Combinatorial genetic analysis of a regulatory network reveals the importance of higher order epistasis for gene deletion phenotypes

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Abstract

A key challenge in biology is to understand how mutations combine to alter phenotypes. Each genetic variant in a genome can have diverse effects, for example decreasing, increasing, inactivating, or changing the function of a protein or RNA. In contrast, systematic analyses of how mutations interact have typically used a single variant of each gene, most often a null allele. We therefore lack an understanding of how the full range of genetic variants that occur in individuals can interact. To address this shortcoming, we developed an approach to combine >5000 pairs of diverse mutations in a model regulatory network. The outcome of most mutation combinations could be accurately predicted by simple rules that capture the ‘stereotypical’ genetic interactions (epistasis) in the network. However, for individual genotypes, additional, unexpected pairwise and higher order genetic interactions can be important. These include ‘harmonious’ combinations of individually detrimental alleles that reconstitute alternative functional switches. Our results provide an overview of how the full spectra of possible mutations in genes interact and how these interactions can be predicted. Moreover, they illustrate the importance of rare genetic interactions for individuals, including the impact of higher order epistatic interactions that dramatically alter the consequences of inactivating genes.
Main text

Human genomes contain millions of genetic variants. Each of these variants can have diverse effects, for example quantitatively increasing, decreasing or changing the activity of individual genes. Understanding and predicting how the particular combination of variants present in each individual affects molecular processes and phenotypic traits is a fundamental challenge for human genetics and evolutionary biology. To date, however, systematic analyses of how mutations in different genes combine to influence phenotypes have used only one or a few mutations in each gene, most often an inactivating null allele. A more complete understanding of how mutations combine in individuals will require functionally diverse mutations in individual genes to be combined in large numbers.

The GAL regulatory (GALR) system from yeast is a promising model to begin such studies because it is mechanistically well-understood, has relatively few molecular players, and is an important model of gene network function and evolution. This network is required for sensing the sugar galactose and then inducing transport (via Gal2) and the Leloir pathway proteins Gal1p, Gal7p, and Gal10p necessary to metabolize this sugar as a carbon source for growth. The core of this network consists of three regulatory genes and their protein products: GAL4, a transcriptional activator; GAL80, its repressor; and GAL3, which acts as a GAL sensor by inhibiting Gal80p as an activated Gal3p-Galactose-ATP complex. The GAL1 locus, encoding the first Leloir enzyme galactokinase (GALK) Gal1p, is a paralog of GAL3 that can also effect galactose sensing. The ancestral GALK locus likely encoded a bifunctional protein that was duplicated to give rise to the main sensor Gal3p and the kinase Gal1p.

To systematically explore how mutations in the GALR genes GAL4, GAL3, and GAL80 affect GAL pathway output we first tested all combinations of GALR wild-type (WT) or coding sequence deletions (Δ) to give $2^3 = 8$ unique genotypes (Fig 1C and Methods). GAL pathway activation and growth were quantified using flow cytometry to measure cell density and the expression of a Gal1p-YeCitrine (YFP) reporter in two environments: ‘uninducing’ conditions with glucose as the only carbon source and after 12 hours of induction in galactose. As expected, across these 8 genotypes, the GAL pathway exhibits three phenotypic classes: Inducible, Uninducible, and Constitutive (Fig 1D-F, Fig S1, and Supplementary Dataset 1). The WT GALR system is Inducible because it is repressed in glucose and activated in galactose. The single mutants ΔGAL3 and ΔGAL4 result in Uninducible phenotypes because they do not activate GAL expression in any conditions, and ΔGAL80 drives Constitutive phenotypes because GAL expression is activated in glucose. The triple mutant and all double mutant genotypes yielded Uninducible phenotypes except the ΔGAL80 + ΔGAL3 mutant, which is Constitutive because WT GAL4 is not repressed in the ΔGAL80 background. GAL1-YFP expression was highly predictive of growth rate across these genetic backgrounds (Fig 1F).
Next, to investigate how richer spectra of mutations in the three GALR genes interact, we combined together diverse alleles generated by random and directed mutagenesis (Supplementary Tables 1-3 and Methods). These included weak and strong loss-of-function (lof) alleles, gain-of-function (gof) alleles, and phenotypically WT-like alleles. We used these alleles to perform a pairwise combinatorially complete experiment where all pairs of GALR variants were combined together and the resulting phenotype quantified. In total we combined 46 alleles of GAL3, 39 of GAL80 and 43 of GAL4. After transformation, this library included 98% of the possible pairwise genotypes, giving 5317 unique double mutant combinations (Fig 2A and Methods). We quantified GAL pathway expression and growth characteristics for each genotype as before (Fig S1 and Supplementary Dataset 2).

The resulting phenotypes were more diverse than when combining gene deletions (Fig 2B), but most genotypes still grouped into a limited number of phenotypic classes (Fig 2C, Fig SA-D and Methods), with 92% of the 5317 genotypes falling into the Inducible, Constitutive and Uninducible expression classes observed when combining gene deletions (Fig 2C). A further ~5% of 'Leaky' genotypes were closest to Inducible profiles but in glucose exhibited a detectable fraction of ON cells with low mean expression (Fig S3B,C). Finally, 3% of samples fell into a 'Weak expression' class with low maximal expression in glucose and galactose. Neither of these behaviors was observed when combining null alleles.

To test whether the expression phenotypes of the single mutants could predict double mutant galactose growth phenotypes, we first computed the mean growth rate value of each of the 23 unique locus-cluster combinations present in the dataset (Fig 2D). These 23 mean values were used as independent variables to predict the growth rate measurements for all unique double-mutant genotypes whose single mutants matched these locus-cluster pairs. So, for example, the mean of all double mutants of an Uninducible GAL3 plus a Constitutive GAL80 were used to predict each individual measurement of double mutants whose single GAL3 variant was Inducible and whose single GAL80 variant was Constitutive. These mean values explained 89%, 90%, 91% of total growth rate variance for double-mutants in GAL3 vs. GAL4, GAL80 vs. GAL3 and GAL80 vs. GAL4 pairings, respectively (Methods). Across all double mutants, this model explained 91% of growth rate variance, compared to 55% for a standard multiplicative growth rate model. A model considering coarser classifications of phenotypes (Inducible, Constitutive, and Uninducible, 14 parameters) explained 89% of growth rate variance (Fig 2D, Fig S3E-H, Methods, and Supplementary Dataset 2). Thus, despite the strong epistasis in this system, simple models that capture the main ‘stereotypical’ genetic interactions between loci can accurately predict how pairs of diverse mutations interact.
Beyond these stereotypical outcomes, particular combinations of mutations sometimes had very different phenotypes to those expected from their individual phenotypic effects. For example, rare pairings of Constitutive GAL4 alleles and Uninducible GAL80 variants yielded Inducible phenotypes, an example of reciprocal sign epistasis\(^\text{10}\). Similar interactions have been previously described for distinct GAL4-GAL80 allele combinations\(^\text{11-13}\). Other combinations yielding viable Inducible or Leaky phenotypes included pairings of Uninducible GAL3 with Leaky GAL80 variants (Fig 2E and Fig S2). These alternative combinations of GAL80 and GAL4 genetic variants that permit a WT-like phenotype represent examples of ‘harmonious’ combinations\(^\text{14}\): alternative genetic solutions to the core Inducible phenotypic characteristic of the GAL pathway.

We hypothesized that the harmonious combinations of GAL80 and GAL4 alleles could simply reflect reconstitution of the original regulatory network that exists in WT cells, with all GALR genes functioning as in the WT network. Alternatively, the functionality of these mutants might reflect different solutions to the same regulatory task, with the roles of other genes changed\(^\text{15,16}\). To distinguish between these two possibilities, we tested whether mutating the additional GALR genes, GAL3 and GAL1, had the same effect in these functional combinations of GAL80 and GAL4 alleles as in the WT system.

We tested a subset of GAL80 and GAL4 alleles in a combinatorially complete set of genotypes incorporating additional third- and fourth-order deletions in the potential galactose sensors GAL3 and GAL1 (Fig 3A). All Inducible allelic combinations depended on GAL1 or GAL3 for robust growth, reflecting the importance of galactose sensing for pathway induction (Fig 3B,C). However, the consequences of deleting GAL3 or GAL1 sensing activity varied extensively across the different genotypes. For example, whereas the WT switch is completely dependent upon GAL3 (Fig 1D and Fig 3), GAL3 was not required for induction in the Leaky GAL80.07 mutant (Fig 2D and Fig 3), nor was it required for Constitutive expression in GAL4C mutants (Fig 3D). Moreover, GAL80S-1 + GAL4C double mutants were still Inducible when GAL3 was deleted (Fig 3D,E, Fig 4A, Fig S4, Fig S5, and Supplementary Dataset 3).

