

On the illusion of auxotrophy: *met15*∆ yeast cells can grow on inorganic sulfur, thanks to the previously uncharacterized homocysteine synthase YII058w

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Organisms must either synthesize or assimilate essential organic compounds to survive. The homocysteine synthase Met15 has been considered essential for inorganic sulfur assimilation in yeast since its discovery in the 1970s. As a result, MET15 has served as a genetic marker for hundreds of experiments that play a foundational role in eukaryote genetics and systems biology. Nevertheless, we demonstrate here through structural and evolutionary modeling, in vitro kinetic assays, and genetic complementation, that an alternative homocysteine synthase encoded by the previously uncharacterized gene YLL058W enables cells lacking Met15 to assimilate enough inorganic sulfur for survival and proliferation. These cells however fail to grow in patches or liquid cultures unless provided with exogenous methionine or other organosulfurs. We show that this growth failure, which has historically justified the status of MET15 as a classic auxotrophic marker, is largely explained by toxic accumulation of the gas hydrogen sulfide because of a metabolic bottleneck. When patched or cultured with a hydrogen sulfide chelator, and when propagated as colony grids, cells without Met15 assimilate inorganic sulfur and grow, and cells with Met15 achieve even higher yields. Thus, Met15 is not essential for inorganic sulfur assimilation in yeast. Instead, MET15 is the first example of a yeast gene whose loss conditionally prevents growth in a manner that depends on local gas exchange. Our results have broad implications for investigations of sulfur metabolism, including studies of stress response, methionine restriction, and aging. More generally, our findings illustrate how unappreciated experimental variables can obfuscate biological discovery.

The metabolism of organic sulfur-containing compounds ("organosulfurs") is critical for all domains of life. Plants and microbes, unlike animals, often synthesize organosulfurs from inorganic sulfates via the sulfate assimilation pathway (SAP) (1). Enzymes in the SAP reduce inorganic sulfates into sulfides and ultimately into organosulfurs. In yeast, the terminal step of the SAP generates homocysteine from O-acetyl-L-homoserine (OAH) and hydrogen sulfide (H_2S) . Homocysteine can then be converted either directly to methionine (Met) or indirectly to cysteine (Cys). These amino acids may be incorporated into nascent polypeptides; alternatively, they may be modified to produce other essential organosulfurs, including the "universal methyl donor" SAM and the important buffer against oxidative stress glutathione. Reactions downstream of homocysteine biosynthesis are reversible; thus, all these organosulfurs may be recycled back to homocysteine (1).

In Saccharomyces cerevisiae, Met15 (also referred to as Met17 and Met25) catalyzes homocysteine biosynthesis (2-4). The first precise and complete deletion of MET15 (met15 $\Delta 0$) in S. cerevisiae was made in 1998 (5). Many commonly used laboratory strains contain this deletion, including BY4741, which serves as the genetic background of the haploid MATa version of the yeast deletion collection that has been extensively utilized for functional genomics and other purposes (6, 7). The use of this deletion as an auxotrophic marker, and the corresponding assumption that MET15 is required for growth in organosulfur-free media, has been applied to the interpretation of a wide range of studies (8-10). However, some evidence suggests that MET15 may not be strictly required for growth without exogenous organosulfurs. For example, the study that first identified *MET15* (11) reported that the organosulfur requirement of met15 mutants was alleviated at a critical methylmercury concentration. Papillae have also been seen when replica plating thick patches of $met15\Delta 0$ cells to medium lacking organosulfurs (12).

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Here, we demonstrate that $met15\Delta$ cells can synthesize organosulfurs from inorganic sulfates despite a long-held assumption to the contrary. We characterize a novel homocysteine synthase, Yll058w, and show that its activity enables $met15\Delta$ cells to grow in the absence of exogenous organosulfurs in certain experimental conditions. Specifically, stable growth is observed with automated colony transfer, but very low or no growth is observed when using traditional techniques such as thick patches or liquid cultures. We show that this appearance of auxotrophy is explained in part by toxic levels of accumulated H₂S. Our results demonstrate the existence of an alternative pathway for homocysteine production in *S. cerevisiae*, with implications for a broad range of studies that presume organosulfur auxotrophy in $met15\Delta$ cells.

Results

met15∆ cells show unexpected growth in the absence of exogenous organosulfurs

We serendipitously observed robust growth of BY4741 when cells were automatedly transferred in a 384-colony array (Fig. 1A and Table S1) from a rich medium (yeast peptone dextrose adenine [YPDA]) to a "synthetic defined" (SD) medium lacking the sulfur-containing amino acids Met and Cys, as well as any other organosulfur, and containing glucose (Glu) as the carbon source (hereafter referred to as SD-Met-Cys+Glu) (Fig. 1B). This growth was apparent from the first days of incubation, with colonies reaching saturation after \sim 360 h (Fig. S1). Both BY4741 and BY4742 (a MET15⁺ control) harbor complete deletions of URA3 and LEU2 (5), rendering them auxotrophic for uracil (Ura) and leucine (Leu). There was a large and statistically significant difference in relative colony size between BY4741 colonies that grew on SD-Met-Cys+Glu and the very small colonies detectable on SD+Met-Cys-Ura+Glu or SD+Met-Cys-Leu+Glu (Fig. 1C; effect size = 0.627, p = 0.021 [Kruskal-Wallis test] and effect size = 0.757, p = 0.007 [Kruskal–Wallis test], respectively). In this automated colony transfer procedure, BY4741 does not behave as an organosulfur auxotroph despite lacking MET15.

To confirm this unanticipated observation, we first ensured that our media were devoid of organosulfur contamination using mass spectrometry (Table S2). We next performed a cross between BY4741 and BY4742 and dissected nine fourspore tetrads. When patched to SD-Met-Cys+Glu, we observed 2:2 segregation in all tetrads, with two patches growing normally and two patches displaying extremely weak growth, consistent with a monogenic phenotype and indicating that our observations were likely not the result of a secondary mutation (Fig. S2). The extremely weak growth of patches of cells lacking MET15 was consistent with auxotrophy, in contrast to the robust growth observed with automated colony transfer. To further investigate the impact of cell propagation technique on growth, we grew BY4741 and BY4742 in liquid SD-Met-Cys+Glu medium and used drops from these cultures to make spots and patches on SD-Met-Cys+Glu solid medium (Fig. 1D). BY4741 cells grew very little in liquid cultures and not at all in spots or patches, whereas BY4742 cells grew

normally (Fig. 1, E and F). Hence, BY4741 displays auxotrophlike growth when propagated using traditional techniques but not when automatedly transferred in a grid to seed colonies.

