutants that delay the onset of the stationary phase.

**Mechanistic categories of suppression interactions**

We classified the suppression interactions into distinct mechanistic categories on the basis of the functional relationship between query and suppressor. Most queries (54%) reported in the literature or identified by our systematic analysis are suppressed by mutations in functionally related genes (class “A”) (Fig. 6, A and B). These functional connections can be further divided into four subclasses. Subclass “A1” includes 135 interactions from the literature and systematic networks, in which both the query and the suppressor genes encode members of the same protein complex. These particular interactions can reflect a mechanism whereby the suppressor represents a gain-of-function mutation (fig. S2A). Subclass “A2,” to which 201 interactions from our
network were assigned, describes cases where the query mutant growth defect is suppressed by a mutation in a gene that is annotated to the same pathway. In the case of loss-of-function suppressor mutations, the suppressor gene often has antagonistic effects compared with the query gene (e.g., Fig. 4). Subclass “A3” involves suppression by a different, but related, pathway and explains 195 interactions in our networks. In this scenario, the growth phenotype caused by absence of a specific cellular function required for normal cell growth is suppressed when an alternative pathway is rewired to re-create the missing activity (e.g., Fig. 3). Finally, subclass “A4” consists of gene pairs that are annotated to the same biological process but for which pathway or complex annotation data were not available for both genes.

In addition to suppression interactions between functionally related genes, suppression interactions involving hypomorphic (partial loss-of-function) alleles—such as conditional TS alleles of essential genes—revealed a different and more general class of suppressors that affect expression of the query gene. This type of suppression (Figs. 2A and 3A) can be achieved by stabilizing a mutant mRNA or protein through the perturbation of pathways or complexes that regulate mRNA or protein turnover (Fig. 6A, class “B” and “C”). Although this type of suppression is rarely described in the literature, 48% of the hypomorphic queries in our experimental data set are suppressed by mutations in protein degradation or mRNA decay genes (Fig. 6B); this indicates that this type of allele-specific suppression is one of the main routes through which partial loss-of-function alleles can be suppressed. Of the suppression interactions, 60 to 70% fall into one of these mechanistic classes, as compared with only 34% of positive genetic interactions identified by SGA (6) and 11% of passenger-query pairs. Thus, positive genetic interactions that are true suppression interactions often show high functional and mechanistic specificity.

Discussion

A global, literature-curated network of genetic suppression interactions (Fig. 1) showed that the majority of suppression interactions linked functionally related genes. Moreover, suppression interactions overlapped significantly with other types of genetic interactions (Fig. 2). Systematic suppression analysis confirmed these general properties of suppression and further showed that suppression of hypomorphic alleles often occurs via loss of protein or mRNA degradation, a finding that was less obvious in literature-curated data (Fig. 6). The underrepresentation of this class of interactions in the literature is consistent with what has been reported for dosage suppression interactions (13) and may reflect that mechanistic studies focused on the functional analysis of a particular gene or pathway are less likely to

Fig. 5. Characterization of potential passenger mutations. (A) Distribution of suppressor and potential passenger mutations over variant effect classes. Only SNPs are considered, as reliable structural variant calls (deletions, insertions, or inversions involving >5 base pairs) were only available for suppressor mutations. The RNA class refers to mutations in an RNA species such as a noncoding, ribosomal, or transfer RNA. (B) The fraction of all suppressor or potential passenger missense mutations that map to an essential gene, at a protein-protein interaction (PPI) interface, or at a disordered region of a protein, and the predicted deleteriousness of these mutations (SIFT scores: 0 = extremely deleterious and 1 = benign). P values were calculated using Fisher’s exact test, except for the SIFT analysis, in which a Mann-Whitney test was used. (C) The percentage of strains in which a particular gene carries a passenger mutation is plotted against the chromosomal position of the gene. Genes that are recurrently mutated in >2% of the sequenced strains are highlighted, and the distribution of the mutations over variant effect classes is shown. (D) Differentially fluorescently labeled cells of the indicated mutants [labeled with red fluorescent protein (RFP)] and wild type (GFP) were mixed, and the ratio of RFP to GFP was followed for six rounds of serial passaging on agar plates. Shading represents the SD, n = 12.
report nonspecific suppressors. Nevertheless, an understanding of the prevalence of this form of suppression could be important when interpreting a genotype-to-phenotype relationship. Even though the genes encoding proteasome or mRNA-decay components are essential in human cell lines (40–42), we anticipate that genetic variation that subtly modulates the activity of these modules may exhibit genetic interactions associated with a decreased disease risk for a variety of human disorders. As in yeast, these processes may thus buffer a range of detrimental mutations in humans and, thereby, modify numerous different disease phenotypes.