GAL1’s galactokinase activity is required for growth in galactose, so the dependency of a network on GAL1’s sensing activity cannot be determined by simply deleting the gene. To test the dependency of each genotype on GAL1 sensing activity, we therefore used a strategy in which GALK genes from other species with different galactose sensing mechanisms were expressed from the GAL1 promoter (‘GAL1::GALK’, Fig 3A, Methods). Consistent with GAL1 sensing activity not being required for induction in WT cells, replacing GAL1 with GALK from other species had no effect on gene expression or growth (Fig 3B-C and Fig S4, Fig S5). In the Leaky GAL80.07 background, replacing GAL1 by GAL1::GALK also had no effect. However, it completely prevented growth and expression when GAL3 was also deleted. In
contrast, in GAL4C mutants, GAL1::GALK reverted the Constitutive expression phenotype to a Leaky or Inducible phenotype (Fig 3D and Fig 4A). This would be consistent with GAL4C single-mutant variants being de-repressed in glucose not only because of reduced GAL80 repression (Fig 3D), but also due to positive feedback via GAL1, which, through a baseline level of leaky expression (Fig 3D) and ability to repress GAL80\(^9\), could reach sufficient abundance to constitutively activate the system (Fig 4A). Finally, in GAL4C + GAL80S-1 double mutants, GAL1::GALK reduced growth rate and expression more than deleting GAL3, including one combination where GAL1 sensing activity was essential (Fig 3B-E and Fig 4A).

In this dataset, there therefore exist four 'harmonious' combinations of GALR genes with Inducible phenotypes but differing dependence on the sensing genes GAL3 and GAL1 (Fig 4A). The first combination is the WT itself, which depends on GAL3 completely but does not require GAL1 sensing activity. The second combination is GAL80.07 + WT GAL4, where GAL3 is no longer essential and where deleting GAL1 has no effect except in combination with a GAL3 deletion. The third combination is the Inducible GAL4C + GAL1::GALK background, where GAL3 serves as a single essential sensor, and where GAL1 sensing activity is deleterious, causing Constitutive expression in glucose. Finally, the combination of GAL80S-1 + GAL4C has a flipped dependence on GAL3 and GAL1 compared to the WT system, with GAL3 no longer essential and growth and expression dependent on GAL1.

In summary, by combining mutations with diverse individual effects, our approach provides a more complete view of how genetic variants in different genes combine to alter the activity of a model regulatory network. We found that the phenotypes of thousands of pairs of mutations fell into a small number of phenotypic classes, with the classes of individual mutants predicting very well their combined effects. In other words, once the phenotype of a single mutant is measured, its effect in combination with many other mutations is normally straightforward to predict, even when there is strong epistasis between loci. If this is also true for other systems including human disease, it will mean that accurate phenotypic prediction will often be possible without the need to resort to detailed mechanistic (dynamical) models.

Rare combinations of mutations in the GALR genes did, however, have unexpected outcomes, including combinations of individually detrimental mutations that reconstitute a functional regulatory switch. These and other ‘harmonious’ combinations of mutations constituted a functional system not because they resuscitated the original WT switch but because they formed alternative ‘re-wired’ regulatory networks with altered ‘functions’ for additional components (Fig 4B). For example, combining the individually deleterious GAL4C with GAL80S-1 variants not only restored an Inducible system, an example of reciprocal sign epistasis, but also switched the system’s dependence on the other two GALR genes,
GAL3 and GAL1, an example of higher order epistasis. Such changes in gene essentiality have been widely observed between and within species \(^{17-21}\) but the genetic causes are poorly understood. The altered requirement for GAL1 and GAL3 across combinations of mutations in GAL4 and GAL80 suggests that selective pressures on paralogous genes can substantially change with variation in other molecular players.

Together, our results illustrate how the genetic interactions between diverse alleles can be accurately predicted using models that capture the ‘stereotypical’ epistasis in a system. However they also demonstrate the importance of rare unexpected pairwise and higher order epistasis for the fitness of individual genotypes, and illustrate how higher order epistasis can rapidly change the essentiality of genes.
Figure legends

**Figure 1. Combinatorial genetic analysis of the GAL pathway.**
(a) Overview of GAL pathway regulation. (b) *in vivo* homologous recombination to combine alleles of 3 GALR genes into a chassis strain with all three genes deleted. (c) Cubic representation of combinatorially complete genetics experiments combining alleles in multiple loci. (d) Three phenotypic classes of GALR deletion mutants. Lines and shading are the mean and ± 1 SD of genotypes falling into the given expression class (N=4 for N=2 independent transformations). Blue lines are samples growing in glucose and red lines are samples after 12 hours of growth in galactose. (e) Expression distributions summarized by the fraction of cells ON in galactose and glucose. Points are mean values for each unique genotype, and are coloured to reflect the number of mutations. (f) Fraction ON in glucose and galactose predicts growth rate. Blue line and shaded region are the expectation and 95% confidence interval of a linear model.

**Figure 2. Phenotypic variation from pairwise GALR allele combinations.**
(a) Alleles obtained by mutagenic PCR of a given GALR locus were paired with alleles of the other two loci. (b) Single and double mutants have more diverse phenotypes than gene deletions. Error bars represent the 95% confidence interval of N=2 to N=36 (median N=2) independent replicates across N=1 to N=18 independent transformants (median N=1). Grey line is Y = X. (c) Five classes of expression distributions. Lines and shaded regions are density means and ± 1 SD across genotypes of each class and genotype. Integers and percentages indicate the number of genotypes in each panel. (d) The 23 mean growth rates corresponding to all pairwise combinations of single-mutant expression classes (horizontal axis) were used to predict 5152 double mutant growth rates (vertical axis). (Methods). (e) The fraction of single-mutant combinations leading to a given double-mutant phenotype shown as stacked bar plots. Single-mutant combinations that comprised > 10% of genotypes leading to the indicated double-mutant phenotype are labeled on the chart.

**Figure 3. Higher-order genetic interactions in the GALR pathway.**
(a) Experimental design to determine the requirement for GAL1 and GAL3 sensor activity for GAL pathway induction across combinations of GAL80 and GAL4 alleles (Methods). (b) The effects of deleting GAL1 and GAL3 change depending on GAL80-GAL4 combinations. The mean for the WT GAL80-GAL4 pairs is indicated as a blue circle. Samples deviating significantly from the grey line where Y = X are coloured red (two-tailed t-test, FDR < 0.05 and mean effect ≥ 0.03 µ hr⁻¹). These points reflect harmonious genetic combinations that depended on GAL1 (left facet panel) or GAL1 and GAL3 (right facet panel) for high rates of growth. (c) Requirement for GAL1 or GAL3 for high growth rates from an
initially uninduced state. In the $\Delta GAL3 + GAL1::GALK$ double mutants, the mean expression level for each $GAL4$-$GAL80$ pair in glucose is used as a measure of $GAL$ pathway 'leakiness' or constitutivity to generate a null expectation for growth rate (black line, logistic fit). Samples coloured red grew significantly faster (1-tailed t-test FDR < 0.05) than this expectation. Each point in (b) and (c) is the mean growth rate observed for a $GAL80$-$GAL4$ pair in a given $GALK$ and $GAL3$ background (N=8 across 4 independent transformations). (d) Distributions of $GAL1$pr-YFP expression for alleles across four genotypic dimensions. Lines and shading reflect within-genotype mean and SD. $GAL1::GALK$ backgrounds bear the $E. coli$ $GALK$ construct. (e) Effects of $GAL1::GALK$ and $\Delta GAL3$ on growth rate across $GAL4$-$GAL80$ pairings. Lines originate and terminate at mean values for the given genotype and grey bars indicate the 95% confidence interval.