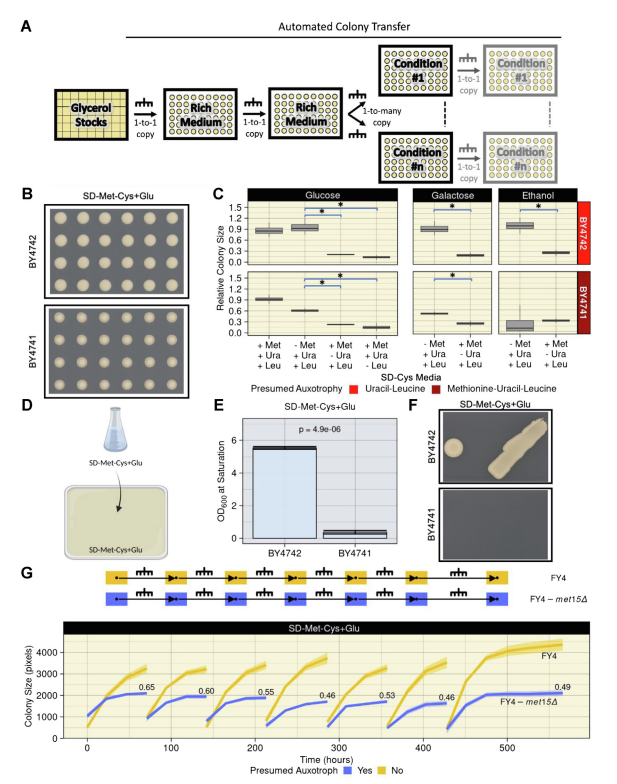
We next interrogated the extent to which the unexpected growth of BY4741 colonies depends on the specific composition of the growth medium. We repeated the automated colony transfer procedure using SD-Met-Cys and SD+Met-Cys-Ura media with either galactose (Gal) (SD-Met-Cys+Gal and SD+Met-Cys-Ura+Gal) or ethanol (EtOH) (SD-Met-Cys+EtOH, SD+Met-Cys-Ura+EtOH) as the carbon source (Fig. 1*C*). As with Glu, when Gal was used as a carbon source, BY4741 colonies grew to much larger sizes on SD-Met-Cys+Gal than on SD+Met-Cys-Ura+Gal (effect size = 0.561, p =0.021 [Kruskal–Wallis test]). In contrast, BY4741 displayed miniscule growth on SD-Met-Cys+EtOH. Robust growth was observed in "synthetic complete" (SC) medium containing all amino acids except Met and Cys (SC-Met-Cys) with either Glu or Gal as carbon source (Fig. S3*A*).

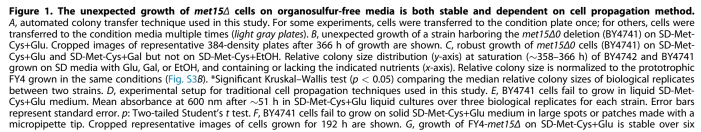
To study the consequences of *MET15* loss independent of the other auxotrophies present in BY4741, we used CRISPR to make a scarless deletion of *MET15* in the prototrophic FY4 background (FY4-*met15* Δ). We used this strain for all subsequent experiments. Like BY4741, FY4-*met15* Δ cells grew robustly when automatedly transferred to SD-Met-Cys with Glu or Gal, but not EtOH, as carbon sources (Fig. S3*B*). FY4-*met15* Δ also grew similar to BY4741 on SC-Met-Cys medium with Glu or Gal (Fig. S3*A*). Thus, the unexpected growth of *met15* Δ cells propagated *via* automated colony transfer is not dependent on the genetic background particular to BY4741.

To determine whether the robust growth of *met15* Δ colonies may be explained by *met15* Δ recycling and/or scavenging of pre-existing organosulfurs, we transferred FY4-*met15* Δ from and to SD-Met-Cys+Glu six consecutive times. FY4-*met15* Δ cells displayed robust growth on SD-Met-Cys+Glu over repeated transfers (Fig. 1*G*). Relative to FY4, the fitness of these colonies was 0.65 on the initial SD-Met-Cys+Glu plates and stabilized around ~0.50. The amount of new biomass generated throughout this experiment strongly suggests that the growth of FY4-*met15* Δ cells cannot be explained by recycling or scavenging. Despite a long-standing assumption of auxotrophy, *met15* Δ strains can grow stably in media lacking organosulfurs when propagated *via* automated colony transfer.

Stable growth of met15 Δ cells on organosulfur-free media requires inorganic sulfates and is not seen in other presumed auxotrophs

To investigate whether the unexpected growth phenotype of $met15\Delta$ strains extends to other key enzymes in the organosulfur biosynthesis pathway, we again used CRISPR to make complete deletions in the FY4 background of nine additional genes (Fig. 2A). These mutants, along with FY4 and the FY4-*met15* Δ strain, were automatedly transferred (Fig. 1A) to either YPDA or SD-Met-Cys+Glu plates. All mutant strains displayed growth (or lack thereof) on SD-Met-Cys+Glu that





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was consistent with current knowledge of their roles in organosulfur biosynthesis, with the exception of FY4-*met3A* (Fig. 2B). Met3 catalyzes the first step in the reduction of intracellular sulfate and is thought to be essential for growth when unreduced sulfates are the sole sulfur source (2); however, the FY4-*met3A* mutant displayed growth on SD-Met-Cys+Glu that was roughly equivalent to that seen with the FY4-*met15A* strain (effect size = 0.651, p < 0.001 [Kruskal–Wallis test] and effect size = 0.618, p < 0.001 [Kruskal–Wallis test], respectively; Fig. 2B). The growth of FY4-*met3A* was largely unaffected by carbon source (Fig. S4).

This surprising growth of *met3* Δ cells led us to examine the level of free sulfides present in our mutant strains. To do so, we utilized a bismuth assay (13), where the darkness of the colonies is a proxy readout for sulfide accumulation. On a complete bismuth-containing medium (BiGGY), we observed colony colors (Figs. 2C and S5B) consistent with the known roles of the deleted genes in organosulfur biosynthesis (Fig. 2A). In particular, $met15\Delta$ colonies were very dark, consistent with an accumulation of sulfides expected from impaired homocysteine biosynthesis activity, and $met3\Delta$ colonies were very light, confirming that the SAP is disrupted in these cells as expected. We saw consistent results in a bismuth-containing medium lacking organosulfurs (SD-Met-Cys+Glu+Bi; Fig. S5A). In this medium, FY4 and FY4-*met15* Δ colonies were darker than in BiGGY (Fig. S5B), likely resulting from the well-characterized upregulation of the SAP triggered by Met restriction (8, 14). In contrast, FY4-met3∆ remained as light as in BiGGY, confirming again that the SAP is disrupted in these cells as expected (Fig. S5B). Therefore, the unexpected growth of FY4-met15A and FY4-met3A cells is not explained by previous mischaracterizations of the positions of the Met15 and Met3 enzymes in the organosulfur biosynthesis pathway.

To determine whether the surprising growth of FY4-met15 Δ and FY4-met3∆ cells in media lacking organosulfurs depends on the utilization of inorganic sulfates, we examined how these mutants grew over repeated transfers onto plates entirely devoid of both organosulfurs and inorganic sulfates. The growth of FY4-met15∆ cells was substantially reduced at first and essentially abolished upon a second transfer; FY4-met3 Δ cells failed to show any substantial growth upon a second transfer to medium lacking organosulfurs, irrespective of the presence or the absence of inorganic sulfates (Fig. 2D). Growth was restored for both the FY4-met15 Δ and FY4-met3 Δ strains that had stopped growing when we transferred them to SD medium containing both Met and sulfates (Fig. 2D). An increase in colony size was also observed for FY4, which had surprisingly continued to grow on media lacking both organosulfurs and inorganic sulfates even after five successive transfers, albeit to a significantly reduced colony size (Fig. 2D). Therefore, these cells, when starved for inorganic sulfates, tend to arrest rather than die, consistent with previous studies (15). Altogether, these results show that deletion of neither Met3

nor any other critical enzyme in the pathway generates the same unexpected growth phenotype as deletion of Met15, and that the unexpected stable growth of $met15\Delta$ cells requires utilization of inorganic sulfates.