Despite the prevalence of these general suppression mechanisms, most suppression gene pairs showed a close functional relationship (Fig. 6), so that genetic suppression can be used to assign function to a previously uncharacterized gene (Fig. 4). The suppressor interactions identified in our systematic screen resulted from the direct selection of spontaneous mutations during standard laboratory growth of a query mutant whose fitness was compromised. In total, ~3% of strains in the yeast nonessential deletion mutant collection and ~4% of the strains in the hypomorphic essential gene mutant collections showed evidence of a suppressor locus when screened by SGA. Whole-genome sequencing of 251 potential suppressor strains did not reveal any instances of suppression via aneuploidy, a mutational event involving copy number variation of many genes, possibly because aneuploides are not necessarily revealed by SGA genetic mapping or because these events come at a fitness cost (43). Although SGA suppressor mapping can theoretically identify multiple suppressor mutations within one strain (20), no query strains with multiple suppressor linkages were identified. This suggests that the direct selection for spontaneous suppressors does not mimic adaptive evolution of wild-type strains in nutrient-limited conditions, in which aneuploides and mutations in multiple genes, each contributing small fitness increases, combine to collectively produce a robust suppression phenotype (35, 44). In contrast, we found that there is often a single direct suppression strategy because for most (~67%) of the queries for which we isolated several independent suppressor mutations, these recurrently occurred within the same single suppressor gene or within genes that encode subunits of the same complex. For example, we found that large increases in fitness are mainly achieved by mutations in genes that have a close functional relationship to the query gene (fig. S4). Thus, only a few, very specific mutational events appear to be able to substantially increase the fitness of a particular query mutant.

Besides the suppressor mutation, each strain also carried, on average, eight additional passenger mutations that did not have a measurable effect on exponential growth rate. In a previous, but relatively limited, study, it was suggested that the deletion of a query gene in the deletion mutant collection often selects for further genetic changes (32). Although this is true for suppressor

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Fig. 6. Mechanistic classes of suppression. (A) Suppressors and query genes often have a functional relationship (class “A”). In a situation where a query (protein A) activates a protein B, which is required for normal growth, suppression can take place in multiple ways. For example, the suppressor (protein C) can be part of the same complex as the query, and gain-of-function mutations in C can restore the activity of B (class “A1”). Alternatively, the suppressor and query may be members of the same pathway, and the suppressor (protein D) may inactivate or inhibit B. Loss of D may thus suppress by partially restoring the activity of B (class “A2”). The suppressor (protein E) can also function in an alternative, but related, pathway, whose activity can be slightly altered to restore the activity of B (class “A3”). Suppression interactions can also occur among pairs of genes that do not share a close functional relationship. For example, partial loss-of-function query alleles may carry mutations that destabilize the protein or mRNA, leading to a fitness defect caused by reduced levels of the query protein. This can be suppressed by a loss-of-function mutation in a member of the protein degradation (class “B”) or mRNA decay (class “C”) pathway, which may partially restore the levels of the query protein. NMD, nonsense-mediated mRNA decay. (B) Distribution of suppression interactions, positive genetic interactions (6), and passenger-query pairs across different mechanistic suppression classes.
mutations, we could not find any substantial evidence connecting the query or suppressor mutations to the occurrence of most passenger mutations. Because we did not observe a significant enrichment for functionally related gene pairs among queries and passengers (fig. 5B), we conclude that the occurrence of query-driven non-suppressor mutations is likely rare.

In a mathematical model of bacterial serial passaging, de novo mutations that delay the onset of stationary phase were more likely to fix in a population than mutations that decrease lag time or increase growth or survival rates (45). This may be true for yeast as well, as the growth history of laboratory-grown yeast strains follows a similar pattern of repeating cycles of lag phase, exponential growth, and stationary phase. Indeed, we observe selection for mutations that likely delay the onset of stationary phase in 26% of the sequenced strains (Fig. 5C). These stationary-delay mutations are thus not true “passenger” mutations but are adaptive. However, in contrast to suppressor mutations that cause adaptation to the query mutation, the stationary-delay mutations are adaptive to laboratory passaging. These mutations could come at a cost, as they probably decrease viability during longer periods of starvation (35, 36).