**Figure 4. Rewiring of regulatory networks in harmonious genetic combinations.**

(a) Snowflake fitness landscape showing the dependency of $GAL4$-$GAL80$ genotype pairings on $GAL1$ and $GAL3$ sensor activity. The shape of each node indicates expression phenotype and shading the growth rate in galactose. Harmonious combinations (HC) with altered galactose sensing requirements are indicated. While all harmonious combinations require some combination of $GAL1$ or $GAL3$ for growth and expression in galactose, the dependence on $GAL3$ and $GAL1$ varies across every $GAL4$-$GAL80$ background. (b) Rewiring of the GALR network across harmonious combinations. Each network’s inferred wiring is illustrated for a given allelic combination, including Leaky $GAL80.07$ (‘80L’), super-repressor $GAL80S-1$ (‘80S’), the $GAL1$ sensor deletion construct $GAL1::GALK$ (‘K’), and Constitutive $GAL4-L868G$ (‘4C’). 'Inducible' classes include Leaky phenotypes which show low but detectable expression levels in glucose and high expression in galactose.
Supplementary Figure Legends

**Figure S1: Reproducibility of the five key phenotypic values within and between experiments.**
Panels are faceted by phenotype measurements (horizontally) and experiment or experiment comparison. The information above each panel includes the phenotype and the variance explained between the data underlying the two dimensions. The grey lines are where X = Y. a) The mean phenotype of each unique genotype in the dataset ('within-genotype mean'; horizontal axis) is plotted against each individual observation (vertical axis). b) Within-genotype mean values for a given experiment were paired with the within-genotype means of the same genotypes from the other two experiments and are plotted as circles. Error bars are the 95% confidence interval. c) A figure exemplifying the gating strategy based on cell size (FSC) and shape (SSC) features is illustrated. Two filtering steps, a rectangular and next a data-driven filter that selects high-density regions in two dimensions via kernel density estimation (“curve2filter”) were used downstream of the original data (colours). A contour plot is shown over the 10000 data points to illustrate the overall distribution of a typical experiment’s FSC and SSC values.

**Figure S2: The distribution of Gal1p-YFP expression across all single and double mutants from the second pairwise mutant experiment.**
The figure is divided into three sections a-c, each with a GALR locus pairing (GAL80 vs GAL3, GAL3 vs GAL4 and GAL4 vs GAL80, respectively). The genotypes of the GALR allele are indicated in the strip text at the top or right-hand side of each section’s panel sets. Each panel is a unique genotype. The single mutant allele ordering along the facetting dimensions were chosen first by calculating a 'phenotypic index' equal to fraction ON in glucose + fraction ON in galactose as a gross measure of how active the pathway was for a given clone. Following this statistic, roughly, a phenotypic index value of 0 = Uninducible, a value of 1 = Inducible and a value of 2 = Constitutive. Single alleles were then ordered according to the mean within-class phenotypic index value across single alleles within the pairing, followed by the mean within-genotype phenotypic index of single mutants. Finally, the WT and Δ control strains were added to the top of the order in order to help the eye examine the single-mutant effects in a WT background and expectation of clone behavior when the other GALR locus was deleted. Lines and shading are the mean and ± 1 SD of genotypes falling into the given genotype (N=4 for N=2 independent transformations). Blue lines are samples growing in glucose and red lines are samples after 12 hours of growth in galactose.
Figure S3: Distribution of phenotypes across classes, further subdivisions of gene expression distribution classes, and predictability of growth rate based on these classes and a geometric model of single-mutant effects.

All data presented in this figure arises from the second pairwise experiment. a) Transformed expression data used for clustering by the HDBScan* algorithm (Methods). The matrix used for clustering was of dimensions genotype (rows) x 120 A.U. fluorescence bins (cols). Each genotype’s mean log10 density value + 0.001 for each A.U. fluorescence bin across in the glucose and galactose environments were used as values for clustering, and are coloured according to these values in the tile graph. The samples falling to the right and left of the white line delineates the measurements in glucose and galactose, respectively. The ordering of genotypes along the vertical axis was determined by mean within-cluster growth rate in galactose. The log of density values was used to exaggerate small differences for detection by the algorithm, e.g. the weak signal around leaky clones’ expression in glucose. The “single mutant locus” points indicate where single mutants fell along the spectrum of expression profiles (vertical dimension, with horizontal jitter added to differentiate overlapping points), and are coloured according to which locus was mutated. The five classes reported in the text fell in the Intermediate set of classifications of a three-level hierarchy of classes (rightmost coloured lines, Methods). Most of these classifications arose “out of the box” depending on the cluster size setting used with the HDBScan* algorithm, however certain genotypes falling into infrequently observed or intermediate classes were assigned to clusters by hand (Methods). The rightmost three vertical lines are coloured corresponding to final classifications assigned to each clone for Intermediate, Broad and Narrow classifications. Black lines highlight transitions between intermediate classification groups.

b) and c): The distribution of phenotypic values across classes are shown as violin plots for the five key phenotypes, as well as the ‘fold induction’ measure used to sub-classify clones after initial unbiased clustering (Methods). Panel b) is how these characteristics broke down by intermediate classification, and c) is the same data broken down by narrow category. In c) the category labels follow class hierarchies as the broadest: intermediate: narrowest. d) the broadest and narrowest categories are plotted as in Fig 2C across controls, single and double mutants.

(e-g) The predicted vs observed values for prediction of double mutant phenotypes from single-mutant allele phenotypes for classes from the three different hierarchies and (h) a geometric model of growth rate variation (Methods). Datasets are facetted by the locus pairing and data points are coloured according to the Narrow classification scheme. In (h), as explained in the Methods, the low variance explained stems from the fact that pathway-level epistasis leads to Constitutive and Leaky expression profiles (Fig 1D). For example, all pairwise combinations of Constitutive GAL80 and Uninducible GAL3 are Constitutive and therefore grow at high rates, when the prediction is that they will grow slowly due to
GAL3’s low growth rate. Similarly, fast-growing Constitutive GAL4 variants are predicted to grow slowly in \( \Delta \text{GAL3} \) backgrounds, when the double mutant remains Constitutive. Harmonious combinations of Uninducible GAL3 paired with Leaky GAL80 lead to Leaky or Inducible double-mutant phenotypes which grow quickly. Similarly, harmonious combinations of GAL80S-1 and GAL4C permit high rates of growth, when they are predicted to grow slowly due to the dominant repressive single GAL80S-1 backgrounds.

Figure S4: The distribution of \( \text{GAL1pr-YFP} \) expression across all single, double, triple and quadruple mutants from the third higher-order combinatorial mutant experiment.

The genotypes of the GALR allele are indicated in the strip text at the top or right-hand side of each section’s panel sets. Each panel is a unique genotype. Lines and shading are the mean and \( \pm 1 \) SD of genotypes falling into the given genotype (N=4 for N=2 independent transformations). Blue lines are samples growing in glucose and red lines are samples after 12 hours of growth in galactose.

Figure S5: Complete landscape of the effects of \( \text{GAL1::GALK} \) and \( \Delta \text{GAL3} \) on growth rate across \( \text{GAL4-GAL80} \) pairings

Lines originate and terminate at mean values for the given genotype and grey bars indicate the 95% confidence interval.

Figure S6: Pairwise comparisons of GALK backgrounds.

All data presented in this figure arises from the third higher-order genetic interaction dataset. a) All within-genotype mean measurements were scaled (z-scores) within phenotypes. Each of the five phenotypes’ mean scaled within-genotype values are plotted as points and error bars equal to 95% confidence interval. Each axis is a pairwise comparison between \( \text{GAL1pr-GALK} \) ortholog or the \( \text{GAL1pr-HIS5} \) from \( \text{S. pombe} \) constructs, with coloring of points and error bars corresponding to phenotypes. The width of error bars are equal to the 95% confidence interval. b) Phenotypes between \( \text{C. albicans} \) and \( \text{E. coli} \) GALKs were indistinguishable, except that \( \text{C. albicans} \) clones with high growth rates grew more slowly than genotype-matched \( \text{E. coli} \) clones (across individual observation, AD test statistic 5.273, \( p < 0.002 \), rejecting that samples come from the same distribution, whereas \( p > 0.05 \) for other phenotypes). Since \( \text{C. albicans} \) has a systematically lower growth rate, in plots and cartoons in figures 3B, 3D, 3E, and 4 we illustrate our data using backgrounds bearing \( \text{GAL1::GALK} \) from \( \text{E. coli} \).
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Endnotes

Supplementary Information is available in the online version of the paper https://github.com/AaronMNew/HarmoniousCombinations.

Code and data availability statement All code and raw data used to generate the analyses and figures presented in this document can be found at https://github.com/AaronMNew/HarmoniousCombinations

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Author contributions AMN conceived the study, conducted experiments, analyzed data and wrote the manuscript. BL conceived the study and wrote the manuscript.
Methods

Yeast and *E. coli* strains

Primers, strains and specific construction notes for plasmids are in Supplementary Tables 1, and and 2, respectively. All *S. cerevisiae* yeast strains were generated starting from BY4741, (MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0). All cloning in *E. coli* was performed using DH5-alpha or its commercial derivative NEB 10-Beta (New England Biolabs product number C3019I or C3020). Genomic DNA of BY4741, DH5-alpha and *C. albicans* strain SC5314 were used as templates to generate GALR and GALK constructs.

PCR

For high fidelity PCR reactions we followed manufacturer’s instructions using either Extaq (TaKaRa # RR001C), KOD hotstart (Merck Millipore # 71086) or Q5 (NEB # M0491) with 0.6 µM final concentration primers.