YLL058W encodes an alternative homocysteine synthase

Our observations led us to hypothesize that the *S. cerevisiae* genome encodes at least one enzyme other than Met15 capable of catalyzing *de novo* homocysteine biosynthesis. We therefore examined the targets of the sole known transcriptional activator of the sulfur metabolic network, Met4 (16). Two of these targets—YHR112C and YLL058W—do not have a known function according to the Saccharomyces Genome Database (17) but are predicted by sequence analysis to belong to the same pyridoxal 5'-phosphate (PLP)-dependent transferase superfamily as Met15 (18). Of these two, YLL058W was the more compelling candidate because it has recently been implicated in sulfur metabolism by quantitative trait locus mapping (19) and encodes a protein predicted to have the ability to catalyze homocysteine biosynthesis by a yeast genome-scale metabolic model (20).

YLL058W is located near the telomere of chromosome XII within a cluster of genes that are transcriptionally regulated by Met4, several of which have a direct role in sulfur metabolism (Fig. 3A) (16). There has been speculation that it may have arisen as a result of horizontal transfer (16)-a relatively rare occurrence in yeast. YLL058W has two close paralogs in the S. cerevisiae genome: STR2, which encodes an enzyme that converts Cys into cystathionine (Fig. 2A), and the uncharacterized YML082W, which is a copy of STR2 produced during the whole-genome duplication event (21). We generated and clustered a collection of YLL058W, STR2, and YML082W homologs from bacterial and fungal genomes on the basis of sequence similarity. If YLL058W were derived from a recent horizontal transfer event, we would expect it to be more similar to some of its homologs outside Ascomycota than to its Ascomycota homologs. Instead, all homologs outside Ascomycota grouped together along with all budding yeast homologs more distantly related to S. cerevisiae than the Hanseniaspora genus. The homologs found in all descendants of the common ancestor of Saccharomyces and Hanseniaspora comprised two additional groups, one containing YLL058W and the other containing both STR2 and YML082W (Fig. S6). The genes belonging to the YLL058W class and the STR2-YML082W class appear at approximately the same point in the phylogenetic tree (Fig. 3B), prior to whole-genome duplication. Therefore, we infer that an ancestral gene duplicated into the YLL058W and STR2-YML082W ancestors, followed by substantial sequence divergence of both copies, and later by duplication of the STR2-YML082W ancestor. Interestingly, in all species that contain genes of the YLL058W class, the YLL058W homolog is located near the telomeres among a

consecutive automated colony transfers. *Top*, experimental design. *Bottom*, colony size (*y*-axis) over time (*x*-axis). *Lines* represent the mean colony size at each time point. *Shaded area* represents one standard deviation. Numbers next to the *blue curves* indicate the fitness of FY4-*met15* relative to FY4 as determined by dividing median colony sizes at the final time point of each curve. Cys, cysteine; EtOH, ethanol; Gal, galactose; Glu, glucose; Met, methionine; SD, synthetic defined.

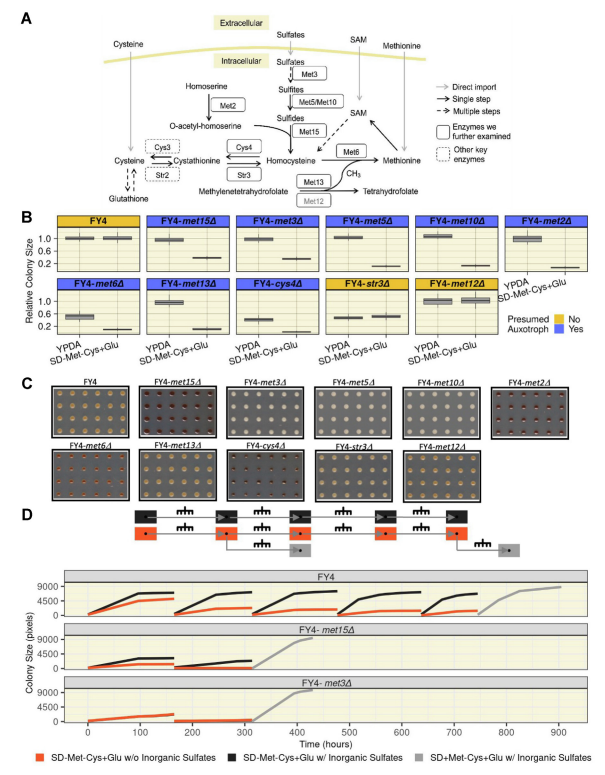


Figure 2. Growth of *met15* Δ colonies in organosulfur-free media is specific and dependent on the utilization of inorganic sulfates. *A*, The *Saccharomyces cerevisiae* organosulfur biosynthesis pathway. Exogenous organosulfurs such as Met, cysteine, and SAM can be directly imported (*gray arrows*); inorganic sulfates can also be imported and reduced to sulfides *via* the SAP. Sulfides are converted into organosulfurs by Met15, which catalyzes *de novo* homocysteine biosynthesis. Homocysteine is then converted to critical downstream organosulfurs, a selection of which are indicated. *In vivo* Met biosynthesis requires both the Met synthase Met6 and the methylenetertahydrofolate reductase Met13. The Met13 isozyme, Met12, is shown in *gray*. Enzymes are indicated within *boxes*, next to the reactions they catalyze, which are illustrated with *arrows* indicating reaction directionality. Draving inspired by (52). *B*, *met3* Δ cells also show unexpected growth. Box plots show relative colony size data (*y*-axis, normalized to FY4) at saturation (~160–173.5 h in SD-Met-Cys+Glu; 83.5–89 h in YPDA). *C*, disruptions of the organosulfur biosynthesis pathway result in a range of H₂S accumulation in cells. Representative crops of plate images of strains transferred to complete medium containing bismuth (BiGGY) for ~160 to 173.5 h. *D*, growth of FY4-*met15* Δ colonies requires inorganic sulfates. *Top*, experimental design. *Bottom*, growth curves (colony size, *y*-axis) of the indicated strains (subpanels) through the entire experimental procedure (time, *x*-axis) with the mean colony size depicted. The *orange* (SD-Met-Cys+Glu without inorganic sulfates) and *black curves* (SD-Met-Cys+Glu with norganic sulfates) in the lower subpanel (FY4-*met3* Δ) overlap. Cys, cysteine; Glu, glucose; H₂S, hydrogen sulfide; Met, methionine; SAP, sulfate assimilation pathway; SD, synthetic defined; YPDA, yeast peptone dextrose adenine.