As most (78%) suppression interactions did not overlap with any previously identified genetic interactions, additional suppression mapping will help complete the yeast genetic interaction landscape. Conditional alleles have been developed for nearly all essential yeast genes (6), and thus, suppression interactions could be mapped for the full set of essential genes. Similarly, suppressors of nonessential genes could be identified in a conditional or synthetic lethal context in which the nonessential query has a fitness defect. Although we focused on mapping suppression interactions in yeast, similar suppression studies should be possible in mammalian cells and model systems and may identify new drug targets for query mutations related to human disease (46). As ~6% of human pathogenic variants are fixed in other mammalian species (47), compensatory mutations may be present at a high frequency in natural populations. Understanding genetic suppression may provide insight into how genetic variance accumulates during evolution and more specifically how modifier genes determine the severity of genetic traits, including human disease.

Materials and methods

Detailed materials and methods are available in the supplementary information.

Literature curation

The Saccharomyces cerevisiae “synthetic rescue” data set was downloaded from the BioGRID (9) on 9 November 2012 (version 3.1.49) and on 31 March 2014 (version 3.2.110). In total, these data sets consisted of 5985 interactions described in 1667 papers. Each paper was read in detail, and an interaction was considered a suppression interaction if the double mutant grew substantially better than at least one of the single mutants. For each interaction, suppressor and query allele type and specific conditions were annotated (27). The final data set consisted of 1642 unique interactions, involving 1304 genes (table S1).

Systematic suppressor identification

All suppressor strains were part of either the BY4741 nonessential deletion mutant collection (MATa αααα: kanMX4 his3Δ1 leu2Δ0 ura3Δ0 met15Δ0 Euroscarf), the SGA nonessential deletion mutant collection (MATα: αααα:kanMX4 can1Δ0 can2Δ0 can3Δ0 his3Δ1 his4Δ1 leu2Δ0 ura3Δ0 met15Δ0) (2) or the corresponding MATa and MATα collections of DAmP or TS mutants of essential genes (6). The presence and genomic location of a spontaneous suppressor mutation were identified by the occurrence of a suppressor linkage group upon screening strains in these collections by SGA analysis (20) (table S3). Potential suppressor strains were subsequently sequenced whole-genome on the Illumina HiSeq 2500 platform using paired-end 100-bp reads. Read mapping and single-nucleotide polymorphism (SNP), as well as indel calling were performed using standard methods (27). Candidate suppressor mutations were confirmed by amplifying the corresponding gene and flanking sequences by polymerase chain reaction, followed by Sanger sequencing (table S2). Suppression interactions were confirmed using plasmid-based complementation assays and tetrad analysis of meiotic progeny derived from crossing each suppressor strain to either a wild-type strain, a strain with a marked deletion that was genetically linked to the candidate suppressor, or a strain carrying a deletion or hypomorphic allele of the suppressor gene (table S2).


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SUPPLEMENTARY MATERIALS

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Materials and Methods

Figs. S1 to S7

Tables S1 to S7

References (48–62)

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Exploring genetic suppression interactions on a global scale
Jolanda van Leeuwen, Carles Pons, Joseph C. Mellor, Takafumi N. Yamaguchi, Helena Friesen, John Koschwanez, Mojca Mattiazi Usaj, Maria Pechlaner, Mehmet Takar, Matej Usaj, Benjamin VanderSluis, Kerry Andrusiak, Pritpal Bansal, Anastasia Barzyshnikova, Claire E. Boone, Jessica Cao, Atina Cote, Marinella Gebbia, Gene Horecka, Ira Horecka, Elena Kuzmin, Nicole Legro, Wendy Liang, Natascha van Lieshout, Margaret McNee, Bryan-Joseph San Luis, Fatemeh Shaeri, Ermira Shuteriqi, Song Sun, Lu Yang, Ji-Young Youn, Michael Yuen, Michael Costanzo, Anne-Claude Gingras, Patrick Aloy, Chris Oostenbrink, Andrew Murray, Todd R. Graham, Chad L. Myers, Brenda J. Andrews, Frederick P. Roth and Charles Boone

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A global genetic suppression network
The genetic background of an organism can influence the overall effects of new genetic variants. Some mutations can amplify a deleterious phenotype, whereas others can suppress it. Starting with a literature survey and expanding into a genomewide assay, van Leeuwen et al. generated a large-scale suppression network in yeast. The data set reveals a set of general properties that can be used to predict suppression interactions. Furthermore, the study provides a template for extending suppression studies to other genes or to more complex organisms.

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