For mutagenic PCR, 25 µl reactions of standard Taq (New England Biolabs # M0273) were used with plasmids bearing WT templates of the GAL genes (pAMN14 (GAL3), pAMN15 (GAL80) or pAMN31 (GAL4)). Initial template concentration was varied to allow 8-12 duplications of template.

Components added to mutagenic PCR (order: *Material, volume in µl*): Standard Taq buffer (10X), 2.5; 50mM MgCl2, 2.75; dCTP (100mM), 0.25; dTTP (100mM), 0.25; dATP (100 mM), 0.05; dGTP (100 mM), 0.05; MnCl2 (50mM), 0.25; oligo1 (20 µM), 0.625; oligo2 (20 µM), 0.625; template (5µl per 50 µl reaction), variable; Taq DNA pol. (5U/µl), 0.25; Water, 14.9.

Generation of yeast chassis strains AN612 and AN634

Primers and strains are in Supplementary Tables 1 and 2. AN612, a Gal1p-YFP fusion, was used for the first two experiments, and AN634, a GAL1pr-YFP transcriptional fusion was used for the final experiment. Standard lithium acetate transformation was used in all stages. BY4741 (S288c MATa ΔHIS3 ΔLEU2 ΔMET15 ΔURA3) was used as a starting strain. Starting from BY4741, we first integrated the YeCitrine-KANMX cassette from pKT140 as a fusion protein with GAL1 protein (AN612) or as a disruption of the GAL1 reading frame (AN634). KANR clones were screened at the cytometer, and those that exhibited high FITC-A measurements in galactose and autofluorescent-equivalent levels of FITC-A signal in glucose were saved. For deletion of GALR genes, full loci, including promoter and terminator sequences.
were deleted (the deleted sequences was exactly the same as the sequence of the genes used as complementation cassettes in plasmid complementation constructs). Upon transformation, potential deletion transformant clones were screened by phenotype. For GAL3, clones with GAL80 + GAL4 WT backgrounds were screened by inoculating a 2-day old colony (1 colony = 10^7 cells) into 200 µl in 96-well plates containing YP+2% galactose media and making four 4-fold serial dilutions. After 12-18 hours, those with no expression of the GAL1-YFP construct were selected and saved as the correct deletion strains. For ∆GAL80 strains in the GAL4.WT background, after integration of the deletion cassette, clones were screened for constitutive expression of GAL1-YFP. For deletion of GAL4, clones were screened for lack of GAL1-YFP expression similarly to the GAL3 clones. To make the combinations of allele deletions, clones were generated in the appropriate order to allow phenotypic characterization. For ∆GAL3 GAL80.WT ∆GAL4 strains, after deletion of GAL3 locus, ∆GAL4 integrants were screened for integration of the deletion cassette at the GAL4 locus by traditional PCR screening of 5’ junctions. The final resulting strains, AN612 and AN634 were screened by PCR at the end for proper integrations of all constructs. To complement GALK activity in AN634, PCR products from plasmids pAMN50 (GAL1pr-GALK *E. coli*), pAMN51 (GAL1pr-GALK *C. albicans*), pAMN52 (GAL1pr-GAL1) and pAMN53 (GAL1pr-HIS5 *S. pombe*) were transformed into the GAL4::URA3 locus.

Generation of screening plasmids for mutagenesis

Primers and strains are in Supplementary Tables 1 and 2. Table 3 includes information specific to the PCR products and transformations used. To ensure that all phenotypic variation observed in the PCR mutagenesis experiment were due to the locus targeted by mutagenesis and not the result of mutations in the other two GALR loci (potentially generated in the assembly or PCR process), we generated a screening plasmid with unique cutting sites between the GALR genes. These plasmids were generated by assembling three-GALR plasmids with the targeted GALR flanked by NotI sites and linker sequences. One plasmid was assembled per GALR locus. After in vivo gap-repair assembly of these plasmids, they were prepped and transformed into *E. coli* and screened for correct digestion patterns. These plasmids were named pAMN26, pAMN27 and pAMN28. After identifying correct banding patterns and phenotypic behavior, plasmids were prepared for downstream analysis by removal of the gene of interest. The locus of interest for each one (GAL4, GAL3 and GAL80, respectively) could be liberated with a NotI digest. Recircularization of these plasmids with a 5-minute room-temperature T4 ligase reaction and transformation into *E. coli* yielded plasmids pAMN32, pAMN33 and pAMN34 with deletions of their locus of interest.
Each GALR-specific screening plasmid (pAMN32, pAMN33 and pAMN34) was digested at a concentration of 100 ng / µl overnight in 7 ml in a 15 ml Falcon tube at 37ºC with NEB Cutsmart Buffer + 10 units of NotI-HF (New England Biolabs # R3189) per ml. After at least 12 hours’ incubation, the reaction was brought to 30ºC, and 10 units of Mungbean nuclease added to chew away overhanging ends. After 1 hour incubation at 30ºC, reactions were terminated by extraction of enzyme and other protein with one volume of TE-buffered phenol/chloroform/isoamyl alcohol mixture at pH 8.0 (Sigma Aldrich P-2069). Extracted samples were washed twice with 1 volume of ether to remove excess phenol. Ether was dried off under a flow hood for 30 minutes. DNA was precipitated by addition of 1/10 volume of 3M sodium acetate and 1 volume of isopropanol, frozen at -80 for 20 minutes to overnight, and then centrifuged 20 minutes in a bucket centrifuge at 4ºC. Pelleted DNA was washed twice with 2 ml of room-temperature 70% EtOH (10 minute spins each). EtOH was pipetted off and tubes allowed to dry. DNA was resuspended in 1 ml of TE buffer and purity and concentration checked on a Nanodrop and digest confirmed by running on an agarose gel. These vectors were used downstream as recipient vectors for mutagenic PCRs.

Primers used to generate mutagenic PCR fragments contained blunt-ended NotI scar sites + novel PmeI digest sites + linker sequences, allowing in vivo gap-repair assembly and downstream confirmation of plasmid structure (via PmeI dual cuts + a mid-vector Ndel site). Vectors for the single GALR mutant screen were co-transformed with mutagenized PCR product using high-efficiency transformation into yeast and clones selected and screened as described below.

Generation of mutagenic libraries and targeted mutations

Tables 1, 2 and 3 describe strains and primers. See tables 3B, C and D for information specific to the PCR products and transformations used. Primers for these constructs encode a linker, a short barcode, a NotI cloning site, and template binding sites for amplification. Targeted mutations and random mutants generated in this study used NotI-digested plasmids pAMN32, pAMN33, and pAMN34 to receive one or two fragments of the given GALR gene. Each plasmid generated in these steps were used downstream as templates in the combinatorial genetics experiments.

Generation of combinatorial mutant constructs.

Tables 1, 2 and 3 describe strains and primers. See table 3E and F for information specific to the PCR products and transformations used. All assemblies for combinatorial genetics used four PCR products (the vector + one allele of each of the three GALR loci) and were co-transformed into chassis strain AN612 or AN634. pRS415 (LEU2 marker) was used for combinatorial genetics experiments.
Generation of GAL1pr-GALK constructs.

Tables 1, 2 and 3 describe strains and primers. See table 3G, H and I for information specific to the PCR products and transformations used. Plasmid pAMN45 (pGAL1pr-MET15) was first generated, and GALK orthologs or HIS5 from S. pombe were cloned downstream of the GAL1 promoter. MET15 was used as a marker because all usual markers in the final chassis yeast strain AN634 were used. The final constructs were designed to disrupt the CaURA3 cassette that was used to delete the GAL4 locus. pAMN45 was made by 6-fragment PCR product assembly and was used downstream as a template for parts to assemble the final GALK constructs. Clones bearing plasmids assembled in these transformations were screened for the ability to grow in 0.2% galactose and GAL-inducible growth in SC-HIS + 0.5, 0.1, 0.2, and 2% glucose ± 0.2% galactose media. Correct clones were used downstream to transform strain AN634 with a GAL4::URA3::GAL1pr-GALKxx-Met15 PCR product.

High efficiency gap repair transformation in yeast.