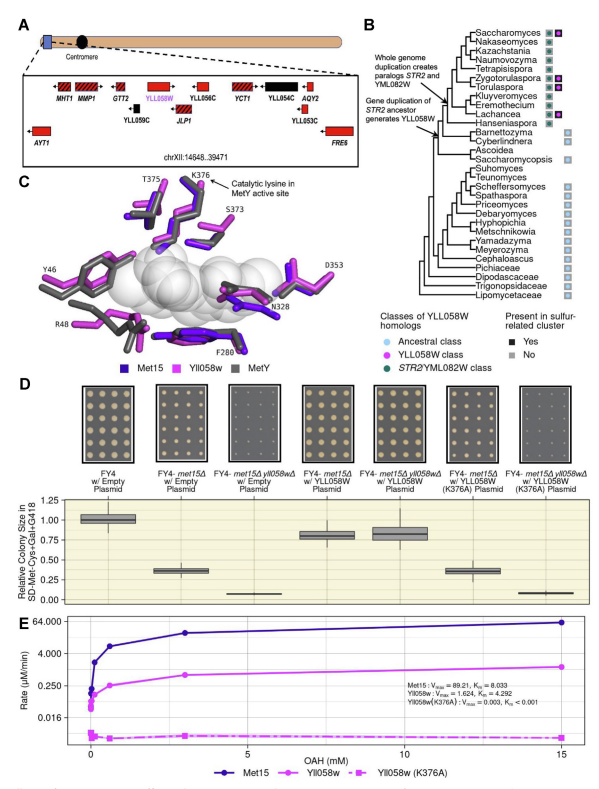


Figure 3. YII058w functions as an inefficient homocysteine synthase. *A*, genomic context of YLL058W (17). In *Saccharomyces cerevisiae*, YLL058W (*magenta*) is located near a cluster of genes that are transcriptionally activated by Met4 ((16), *red boxes*), several of which have a direct role in sulfur metabolism (*red boxes* with *black stripes*). *Black boxes* indicate ORFs that fall into neither category. *B*, distribution of YLL058W homolog clusters across the budding yeast phylogenetic tree. The tree topology follows (47). *Circles* adjacent to each taxon indicate whether YLL058W homologs belonging to each of three classes are found in the genome of that taxa, whereas *black boxes* indicate the presence of a cluster of sulfur-related genes, as depicted in *A* for *S. cerevisiae*. *C*, YII058w has structural similarity to a homocysteine synthase. Overlap of the active site residues of MetY from *Thermotoga maritima* (Protein Data Bank ID: 7KB1) (*gray*) with the predicted structures of YII058w (*magenta*) and Met15 (*violet*), calculated using AlphaFold, is shown. A reaction intermediate captured within the MetY structure (23) is shown in *semitransparent spheres*. Residue labeling corresponds to the YII058w sequence. *D*, YII058W is critical for growth in the absence of *MET15*. The indicated strains bearing high-copy (2µ ORI) plasmids either empty or overexpressing YLL058W or YLL058W (K376A) were grown on SD-Met-Cys+Gal+G418 medium. The *box plots* show relative colony sizes at saturation (296 h). Cropped representative images of the plates corresponding to each strain are shown above. *E*, YII058w is a less efficient homocysteine synthase. Enzyme kinetic assay was carried out using



collection of other genes related to sulfur metabolism (Fig. 3, A and B and Table S3). The preservation of this pattern in the genomically unstable near-telomere region is consistent with a strong selective constraint keeping YLL058W in proximity to other genes involved in sulfur metabolism.

We next examined the predicted structure of the putative Yll058w enzyme. To this aim, we compared structures predicted by AlphaFold2 (22) for Yll058w and Met15 to the experimentally determined crystal structure of MetY (Fig. S7A), a bacterial Met15 ortholog that catalyzes homocysteine biosynthesis from OAH and bisulfide (23). The MetY crystal structure captured a reaction intermediate, allowing for a precise determination of active-site residues. The predicted structures of both Met15 and Yll058w are highly similar to MetY within the active site, including the catalytic lysine K376 (Figs. 3C and S7B). MetY and Met15 function as homotetramers (23, 24), with multimerization domains allowing the distinct monomers to come together and complete the active site (Fig. S7C). In contrast, Yll058w lacks a multimerization domain and is predicted to be monomeric. However, Yll058w contains an approximately 180 amino acid region that has no homology to MetY or Met15. This region is predicted to fold back onto the core of the protein, supplying the missing amino acids to complete the active site (Fig. S7D). Altogether, these comparative structural analyses suggest that a Yll058w monomer may be capable of carrying out catalysis similar to that of the MetY and Met15 tetramers.

We next employed a genetic approach to determine whether the putative Yll058w enzyme is necessary and sufficient to support the growth of *met15* Δ cells in the absence of exogenous organosulfurs. We used CRISPR to delete YLL058W and to make a met15 yll058w double mutant in the FY4 background. We also constructed two high copy-number plasmids, one overexpressing YLL058W, and the other overexpressing a YLL058W mutant in which the putative catalytic lysine (K376, Fig. 3C) was changed to an alanine (A376). The FY4-yll058w Δ single mutant resulted in only a very modest growth defect on SD-Met-Cys+Glu (effect size = 0.054, p > 0.05 [Kruskal-Wallis test]) (Fig. S8A) and wildtype levels of H₂S accumulation (Fig. S8B). Strikingly however, the FY4-met15 Δ yll058w Δ double mutant showed little to no growth in the absence of exogenous organosulfurs (Fig. 3D), indicating a synthetic lethal interaction between MET15 and YLL058W. Overexpression of YLL058W fully compensated for the loss of MET15, both alone (effect size= 1.238, p < 0.001 [Kruskal–Wallis test]) and in combination with the loss of YLL058W (Fig. 3D; effect size = 10.117, p < 0.001 [Kruskal–Wallis test]). In contrast, the overexpression of YLL058W (K376A) in these same strain background has no rescue effect (Fig. 3D). These results strongly suggest that the growth of $met15\Delta$ cells in the absence of exogenous organosulfurs is mediated by the enzyme encoded by YLL058W.

To directly examine the ability of Yll058w to catalyze homocysteine biosynthesis, we first purified recombinant Yll058w and Yll058w (K376A) proteins from Escherichia coli, along with Met15 as a positive control (Fig. S9). We then carried out in vitro reactions using the canonical OAH and H₂S substrates as well as PLP, a cofactor critical for Met15 activity. Yll058w was able to catalyze homocysteine production significantly above background, albeit much less efficiently than Met15, with the Met15 reactions yielding \sim 9.9 times as much homocysteine at the 15 min time point (Fig. S10A). We then measured enzyme kinetic parameters by performing analogous reactions with varying amounts of the limiting substrate, OAH (Fig. 3E). Yll058w activity was consistently above background and increased with increasing OAH concentrations (V_{max} : 1.62; K_m : 4.29), although remaining less efficient than Met15 activity at all OAH concentrations (V_{max} : 89.21; K_m : 8.03). The putative catalytic dead mutant Yll058w (K376A) had no detectable activity. We therefore conclude that YLL058W encodes an inefficient homocysteine synthase capable of catalyzing the same reaction as Met15.

To investigate whether the inefficient enzymatic activity of Yll058w would be sufficient to support growth of *met15* Δ cells in the absence of exogenous organosulfurs, we adapted an enzyme-constrained version of the consensus genome-scale metabolic model for *S. cerevisiae*, Yeast8 (25). We removed *MET15* from the model, added the Yll058w reaction we observed *in vitro*, and evaluated the flux through the biomass equation at a range of arbitrarily assigned catalytic efficiencies (K_{cat}) for Yll058w. We found that an appreciable reduction in biomass occurred only when Yll058w was assigned a catalytic efficiency ~100× lesser than that assigned to Met15 in the model (Fig. S10*B*). These simulations, together with our structural, *in vivo*, and biochemical findings, demonstrate that it is plausible that Yll058w supports growth of *met15* Δ cells in the absence of exogenous organosulfurs.

Removal of H_2S enhances growth of met15 Δ cells in organosulfur-free media

We hypothesized that the large excess of H₂S present in $met15\Delta$ cells (Figs. 2C and S5) may explain why these cells do not grow without organosulfurs using traditional growth techniques despite having the capacity to synthesize homocysteine through Yll058w. This accumulation of H₂S is expected since the reaction catalyzed by Met15 is one of the major sulfide-consuming reactions in the cell (26) and since Met restriction triggers upregulation of the SAP (8, 14). Although some amount of H₂S is required for cell growth, at high levels, it is toxic in a range of species (27–29), including in *S. cerevisiae* (30). To investigate this hypothesis, we examined the growth of FY4 and FY4-*met15* Δ cells in liquid SD-Met-Cys+Glu cultures with or without the H₂S chelator ferric

the recombinantly expressed purified proteins used for Fig. S9, Met15 *purple solid line*, YII058w *pink solid line*, and the putative catalytic dead mutant of YII058w (*pink dashed line*). Varying concentrations of the substrate, *O*-acetyl-L-homoserine (OAH), are shown on the *x*-axis and the rate of reaction on the *y*-axis. Inferred V_{max} and K_m values of Met15, YII058w, and YII058w (K376A) are shown in the panel and Table S10. Cys, cysteine; Gal, galactose; Met, methionine; SD, synthetic defined.

(Fe)-EDTA, which bonds free sulfides that exit the cell by diffusion (31, 32) (Fig. 4*A*). In the presence of Fe-EDTA, we observed a modest increase in the growth of FY4 cultures (p = 0.041 [Student's *t* test]; Fig. 4*B*) and a striking increase in the growth of FY4-*met15* Δ cultures (p = 0.01 [Student's *t* test]; Fig. 4*B*).

To rule out spontaneous mutations as the reason for the growth increase of FY4-*met15* Δ cultures, we repassaged the

cultures in SD-Met-Cys+Glu medium lacking chelator (Fig. 4*A*). The repassaged FY4-*met15* Δ cultures showed little to no growth (Fig. 4*B*). Interestingly, the repassaged FY4 cultures that had been propagated in the presence of chelator saturated at a higher absorbance at 600 nm than those that had not (*p* = 0.0012 [Student's *t* test]; Fig. 4*B*). We then placed cells from both the original and the repassaged cultures on solid media plates in large drops and patches (Fig. 4*C*). The FY4-*met15* Δ

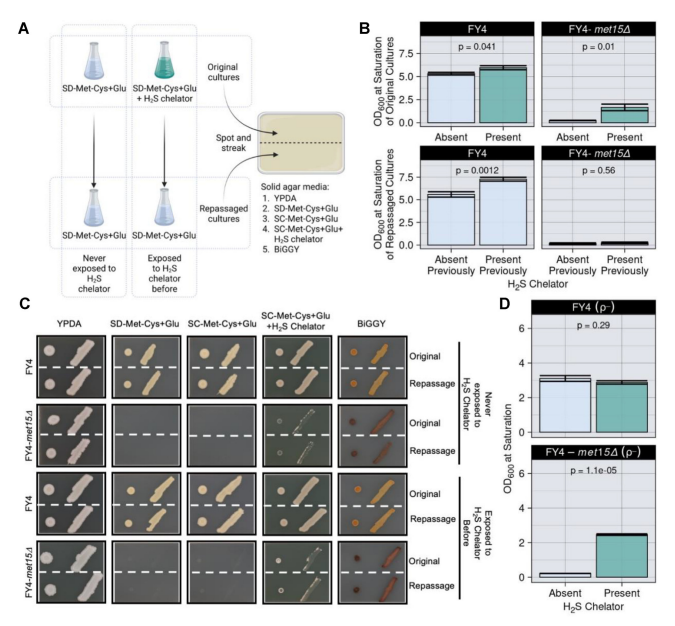


Figure 4. Chelation of H₂S facilitates the growth of *met15* Δ cells in the absence of exogenous organosulfurs. *A*, experimental setup. Strains were grown in either SD-Met-Cys+Glu or SD-Met-Cys+Glu+Fe–EDTA liquid media (top two flasks, original cultures). Once saturation was reached for all cultures, cells were repassaged in SD-Met-Cys+Glu media (bottom two flasks, repassaged cultures) and again grown to saturation. Ten microliters from each culture were then spotted and struck to the indicated solid media using a micropipette tip. *B*, the H₂S chelator Fe–EDTA partially rescues growth for *met15* Δ cells in liquid media lacking organosulfurs. Mean absorbance at 600 nm values of the cultures after ~51 h of growth with or without chelator (*top panel*). Exposure to the chelator does not induce heritable growth-enabling mutations (*lower panel*). Mean absorbance at 600 nm values of the cultures after ~51 h of growth with or without chelator (*top panel*). Exposure after repassage. Error bars represent standard error; *p*: two-tailed unpaired Student's *t* test. Individual values for each replicate, from both independent experiments, are shown in Table S4; outlier FY4-*met15* Δ cultures that showed unexpected growth in SD-Met-Cys+Glu were excluded from the data presented here and indicated in *green text* in the table. *C*, chelation of H₂S partially rescues the growth of FY4-*met15* Δ cells grown in spots or patches (*A*). Cropped representative images of cells grown for 192 h are shown. *D*, growth defect of *met15* Δ petite (ρ -) derivatives in liquid media lacking organosulfurs for each experiment in same as in *B*. Cys, cysteine; Fe, ferric; Glu, glucose; H₂S, hydrogen sulfide; Met, methionine; SD, synthetic defined.

cells failed to grow on SD-Met-Cys+Glu and SC-Met-Cys+Glu in drops or patches, irrespective of their previous exposure to the chelator (Fig. 4*C*). These results demonstrate that the growth of FY4-*met15* Δ cultures in the presence of chelator is not facilitated by heritable mutations. Growth in drops and patches of FY4-*met15* Δ cells was partially rescued by adding Fe-EDTA to the organosulfur-free medium (Fig. 4*C*). Thus, the growth defect (in some settings) and auxotrophic-like behavior (in other settings) of *met15* Δ cells grown without organosulfurs is at least partially attributable to H₂S toxicity.