In vivo homologous recombination ("gap repair") was used for generation of all plasmids except pAMN31, which was made by Gibson cloning. For gap repair, yeast clones were generated using combined with a high efficiency yeast transformation adapted from on a protocol described in 23. Appropriate yeast strain backgrounds were streaked from the freezer and incubated 2-3 days at 30C. A patch of cells were picked from the center of the streaked cells to prevent genetic bottlenecks of single clones. The cells were inoculated into 5-50 ml of YPD and grown 12-16 hours until saturation with vigorous shaking at 30ºC. These overnight cultures were then inoculated into 30ºC YPD media to a 1 cm OD600 absorbance of 0.3, approximately 2*10^7 haploid BY4741 cells per ml. Cells were incubated in Pyrex glass bottles with vigorous shaking every 30 minutes to 1 hour. After 4 hours, cultures were aliquoted, appropriate to the scale, to 50 ml falcon tubes or 250 ml centrifuge bottles and centrifuged at 1250xg for the a the time required to pellet the cells (approximately 2 minutes for 50 ml falcon tubes and 5 minutes for the 250 ml centrifuge tubes). Cells were resuspended in 10-50 ml of SLAT buffer, and centrifuged again in 50 ml falcon tubes for 5 minutes at 1250xg, enough to pellet most cells.

Thereafter, per 50 ml of culture: cells were resuspended in SLAT buffer by agitating the cell pellet with a long pipette, followed by vigorous shaking. The volume of SLAT was adjusted depending on the volume of DNA transformation product being added such that a final volume of SLAT was 2.5 ml per 50 ml original culture. The protocol differed hereafter for library vs. targeted GALR allele clones.
For library generation, DNA was directly added to cells, including 50 µl of recently boiled-then-snap-chilled 10 mg/ml ssDNA (salmon sperm DNA Agilent # 201190), 1 mg vector, and 1 mg each PCR product for in vivo gap repair assembly. The mutagenized PCR product was added directly to the cells. Cells+SLAT+DNA mixtures were left with occasional agitation at room temperature for at least 30 minutes. Thereafter, 10 ml of PLATE mixture was added to cells, and cell-SLAT-DNA-PLATE mixtures were shaken and left to rest at least 30 minutes at room temperature. Then DMSO was added to PLATE + cell mixtures to 8% final concentration, and they were heatshocked in a water bath for 20 minutes. Cells were then centrifuged 5 minutes at 1250xg, resuspended in 0.5 M sorbitol + YPD media and allowed to recover 1 hour at 30ºC. After recovery, cells were centrifuged again at 1250xg, and pellets resuspended in selection media and either recovered in 500 ml of liquid (200 RPM shaking for 48 hours) or spread onto solid plates.

For high-throughput gap repair assemblies with specific combinations of GALR alleles, cell mixtures were resuspended in 1.375 ml SLAT, 50 µl of recently boiled-then-snap-chilled 10 mg/ml ssDNA and 1 mg PCR vector per 50 ml original culture. Then 11 µl of cells+SLAT were aliquoted to 9 µl of DNA-SLAT mixtures pre-aliquoted to 96-well polyethylene PCR plates (Thermo Scientific AB-0700). Each gap repair assembly mix had 0.1 µl of each of the three GALR’s PCR product + 8.7µl SLAT. No shaking or agitation/mixing was used to blend the cells with DNA. After adding cells to 96-well plates, 80 µl of PLATE was added and plates sealed carefully using Biorad Microseal B seals. Samples were shaken vigorously by inversion to mix cells with the PLATE buffer. Cells were centrifuged in a swinging bucket centrifuge at 300 rpm for 3 seconds and then left at room temperature for at least 30 minutes and up to 3 hours. After incubation, plate seals were removed and 8 µl of DMSO was added to cell + PLATE mixtures. Seals were reapplied to the 96-well plates and the samples shaken vigorously by inversion, followed by centrifugation in a swinging bucket centrifuge at 300 rpm for 3 seconds (to bring samples down to the tube bottoms), followed by heat shock at 42ºC for 20 minutes in a pre-warmed PCR block.

After heat shock, cells were centrifuged for 1 minute at 1250xg, PLATE+SLAT mixture dumped out, and cells resuspended in 80 µl YPD + 0.5 M sorbitol and incubated for one hour. Plates were then spun down for 3 minutes at 1205 x g to pellet cells, and cells were resuspended in 500 µl of SC-LEU+2% glucose + ampicillin 100 mg/L selection medium in 96-well plates. Plates were sealed with Biorad Microseal B seals or unsealed and covered as a stack of plates with the same plastic sheaths in which the plates were shipped. Efficiency of transformation of each well could be assessed after 36-48 hours growth by counting colonies on the bottom of the wells. Transformations with fewer than 20 colonies could be accurately counted. Although most wells had > 20 colonies, those with < 10 single colonies were excluded from analysis downstream. Cells were then resuspended and 3 µl inoculated into 75 µl of
selection media and allowed to grow overnight. Cells were resuspended with 75 µl of 50% glycerol and frozen at -80°C until experiment measurement.

Pre Growth of clones before flow cytometry

To begin flow cytometry analysis, plates with combinatorial genetic assemblies were taken from freezer and in total sat at room temperature for 20-25 minutes. Plates were put on orbital shakers once thawed and shaken 1-5 minutes before inoculation of 10 µl into 190 µl (200 µl final volume) of SC-[LEU or HIS] +0.1% glucose + 100 mg/L ampicillin a+ 20 mg / L chloramphenicol and grown without agitation in stacks encapsulated in the plastic sheaths in which the plates were shipped (Sarstedt 82.1581) for 12-24 hours to saturation. Cells were resuspended on the plate shaker and diluted 1/50 into 75 µl SC-[LEU or HIS] +0.1% glucose (no antibiotics here) grown 18-24 hours to saturation in stacks of unsealed plates encapsulated in the plastic sheaths in which the plates were shipped in preparation for inoculation to galactose and measurement at the cytometer the next day.

Inoculation of clones to galactose, measurement of cell densities and GAL-YFP gene expression in glucose and galactose environments by flow cytometry

Plates containing 75 µl 0.1% glucose-grown cultures (either from the single clones picked in the mutagenesis experiment or the clones generated to have targeted allele combinations) were grown 18-24 hours unsealed in stacks of plates encapsulated in the plastic sheaths in which the plates were shipped (Sarstedt 82.1581). After growth, samples were placed on an orbital shaker for 1-5 minutes to resuspend cells, then 150 µl of ddH2O was added to the cells to make a 3-fold dilution of the original cell density, with continued shaking for another 1-5 minutes. 9 µl of the ddH2O-diluted cultures was added to 4ºC plates containing 141 µl of 1.06x concentrated SC-[LEU or HIS] + 0.2% galactose media. Inoculated galactose plates were sealed with Microseal B seals and placed immediately at 4ºC to prevent growth or gene expression prior to beginning of growth experiment. We found that only turbid cultures could be resuspended by shaking on the 2.5 mm-radius orbital shaker. Therefore, while inoculating into galactose, we took care to distribute the cells evenly across the whole well. Plates were then sealed with Microseal B seals and put at 4ºC. At the end of the day all glucose-pregrown cultures that had been inoculated into galactose media were placed at 30ºC in stacks of 1-2 plates to begin growth.

After inoculation of galactose plates with glucose-grown cells, we put the glucose plates at 4ºC until measurement at the flow cytometer (BD FACS Canto; FACS Diva v 5.0.3 Firmware V 1.4). Prior to measurement at the cytometer, plates were put back on the shaker for 2 hours. Plates were visually
inspected to be sure that the cells in all wells were well-suspended. We measured cell density and gene expression (bandpass filters “FITC-A” 530±15 nm and “PE” 585±21 nm were used for YFP signal and 488±5 nm for SSC signal). High-throughput sampling mode was used with no mixing. The median time to complete a plate was 18 minutes. During this time we determined that cell density measurements did not appreciably change. If any problem was encountered during the cytometry and the measurements needed to be stopped, we took the plate out and put back on the plate shaker briefly to resuspend the cells before resuming the cytometry.

After 12 hours of growth at 30°C in SC-[LEU or HIS]+0.2% galactose, samples were placed on ice or on a cold surface in a 4°C room to arrest growth and allowed to cool at least 30 minutes prior to exposure back at room temperature. Prior to measurement at the cytometer, Microseal B covers were removed and samples put on the orbital shaker for 2 hours covered by a breathable plate seal. As mentioned above, samples that did not grow appreciably could not be easily resuspended by the orbital shaker. Therefore, prior to measurement, all cultures were pipetted up and down 5 times with a multichannel micropipette, and then placed immediately in the FACSCanto for analysis. Prior to sampling in the cytometer, wells were scored by eye for high growth or low growth. The cytometer template’s sampling rates were adjusted according to these by-eye scores: high density samples were sampled at 0.5 µl per second, while low density cultures were sampled at 2.0-3.0 µl per second, with occasional intermediate sampling rates for obviously intermediate cell densities. Each sample’s sampling rate can be found in all supplementary tables where we report data for these experiments.

Selection and isolation of plasmids from single yeast clones for combinatorial genetics experiment.