Excess H₂S is often associated with mitochondrial dysfunction (33). Yeast strains with damaged or absent mitochondrial genomes display the so-called "petite" phenotype (34). To determine whether the partial rescue of $met15\Delta$ cells by an H₂S chelator was dependent on the mitochondrial genome, we constructed petite (ρ -) derivatives of FY4 and FY4-*met15* Δ and grew them in SD-Met-Cys+Glu cultures with or without Fe-EDTA. We observed a strong rescue of FY4-met15Δ ρgrowth upon addition of the chelator ($p = 1.1 \times 10^{-5}$ [Student's t test]; Fig. 4D, bottom panel). This result indicates that the chelator-mediated rescue of met15A cell growth in liquid cultures does not require an intact mitochondrial genome. Next, we examined the growth of these petite strains and their respective nonpetite parental strains in our automated colony transfer pipeline (Fig. 1A). The cells were transferred to solid media containing or lacking Met, and with either Glu or EtOH as the carbon source (Fig. S11). In SD+Met-Cys+EtOH, nonpetite FY4-met15 Δ cells grew poorly relative to their MET15⁺ counterparts. In SD-Met-Cys+Glu, the FY4-met15Δ ρ- strain grew almost as well as their nonpetite parental strain. These results suggest that met15A cells experience some degree of mitochondrial dysfunction that impairs their ability to metabolize EtOH, but that neither their unexpected growth in automated colony transfer settings nor the chelator-mediated rescue of their lack of growth in traditional propagation techniques require a functional mitochondrial genome.

Discussion

Here, we show that, contrary to long-standing assumptions, the organosulfur auxotrophy-like phenotype in a $met15\Delta$

background is dependent on cell propagation technique. Our analyses demonstrate that the S. cerevisiae genome, and possibly those of several closely related fungal species, encode another enzyme, the previously uncharacterized Yll058w, that can catalyze the same reaction as Met15, albeit less efficiently. This activity compensates for the loss of MET15 and enables the growth of met15A cells in the absence of exogenous organosulfurs when cells are propagated using automated colony transfer or grown in the presence of an H₂S chelator. Altogether, our observations support a model according to which $met15\Delta$ cells can synthesize enough homocysteine to support growth through the activity of Yll058w but often experience chronic H₂S toxicity that prevents growth when standard laboratory techniques are used (Fig. 5). Given its evolutionary conservation, Yll058w may well have other activities that are not investigated here. For example, YLL058W was recently implicated in S-methylmethionine metabolism by quantitative trait locus mapping (19).

H₂S-mediated dependence of a classic auxotrophy

 H_2S levels are extremely high when *met15* Δ cells are growing without exogenous organosulfurs (Figs. 2C and S5), as a result of the upregulation of the SAP (8, 14) and Yll058w being an inefficient homocysteine synthase. Our results indicate that H₂S toxicity is at least partially responsible for the arrest and/or death that occurs in *met15* Δ cells grown using traditional techniques (Fig. 5). A primary mechanism for the toxic effects of elevated H₂S, seen in many organisms, involves mitochondrial dysfunction (33). The failure of $met15\Delta$ cells to utilize EtOH (Figs. 1C, S3B, and S11) is consistent with mitochondrial dysfunction. However, the ability of the H₂S chelator Fe-EDTA to restore the growth of FY4-met15A ρ -cells grown in liquid cultures (Fig. 4D), as well as the ability of these cells to grow when automatedly transferred to solid SD-Met-Cys+Glu medium (Fig. S11), suggests that H₂S is limiting the growth of *met15* Δ cells in part through a pathway that does not involve the mitochondrial genome. The precise molecular mechanisms at play may be dissected in future studies. Our discovery that automated colony transfer allows for growth of *met15* Δ cells without organosulfurs provides a

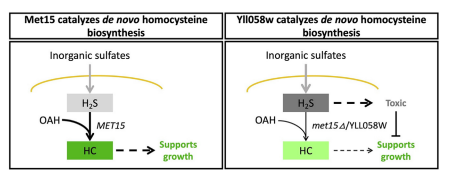


Figure 5. H_2S toxicity inhibits growth when YII058w catalyzes *de novo* homocysteine (HC) biosynthesis. Model representing the hypothesized response to organosulfur starvation in cells containing or lacking Met15. Extracellular inorganic sulfates are imported through the cell membrane (*yellow line*) and reduced to H_2S . When Met15 is present (*left*), cells efficiently convert H_2S and O-acetylhomoserine into HC, leading to a moderate amount of H_2S (*light gray*) and an adequate amount of HC (*dark green*) to support normal growth. When Met15 is absent, HC biosynthesis is catalyzed by the comparatively inefficient YIl058w enzyme (*right*). The resulting lower amount of HC (*light green*) is sufficient to support growth in some contexts, but the high amount of H_2S that accumulates (*dark gray*), exacerbated by an upregulation of the SAP (8, 14), can become toxic and lead to cell death or arrest in other contexts. H_2S , hydrogen sulphide; Met, methionine; SAP, sulfate assimilation pathway.

new tool to investigate this question as well as, more generally, the response to extreme sulfur starvation and associated phenotypes such as aging.

We speculate that, when cells are grown via automated colony transfer, excess sulfides can be eliminated through diffusion between the evenly spaced small colonies, whereas, in thick patches of cells, excess sulfides are secreted and affect neighboring cells, creating group-level toxicity and selflimiting growth. Similarly, we speculate that H₂S toxicity is high in liquid cultures because the semisealed environment limits gas exchange. Interestingly, a recent preprint suggested that when *met15* Δ cells are grown to a high density in liquid cultures, they can keep growing after having exhausted exogenous organosulfurs by utilizing the excess H₂S secreted in the prior phase of the culture (35). This preprint implicated Yll058w as the key enzyme allowing cells to reincorporate the sulfide ions and showed that the equilibrium of H₂S levels between the aqueous (retained in the media and accessible) and gas phases (escaped and inaccessible) determined whether organosulfur auxotrophy could be rescued. Our combined observations hint at a cell-density dependent rescue of organosulfur auxotrophy that is influenced by the interplay between the amount of H_2S that is (1) produced by the cells, (2) retained in the microenvironment, and (3) released into the atmosphere. Differences in cell propagation techniques likely introduce unappreciated differences in the dynamics of gas exchange that determine whether $met15\Delta$ cells grow, arrest, or die. While the conditional essentiality of genes has been well studied in the context of differing genetic backgrounds or environments (36, 37), to our knowledge, MET15 represents the first report of a case in which the essential nature of a gene is dependent on the way cells are propagated.

An expanded yeast sulfur metabolic network

It is of note that even the prototrophic FY4 strain displayed enhanced growth in liquid media when cells were previously exposed to an H_2S chelator (Fig. 4B). While it is possible that Fe-EDTA enhances the growth of even MET15⁺ strains because of some secondary effect unrelated to the chelation of H₂S, previous studies showed that very little of the chelator is taken up by yeast cells (31). Furthermore, our data indicate that wildtype strains accumulate H₂S to a level that is roughly intermediate between that seen in $met15\Delta$ cells and mutants lacking an intact SAP (Figs. 2C and S5). This suggests that the possibility that (1) even wildtype cells suffer some degree of H₂S-induced stress when grown in organosulfur-free media and (2) removal of excess H₂S allows the cells to adapt to this stress, likely at the level of gene expression or epigenetics. This may have implications for anyone seeking to maximize cellular growth in media lacking, or containing low amounts of, organosulfurs, whether in an industrial or a research setting. Also surprising was the limited but continued growth of FY4 cells even after multiple successive transfers to media lacking both organosulfurs and inorganic sulfates (Fig. 2D). This continued growth was unanticipated given that to our knowledge the SAP relies on inorganic sulfates (Fig. 2A).

Perhaps Met15, but not Yll058w, can metabolize derivatives of other sulfur-containing compounds present in our growth media (*e.g.*, biotin). Altogether, our work suggests that there may yet be more unknown factors that contribute to the yeast sulfur metabolic network.

Sulfur-containing amino acids act as cellular sensors for growth control (38, 39). In particular, Met restriction is known to increase longevity in a broad range of organisms (40) and may serve as a therapeutic treatment for certain cancers (41). Mutants lacking MET15 have been widely used in the study of life span extension in yeast (8-10, 42). Our data add a potential layer of nuance to such studies, including those that have explored the role for H₂S specifically in life span extension (e.g., (43)). More generally, our discovery of the alternative homocysteine synthase Yll058w may influence the interpretation of the plethora of published studies, including genomewide screens conducted in the BY4741 background, wherein the results or methodologies assumed $met15\Delta$ cells to be auxotrophic for organosulfurs. Our findings that S. cerevisiae can grow in the absence of exogenous organosulfurs despite very low levels of de novo homocysteine biosynthesis, that the previously observed auxotroph-like phenotypes are dependent on cell propagation techniques, and that H₂S levels control group-level growth behaviors, provide an additional avenue for studying the interplay between sulfur metabolism and its many dependent phenotypes.

Experimental procedures

This article contains supporting information, including Figures S1–12 and Tables S1–S10, Supporting File S1, and Supporting Materials and Methods.

Yeast strains and growth media

See Supporting Materials and Methods and Tables S6–S8 for a list and description of all strains, plasmids, guide RNAs, and repair fragments used for this study. See Table S9 for recipes for the various growth media used in this study. For each strain, we selected four independent colonies from streakouts, which were used to inoculate liquid YPDA medium and grown overnight at 30 °C. These cultures were considered biological replicates. The cultures were then used to create 384-well glycerol stocks (glycerol 25% v/v), with each biological replicate occupying one-fourth of the plate, distributed uniformly.

Automated colony transfer

Using the benchtop RoToR HDA robotic plate handler (Singer Instruments Co Ltd), strains were transferred from glycerol stocks to YPDA agar plates and incubated at 30 °C for 72 h (see Table S1 for a description of the settings used). Colonies on each plate were transferred to two YPDA plates (representing technical replicates) and again incubated at 30 °C for 48 h; this step reduces the variability in colony sizes introduced when cells are transferred from glycerol stocks. Finally, these plates were "one-to-many" transferred onto various solid "condition" media plates. Condition plates were



incubated at 30 °C and periodically imaged until saturation. This protocol, depicted in Figure 1*A*, was carried out at least twice per experiment to control for batch effect. Figures 1, *B*, *C*, 1*G*, 2, *B*, *C*, *D*, 3*D*, S1, S3–S5, S8, and S11 were carried out in this manner.

Relative fitness measurement

A spatially cognizant colony size database was built using colony size estimations from the plate images per the linear interpolation-based detector framework for high-throughput colony-based screening analysis (44). For all experiments, border colonies were removed from the dataset as these tend to grow more because of greater nutrient access, leaving a maximum of 77 colony size observations per strain per time point per plate per biological replicate. Outlier colony size values were then removed using two median-adjusted deviations as a cutoff. These raw colony size values are reported in Figures 1G and 2D. Otherwise, the median colony size per time point of the prototrophic strain FY4 was used to calculate relative colony size of every colony per strain per time point. At saturation, the relative colony size distribution per strain is determined by the median relative colony sizes per plate per biological replicate. This relative colony size distribution is compared amongst strains using the nonparametric Kruskal-Wallis test to determine significant changes in fitness. Effect size values are calculated as the difference in median colony sizes divided by the median colony size of the strain being used as the reference. In addition, in Fig. S11, growth curve analysis was done with the growthcurver (45) package in R (46), using the time series colony size data per colony per strain per technical replicate per biological replicate. These data were first converted into an equally spaced time series using a second-degree locally weighed scatterplot smoothing (loess) function to generate 100 equally spaced data points. This was then fed into the "SummarizeGrowthByPlate()" function of the growthcurver package (45) to estimate the area under the curve (AUC) per colony per strain in the experiment. AUC summarizes a growth curve by integrating the contributions of the initial population size, growth rate, and carrying capacity into a single value, making it a more robust estimation of fitness. Relative AUCs, effect sizes, and significant changes were estimated the same way as the colony size data described previously. The AUCs from the data in Fig. S11 were used to calculate the statistics in Table S5.

Evolutionary origins of YLL058W

Genome sequences of 332 budding yeasts were taken from a published dataset (47). Each genome was scanned for ORFs (consisting of the sequence between an ATG start codon and a stop codon in the same frame) larger than 300 bp. BLASTP was then run with default settings and a $10^{-4} e$ value cutoff on the predicted protein products of these ORFs to identify the strongest match among *S. cerevisiae* annotated genes. All ORFs for which the best match (*i.e.*, the lowest BLASTP *e* value) was either the *S. cerevisiae* YLL058W sequence or the sequence of one of its two *S. cerevisiae* homologs (*STR2* and

YML082W) were retained. For each homolog identified, we also noted the best *S. cerevisiae* matches for the five nearest ORFs in each direction that had a match. To supplement this list with homologs from more distant species, we submitted the *S. cerevisiae* YLL058W sequence to the HMMER homology search tool (48). From these results, we selected the strongest matching sequences (lowest *e* values) from a variety of bacterial and fungal taxa outside the *Ascomycota* phylum (four bacterial and nine fungal outgroups). The full list of YLL058W homologs and their sequences is given in File S1. We next constructed a multiple alignment of all homologs using the MUSCLE aligner (49). We used the R package bio2mds to perform K-means clustering on the sequence alignment and to visualize the results using multidimensional scaling (Fig. S6).

In vitro homocysteine assay

In vitro homocysteine biosynthesis was carried out largely as described previously (3). Aliquots of Met15, Yll058w, and Yll058w (K376A), purified as described (Supporting Materials and Methods and Fig. S9), were dialyzed into a final storage buffer of 20 mM Tris-Cl (pH 8.0), 500 mM NaCl, 20 mM imidazole, 0.2 mM PLP, and 10% glycerol, snap frozen, and stored at -80 $^\circ$ C. Aliquots were thawed on ice, and the enzymes were diluted in the storage buffer to a concentration of 60 nM and preincubated at room temperature for 10 min. To compare enzymatic activity of Met15 with Yll058w at a saturating substrate concentration, the enzyme solution was then mixed with diluted 10×, giving a final enzyme concentration of 6 nM, with a reaction mixture containing 50 mM potassium phosphate buffer (pH 7.8), 1.0 mM EDTA (pH 8.0), 15.0 mM OAH, 5.0 mM NaHS, and 0.6 mM PLP. The reaction components were then mixed by gentle pipetting and incubated at 30 °C. Each reaction was performed in triplicate. Samples were removed at six time points (0, 1, 3, 5, 10, and 15 min), and the reactions were stopped by adding 1.0 M HCl at a ratio of 10 ml for every 100 ml of sample and then snap frozen. To calculate K_m and V_{max} for Met15, Yll058w, and Yll058w (K376A), the reaction was performed for 10 min with a range of OAH from 0 to 15 mM as indicated. In the same buffer as the reaction mixture but without any enzyme or substrate (but with acid added), we also generated a homocysteine standard curve with seven serial 1:5 dilutions ranging from 15 mM to 0.00096 mM, as well as a 0 mM "blank" sample. Homocysteine levels were measured as described in Supporting Materials and Methods.