After screening and phenotypically characterizing mutagenized GALR variants by flow cytometry (see above) we isolated plasmids of interesting phenotypes. Clones were selected to reflect either outlying phenotypes or more typical behaviour. “Outlying phenotypes” included samples where both fracon.glu and fracon.gal measurements were > 0 and less than 1, indicating that the clones had a constitutive character but could not fully induce the GAL pathway. Another rare phenotype we tried to isolate were clones where mean signal in ON cells was less than mean signal of typical inducible ON clones (as discussed in the text this phenotype was quite rare). Although some GAL3 mutants appeared to have constitutive characters in the first screen, we found that none of these phenotypes were recapitulated upon subcloning.

After selection based on phenotype, clones were thawed from the freezer and struck to single colonies. These were inoculated into SC-HIS+2%glucose media in PCR plates and gDNA prepped in 96-well plates as described above. Clones were transformed into electrocompetent 10-Beta cells (New England
Biolabs # C3020) using a 96-well plate electroporator using fresh electroporation plates (BTX 45-0450-M). Cells were recovered in deepwell plates (Thermo Scientific # 260252) and recovered in 0.6 ml SOC media for 1 hour prior to inoculation into 0.6 ml LB+100 mg / ml ampicillin. The next day plasmids were prepped in 96-well plates as described above. Preps were digested with Ndel and Pmel enzymes, which yielded 3 bands in correctly assembled constructs. Plasmids with the correct banding patterns were used downstream in the combinatorial genetics experiment.

Data availability

Scripts and processed data are available at the github link: https://github.com/AaronMNew/HarmoniousCombinations.

Analysis of the raw flow cytometry data.

R was used for all analyses. Scripts and data are available at the github link: https://github.com/AaronMNew/HarmoniousCombinations. FCS3 files were exported from the computer controlling the FACSCanto measurements and sampling rate information extracted from exported .xml files generated from export of “Experiment Template”. Scripts for extracting metadata from these .xml files are found in the supplementary code. Experiment “layout” files were generated including clone information, known genotype information and censorship information (censored either if they had very low transformation efficiency or a contamination), and this was merged with metadata of sampling rates in the .xml file. As a basic overview of the analysis, the Bioconductor FlowCore package tools were used to open the binary FCS files and filter first based on cell shape and size information using first a rectangle including 95% of observations in side scatter (SSC) and forward scatter (FSC), then a centroid algorithm was used to identify the most dense observations in these two dimensions, excluding between 30-50% of outlying original observations. FITC-A signal was used to quantify YFP expression. The predicted FITC-A value of a PE-A reading was predicted by a linear model \(\text{lm}(\log(\text{FITC-A}) - \log(\text{PE}))\), and these predicted values were used in the rare cases where a cell’s FITC-A signal exceeded the machine’s maximal measurement value. Then key parameters of FITC-A distributions were extracted, including the mean YFP signal, fraction ON (the proportion of cells falling above an empirically determined cutoff based on autofluorescent cell controls). Because many lowly expressing cells gave negative values at the flow cytometer, we calculated a pseudo-log10 FITC-A measurement as the log10( raw FITC-A measurement + 1000 )-3. These pseudo-log10 fluorescence intensity values were broken into 60 bins using \text{cut()}\). Cells were counted in the glucose and galactose environments by determining the slope term in of events / second information of each FCS file using a linear model \(\text{lm}(\text{events} - \text{ms})\). Glucose-grown biological replicate clones were then matched with their next-day galactose measurements. We calculated cell densities by multiplying the events/second measurement by the known sampling rates.
extracted from the .xml file. The density of the culture in galactose was calculated as this measured cell density parameter multiplied by the inverse of the dilution factor from glucose the day before (150/9). Log2 change of the culture $\text{Log}_2(\text{Density.GAL grown.culture} / \text{Density.GLU grown.Culture})$ was scored as the number of generations, and the growth rate parameter $\mu$ was calculated as $\ln(\text{Density.GAL grown.culture} / \text{Density.GLU grown.Culture}) / 12$ hours.

Plotting and basic analysis of data

All scripts are online at github https://github.com/AaronMNew/HarmoniousCombinations. Custom functions, data.table and ggplot2 packages were used for summarizing data and plotting. Minor aesthetic changes were made in Adobe Illustrator. The final two panels of figure 4 were made in Adobe Illustrator.

Gene expression distribution clustering.

For gene expression distribution clustering, we used the HDBScan* function (R package dbscan)\textsuperscript{24}. This algorithm establishes hierarchical clusters based upon distance-weighted graphs assembled according to the density of data points across n-dimensions. Its final cluster assignments are mainly sensitive to the “minPts” parameter, which sets the minimum cluster size. This sensitivity arises primarily from a 1) the size of the dataset (because more points for a large dataset will yield the same number of clusters as for a small dataset) and 2) how distantly spread the clusters of data points are in n-dimensional space.

For gene expression distribution clustering we wanted to cluster based on expression in both glucose and galactose. For this we first paired the densities of expression across 60 bins of pseudo-log10-transformed A.U. FITC-A signal for each sample at time 0 (glucose expression) and after 12 hours of growth in galactose (galactose expression) to generate vectors of 120 units for each observation. The mean vector for each unique genotype was then calculated. These mean values were then pseudo-log transformed to exaggerate signal within bins exhibiting low density values, for example such that a small fraction of cells active in glucose would be more salient to the HDBScan* algorithm (Fig S3A). For this, the density values for each bin transformed as follows:

$$V = \log(\text{round}(v,3)+0.001)$$

Where $V$ is the final pseudo-log density value and $v$ is the original density value. A matrix was made comprised of rows corresponding to each genotype’s vector of measurements of $V$, with pseudo-log10
A.U. expression distribution bins as columns. As a final step, each bin’s $V$ across genotypes was scaled by z-score. The scaled 120 dimension matrix was then clustered using HDBScan* algorithm.

For experiment 1, with three clearly defined clusters, a minPts parameter was chosen such that 3 clusters emerged. For experiment 2, which exhibited a larger variation in expression distributions between genotypes, the minPts parameter was tuned to such that 7 clusters, plus a number of unclustered samples, emerged. These clusters were chosen a posteriori based on their ability to predict total phenotypic variance, a parameter that was determined using least-squares regression of the form

$$lm \ (within\_genotype\_mean\_value \sim cluster \ast phenotype\_id)$$

Where phenotype_id was one of the five phenotypes including fraction ON in galactose, fraction ON in glucose, mean YFP expression in galactose, mean YFP expression in glucose, and the growth rate parameter $\mu$.

Across minPts parameters, two levels of clusters were evident: the broadest category included the three main inducible, uninducible and constitutive clusters and explained 84% of total phenotypic variance across all phenotypes. A more narrow set of classes comprised nine categories, explaining 97% of total phenotypic variance. Clusters were then manually curated to five intermediate “constitutive”, “uninducible”, “inducible”, “leaky” and “weak expression” categories, which together explained 96% of total phenotypic variance. We chose the intermediate classifications for discussion in the main text for prediction of double mutant phenotypes from single-mutant phenotypes due to their power to explain total phenotypic variance in the dataset compared to the broad categories and limited number of parameters compared to the more narrowly-defined categories (see Supplemental Analysis Code).

After this first round of clustering, 3-4% of samples remained unclassified. A closer look at the these samples showed that most expression distributions were “flavors” of the other phenotypes, however due to slight differences and their infrequent numbers, they remained unclustered. For example, the three GAL4 clones in the “weak expression” category all showed low max gene expression level, however otherwise behaved WT, for example exhibiting constitutive characters in combination with $\Delta$GAL80. Repeating the clustering with unclustered samples plus these less frequently observed weak expression clones showed that most unclustered samples showed characteristics similar to these GAL4 alleles, and indeed that they mostly included one of these three GAL4 single-mutant backgrounds. Using a similar approach, we took the remaining unclustered samples and repeated their clustering alone with samples from the three archetypal samples identified in the first experiment, leaving all previously uncategorized
expression profiles with a classification. Finally, visual inspection revealed that some clones with constitutive characters exhibited a nonetheless high degree of activation in galactose relative to glucose. To quantify this we took the mean YFP signal in galactose / mean YFP signal in glucose as a measure of a clone’s induction. Using this parameter, we could identify clones mis-classified as constitutive to be “inducible” or “leaky”. Similarly, some “inducible” clones displayed low mean induction values, so were classified as uninducible, and some “uninducible” clones actually showed >40% of cells ON in galactose with high a mode of expression and so were classified as inducible.

Predictive modeling of double mutant growth rates based single mutant gene expression clusters.

The expression clusters of each single mutant was matched to each double mutant. Mean growth rate values for each unique combination of single-mutant expression clusters were then taken as an expectation of the double-mutant’s growth rate. These 23 unique numbers were used to calculate the fraction of variance explained across all individual observations, and within-genotype mean growth rate values were calculated and plotted against predictions for the second figure. Note that these 23 unique combinations do not include all 32 possible combinations of 4 * 4 * 2 GAL4, GAL80, and GAL3 classes. This is because certain triple mutant classes were not measured. Specifically "uninducible" GAL3 was never paired with the three classes of GAL80 and GAL4 that do not include "inducible". These classes number 3 X 3 X 1 GAL4, GAL80 and GAL3 alleles = 9, so 32-9 = 23 unique clusters arising from single-mutant combinations.

Notably, these missing classes are all almost included in the final experiment, including constitutive GAL80, leaky GAL80, uninducible GAL80, uninducible GAL4, constitutive GAL4. The only one not included are the weak expression class of GAL4, three alleles of which all behaved very similarly to WT GAL4 (e.g., uninducible with uninducible GAL80S variants or uninducible GAL3 variants, and constitutive with constitutive GAL80 variants) but with lower peak expression level.

Predictive modeling of double mutant growth rates based on single mutant mutation effects.

For calculating expected growth rate based on single mutant effects, we used a first-order geometric model of phenotypic variation where the expected multiple mutant locus (\( \mu_{MUT} \)) phenotype is the product of the phenotypes of each (single) allele across N loci in the WT background normalized by the WT reference phenotype value:

\[
\mu_{MUT} = \frac{\mu_{mut1} \cdot \mu_{mut2} \cdot \mu_{mut3} \cdot \ldots \cdot \mu_{mutN}}{\mu_{WT}^{N-1}}
\]
Standard errors for these predictions were propagated as

\[ se_{\text{mut}} = \text{abs}(\mu_{\text{WT}}) \times \sqrt{((N-1) \times (se_{\text{WT}}/\mu_{\text{WT}})^2 + se_{\text{mut1}}/\mu_{\text{mut1}} + se_{\text{mut2}}/\mu_{\text{mut2}} + \ldots + se_{\text{mutN}}/\mu_{\text{mutN}})^2)} \]

Phenotype values used for \( \mu \) were equal to growth rate minus the background growth rate observed in the absence of any GAL regulator. These expectation values were used to predict overall variance in the dataset, explaining 52%, 20%, and 85% of variance for pairings between GAL3 vs. GAL4, GAL80 vs GAL3, and GAL4 vs GAL80, respectively. Overall this model explained 55% of variance across the dataset. These low values stem from the fact that pathway-level epistasis leads to constitutive and leaky expression profiles, which dominate signal in the dataset (Fig S3H). For example, all pairwise combinations of constitutive GAL80 and uninducible GAL3 are constitutive and therefore grow at high rates, when the prediction prediction is that they will grow slowly due to GAL3's low growth rate. Similarly, fast-growing constitutive GAL4 variants are predicted to grow slowly in \( \Delta \text{GAL3} \) backgrounds, when the double mutant remains constitutive. Harmonious combinations of uninducible GAL3 paired with leaky GAL80 lead to leaky or inducible double-mutant phenotypes which grow quickly. Similarly, harmonious combinations of GAL80S-1 and GAL4C permit high rates of growth, when they are predicted to grow slowly due to the dominant repressive single GAL80S-1 backgrounds.

Fitting a logistic model to the relationship between GAL1-YFP expression and growth rate

To demonstrate that GALK orthologs from \textit{E. coli} and \textit{C. albicans} did not display signaling activity, we compared expression of the GAL1pr-YFP fusion to growth rates in which all sensors were deleted and found a sigmoidal relationship between mean YFP expression in glucose. We used this latter expression in glucose a measure of “pathway leakiness” or constitutivity to predict the expected growth rate of the culture in galactose. For this, we fitted a logistic curve using R’s \texttt{SSlogis()} function of growth rate as a function of initial gene expression level to demonstrate that inducible or leaky “harmonious combination” mutant backgrounds are able to mount an induction response and high growth rate from an initially OFF state. A one-sided t-test was performed for each within-genotype mean across all backgrounds compared to this null expectation. To account for the differential growth rates observed for \textit{C. albicans} vs. \textit{E. coli} GALKs, we fitted separate curves for calculation of t-test statistics, while the main figure simply shows the logistic curve across all \textit{E. coli} and \textit{C. albicans} backgrounds. See supplemental code for details.
Standard lithium acetate yeast transformation

PLI (Polyethylene glycol 3530 + Lithium acetate + 1x TE buffer) is 50% polyethylene glycol + 0.1 M LiAc in TE Buffer and was prepared ahead. Cells were inoculated 200 µl into 25 ml YPD and grown in Falcon tubes with occasional shaking at 30ºC for 4-5 hours. Cells were pelleted by centrifugation on high in the Falcon tube in which they were grown, washed once with 700 µl 0.1 M LiAc, spun again quickly in the Eppendorf tube to pellet cells, and resuspended 200 µl of 0.1 M LiAc. 10 µl of boiled ssDNA (1 mg / ml) + 25 µl of PCR product was added and samples mixed by flicking. Tubes were optionally incubated at room temperature for up to 30 minutes. 600 µl of PLI was then added to cells and mixed well by vortexing. Tubes were optionally incubated at room temperature for up to 30 minutes. Tubes were then incubated at 42°C for 30 minutes. Cells were pelleted at 1250xg for 3-5 minutes and then resuspended either directly in selection media (for auxotrophic markers) or in 0.5-5 ml YPD media (for dominant drug resistance markers). YPD samples were incubated 3-4 hours at 30ºC. Samples were then spread on solid selective media (1-5 plates depending on expected transformation efficiency). Clones were visible after 2 days and were struck to single colonies on selective media and grown 48 more hours. Clones that grew in the patches were picked for downstream phenotypic screening and genotypically and correct clones were frozen from single colonies.

Small-scale genomic DNA (gDNA) preps

Yeast genomic DNA was isolated by alkaline lysis and isopropanol precipitation using a scaled-down protocol based on that provided by MasterPure™ Yeast DNA Purification kit (#MPY80200). 150 µl of yeast cells were grown overnight in appropriate selective media in 96-well PCR plates covered with breathable seals with no shaking. The next day they were spun for one minute in a swinging centrifuge centrifuge on high (~3200 x g or 4000 RPM on a swinging bucket centrifuge). Spent media was shaken out and immediately after dumping spent media, plates were swabbed on ethanol-soaked paper towel to remove most of the media still clinging to the sides of the plate. Cells were resuspended in 50 µl lysis buffer by pipetting or inversion while covered with Biorad Microseal B seals (catalog number fMSB1001) and inverted several times to mix. Plates were spun briefly for 1 second at 300 RPM to get the lysed cells back into the wells and off of the plate sealer. Samples were incubated at 65ºC for 15 minutes for lysis, and then plates placed on ice for 5 minutes. Plate seals were removed and 25 ul of MPC precipitation buffer was added to the lysed cells. Plates were re-sealed and inverted multiple times to be sure protein and other cellular debris was precipitated. Debris was pelleted by centrifugation on high (~3200 x g, 4000 RPM) in a bucket centrifuge for 10 minutes. Then plate seals were removed and 50 µl of gDNA-containing supernatant was transferred into 50 µl of isopropanol in a new 96-well plate. Plates were re-sealed and inverted several times to mix the DNA and isopropanol.
Samples were then centrifuged on high (~3200 x g, 4000 RPM) in a bucket centrifuge for 10 minutes. After centrifugation, isopropanol was dumped off and while still upside down, the plates gently dabbed on paper towels to absorb more isopropanol clinging to the plate. 60 µl of 70% ethanol was then added, plates resealed and inverted gently one time to mix the remaining isopropanol and ethanol together. Plates were then centrifuged on high (~3200 x g, 4000 RPM) in a bucket centrifuge for 2-5 minutes. An optional second 70% ethanol wash was sometimes performed. Then ethanol was dumped and while still upside down put the plate on a paper towel to absorb ethanol clinging to the plate. Plates were then spun briefly to bring remaining ethanol to the bottom of the wells, and using a Rainin multichannel P10 with LTS tips (very fine tips) the remaining ~4-10 µl of ethanol was removed from the plates. Plates were allowed to dry 10 minutes. DNA was resuspended in 25-50 µl EB buffer, mq H2O or TE buffer (depending on downstream use). Quality of prep was confirmed by measuring purity and estimation of DNA concentration on a Nanodrop and running an agarose gel of 3 µl of 12 preps randomly selected across the plate.

Small-scale plasmid mini preps

To isolate plasmids, we used a simple alkaline lysis miniprep protocol using buffers P1, P2 and P3 from Qiagen (catalog numbers 19051, 19052 and 19053 respectively), either at a “normal” scale, with 1.5 ml of saturated bacterial culture yielding > 10 mg of DNA or scaled-down in a 96-well plate with yields of >1 mg plasmid DNA per sample.

For normal mini-preps we first picked single colonies from a selective plate or inoculated a stab of cells directly from the -80 freezer stock into at least 2 ml of liquid LB containing the selective antibiotic and incubated overnight with vigorous shaking. After incubation, 1.5 ml of cells were pelleted at 13000 rpm for 1 min. Cells were resuspended in 150 µl buffer P1 + RNAse, lysed for 1-5 minutes in 150 µl buffer P2, and then cellular debris precipitated with 150 µl buffer P3. Debris was then pelleted at 13000 RPM for 10 minutes. Plasmid DNA was precipitated by adding 1 volume of isopropanol, the tubes inverted a few times to mix well, and then tubes were centrifuged at 13000 RPM for 10 minutes at 4°C. Isopropanol was dumped from the tubes and 0.5 ml of room-temperature 70% ethanol was added, the tubes gently inverted one time, and spun at 13000 RPM for 2-5 minutes. Sometimes the pellets were washed again with ethanol. Ethanol was dumped, residual ethanol removed by a quick spin and pipetting, and the tubes left to dry for 15 minutes. DNA was resuspended in 1X TE buffer or EB buffer from Qiagen (catalog number 19086).

For 96-well plate mini-preps, a scaled-down version of the protocol used above was followed. Single colonies or pools of transformants from yeast clones were inoculated into LB+selection in a deep-well
96-well plate (Thermo Scientific 260252) and the plates sealed using breathable plate seals (Thermo Scientific # AB-0718) and incubated overnight with vigorous shaking on a 2.5 mm radius orbital plate shaker overnight at 37ºC. After growth, 150-160 ul of turbid cells were pipetted into 96-well PCR plate and centrifuged on high (~3200 x g or 4000 RPM on a swinging bucket centrifuge) for 5 minutes. Spent media was shaken out into an autoclavable or bleachable container, and immediately after dumping spent media, plates were dabbed onto an ethanol-soaked paper towel to remove most of the media still clinging to the sides of the plate. Cells were resuspended in 25 µl P1 buffer + RNAse by pipetting up and down, then 25 µl buffer P2 was added to cells. Plates were sealed with Microseal B seals and inverted several times to mix, then left to let sit for 5 minutes. Plates were spun briefly for 1 second at 300 RPM to get the lysed cells back into the wells and off of the plate sealer. Plate seals were removed and 25 ul of buffer P3 was added to the lysed cells and plates were re-sealed and inverted multiple times to be sure protein and other cellular debris was precipitated.

Debris was pelleted by centrifugation on high (~3200 x g, 4000 RPM) in a bucket centrifuge for 10 minutes. Then plate seals were removed and 60 µl of plasmid-containing supernatant was transferred into 60 µl of isopropanol in a new 96-well plate. Plates were re-sealed and inverted several times to mix the DNA and isopropanol. Samples were then centrifuged on high (~3200 x g, 4000 RPM) in a bucket centrifuge for 10 minutes. After centrifugation, isopropanol was dumped off and while still upside down, the plates dabbed on paper towels to absorb more isopropanol clinging to the plate. 60 µl of 70% ethanol was then added, plates resealed and inverted gently one time to mix the remaining isopropanol and ethanol together. Plates were then centrifuged on high (~3200 x g, 4000 RPM) in a bucket centrifuge for 5 minutes. An optional second wash was sometimes performed. Then ethanol was dumped and while still upside down put the plate on a paper towel to absorb of yet more ethanol clinging to the plate. Plates were then spun briefly to bring remaining ethanol to the bottom of the wells, and using a Rainin multichannel P10 with LTS tips (catalog number 1700587; very fine and flexible tips) the remaining ethanol was removed from the plates. Plates were allowed to dry 10 minutes. DNA was resuspended in 25-50 µl EB buffer, mq H2O or TE buffer (depending on downstream use). Quality of prep was confirmed by measuring purity and DNA concentration at a Nanodrop and running an appropriate digest on an 0.8% agarose TAE-buffered gel.

Methods References

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**Experiment 1:**

Factorially complete WT and ∆GALR mutants

**Combinatorial genetics:**

3-locus gap assembly into ∆ chassis strain

**Growth**

Endpoint average growth rate \( \mu \) measured from cell density

**Gal1p-YFP reporter**

Single-cell distributions via flow cytometry

**Chassis genome:** ∆G3, ∆G4, ∆G80

**Gal1p-YFP**

Single-cell distributions via flow cytometry

**Figure 1**

**GAL regulatory (GALR) genes and pathway measurements**

- Canonical GALRs
- Measured phenotypes

**Galactose intracellular**

- Galactose sensor
- Repressor
- Trans activator

**Transcriptional activation**

- Inhibitory physical interaction
- Stimulatory physical interaction

**Uninducible**

- All ∆GAL4 backgrounds or ∆GAL3 single
- WT GALRs on plasmid or WT genome control

**Inducible**

- ∆GAL80 single or ∆GAL80 + ∆GAL3

**Constitutive**

- PT GAL3 locus
- GAL4 locus

**GAL3**

- WT
- ∆

**GAL4**

- WT
- ∆

**GAL80**

- WT
- ∆

**Number of mutations**

- 0 – WT genes on plasmid
- 1 – Single mutants
- 2 – Double mutants
- 3 – Triple mutant

**Figure 1**

**A**

- Uninducible
- Inducible
- Constitutive

**D**

- Uninducible
- Inducible
- Constitutive

**E**

- Fraction ON in glucose
- Fraction ON in galactose

**F**

- Growth rate \( \mu \) hr\(^{-1}\) in galactose
- R\(^2\) = 0.44
- R\(^2\) = 0.96

**G**

- Growth

**H**

- Endpoint average growth rate \( \mu \) measured from cell density

**I**

- Galactose intracellular
- Galactose sensor
- Repressor
- Trans activator
Figure 2

Experiment 2: 5,319 pairwise combinations of GALR alleles

- Single mutants
- Double mutants

**C**

| Single mutants | Double mutants |
|----------------|----------------|
| Control        | GAL3          |
|                | GAL80         |
|                | GAL4          |
| GAL3 GAL4      | GAL3 GAL80   |
| GAL3 GAL80    | GAL4 GAL80   |
| GAL4 GAL80    |               |

**D**

36 possible combinations - 9 unmeasured triple-mutant combinations = 23 combos total

**E**

- Inducible
- Uninducible
- Constitutive
- Leaky
- Weak Expr

**GAL4** 43 alleles
**GAL3** 46 alleles
**GAL80** 39 alleles

**GAL3 GAL4**

- Variety of expression classes based on single-mutant genotypes

- Predicted $\mu$ hr$^{-1}$ in galactose based on single-mutant expression class membership

**Figure 2**
Figure 3

A

Mother GALK (galactokinase)

Bifunctional GALK

S. cer. GAL3

Sensor ++++

No GALK activity

S. cer. GAL1

Sensor +

C. albicans GALK

Sensor −

E. coli GALK

Sensor −

GAL4 alleles

GAL80 alleles

GAL4-WT

WT GALK

Inducible

Growth in gal

Baseline pathway activity

Expression in glu

Each point is a unique GAL80 + GAL4 combination

GAL1 sensing required

** GAL3 non-essential

** GAL1 sensing required even with GAL3

D

WT GAL80

GALB0 #07

GAL80S-1

WT GAL3

ΔGAL3

E. coli

WT

ΔGAL3

WT

ΔGAL3

WT

ΔGAL3

WT

ΔGAL3

WT

ΔGAL3

WT

ΔGAL3

mean GAL1pr-YFP expression in glucose, A.U.

B

mean µ hr^-1 in galactose

GAL1 pr-YFP fluorescence intensity, A.U.

pseudo−log10

sugar
gal

E

Number of mutations

GAL sensor genetic changes

ΔGAL3

ΔGAL3

ΔGAL3 + GAL1::GALK E. coli

95% CI WT background

95% CI ΔGAL4 or GAL1::HIS5
Figure 4

A

Growth rate in galactose
- High
- None

Expression class
- Constitutive
- Inducible
- Uninducible

Mutations added
- GAL80S-1
- GAL80.07
- GAL4-L868G
- ∆GAL3
- GAL1::GALK

B

Expression class
- Inducible
- Constitutive
- Uninducible

Genetic interactions
- Inhibitory
- Stimulatory
- weak
- strong

Mutations added
- GAL80S-1
- GAL80.07
- GAL4-L868G
- ∆GAL3
- GAL1::GALK