Metabolic modeling

Metabolic modeling simulations were performed with a Python script making use of the COBRApy toolkit (50) and the ecYeastGEM_batch model (25), an "enzyme-constrained" variant (51) of Yeast8. The enzyme-constrained feature of the steady-state model is achieved by treating enzymes as metabolites that are temporarily "consumed" (*i.e.*, removed from the available pool) while they are engaged in catalysis. The script first removes all pre-existing reactions (which have not been experimentally validated) contributing to the "consumption" of

Yll058w (r 0815No1, r 0815REVNo1, and arm r 0815). It then adds the homocysteine-generating reaction that we observe in our in vitro assay, which uses H2S and O-acetylhomoserine as substrates and yields homocysteine, acetate, and hydrogen as products, with Yll058w as the catalyst. This reaction was added with a range of K_{cat} values assigned to Yll058w in order to evaluate the "biomass" generated over a broad range of enzyme efficiencies. The reaction catalyzed by Yll058w was made to be bidirectional; however, flux was only observed in the forward (homocysteine-producing) reaction. MET15 was then removed from the model, eliminating all reactions that the encoded enzyme was associated with. Finally, the model was "solved" to obtain the flux through the biomass equation, a proxy for fitness. The default growth medium associated with the model, which contains sulfates but lacks organosulfurs, was used for all simulations.

Chelation of H₂S with Fe-EDTA sodium salt

For each strain, five single colonies were scraped from a YPDA agar plate, and cells were resuspended in SD-Met-Cys+Glu medium. Absorbance at 600 nm of the suspension was measured, and 25 ml cultures were inoculated at a starting absorbance at 600 nm of 0.1 in SD-Met-Cys+Glu or SD-Met-Cys+Glu supplemented with 0.048 M Fe-EDTA (sodium salt; catalog no.: E6067, MilliporeSigma) in 125 ml Erlenmeyer flasks (optimal Fe-EDTA dosage was determined by a titration experiment [Fig. S12]). For the nonpetite strains, the experiment was performed twice; each time, cultures were prepared in triplicate. The petite strains were tested in a single experiment using four biological replicates. Cultures were incubated at 30 °C with 150 rpm shaking, and absorbance at 600 nm was measured regularly until saturation was reached (\sim 51–53 h). For the repassaging of the nonpetite strains (Fig. 4, A-C), cells were then pelleted to facilitate the exchange to fresh SD-Met-Cys+Glu medium, again using a starting absorbance at 600 nm of 0.1, and incubated as before for \sim 51 h with regular absorbance at 600 nm measurements. A portion of each culture was stored at 4 °C. Ten microliter drops of each original and repassaged culture were then both spotted with a pipette and struck with a sterile pipette tip, on agar plates (YPDA, SD-SC-Met-Cys+Glu, Met-Cys+Glu, SC-Met-Cys+Glu+[Fe-EDTA], and BiGGY) that were incubated at 30 $^{\circ}C$ (Fig. 4C) and imaged regularly with spImager Automated Imaging System (S&P Robotics, Inc).

Data availability

All data generated/analyzed in this study are available in the main text, in the supporting figures and tables, and as supporting data files. Supporting information is available at https://bit.ly/3GIgpld. All images and statistical analyses in this study were generated using code available at https://github.com/sauriiiin/methionine/tree/master/paper.

Supporting information—This article contains supporting information (23).

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Conflict of interest—A.-R. C. is a member of the scientific advisory board for Flagship Labs 69, Inc (ProFound Therapeutics). T. I. is a cofounder of Data4Cure and has an equity interest. T. I. is on the Scientific Advisory Board of IDEAYA Biosciences, Inc, has an equity interest, and receives income. The terms of these arrangements have been reviewed and approved by the University of California San Diego in accordance with its conflict of interest policies. All other authors declare that they have no conflicts of interest with the contents of this article.

Abbreviations—The abbreviations used are: AUC, area under the curve; Cys, cysteine; EtOH, ethanol; Fe, ferric; Gal, galactose; Glu, glucose; H₂S, hydrogen sulfide; Leu, leucine; Met, methionine; OAH, *O*-acetyl-L-homoserine; PLP, pyridoxal 5'-phosphate; SAP, sulfate assimilation pathway; SC, synthetic complete; SD, synthetic defined; Ura, uracil; YPDA, yeast peptone dextrose adenine.

References

- Thomas, D., and Surdin-Kerjan, Y. (1997) Metabolism of sulfur amino acids in Saccharomyces cerevisiae. *Microbiol. Mol. Biol. Rev.* 61, 503–532
- Masselot, M., and De Robichon-Szulmajster, H. (1975) Methionine biosynthesis in Saccharomyces cerevisiae. I. Genetical analysis of auxotrophic mutants. *Mol. Gen. Genet.* 139, 121–132
- Chen, Z., Zhang, X., Li, H., Liu, H., Xia, Y., and Xun, L. (2018) The complete pathway for Thiosulfate utilization in Saccharomyces cerevisiae. *Appl. Environ. Microbiol.* 84, e01241-18
- Yamagata, S. (1971) Homocysteine synthesis in yeast. Partial purification and properties of O-acetylhomoserine sulfhydrylase. J. Biochem. 70, 1035–1045
- Brachmann, C. B., Davies, A., Cost, G. J., Caputo, E., Li, J., Hieter, P., et al. (1998) Designer deletion strains derived from Saccharomyces cerevisiae S288C: a useful set of strains and plasmids for PCR-mediated gene disruption and other applications. Yeast 14, 115

- Giaever, G., and Nislow, C. (2014) The yeast deletion collection: a decade of functional genomics. *Genetics* 197, 451–465
- Douglas, A. C., Smith, A. M., Sharifpoor, S., Yan, Z., Durbic, T., Heisler, L. E., et al. (2012) Functional analysis with a barcoder yeast gene overexpression system. G3 (Bethesda) 2, 1279–1289
- Zou, K., Ouyang, Q., Li, H., and Zheng, J. (2017) A global characterization of the translational and transcriptional programs induced by methionine restriction through ribosome profiling and RNA-seq. *BMC Genomics* 18, 189
- 9. Plummer, J. D., and Johnson, J. E. (2019) Extension of cellular lifespan by methionine restriction involves alterations in central carbon metabolism and is mitophagy-dependent. *Front. Cell Dev. Biol.* 7, 301
- 10. Johnson, J. E., and Johnson, F. B. (2014) Methionine restriction activates the retrograde response and confers both stress tolerance and lifespan extension to yeast, mouse and human cells. *PLoS One* 9, e97729
- 11. Singh, A., and Sherman, F. (1974) Association of methionine requirement with methyl mercury resistant mutants of yeast. *Nature* **247**, 227–229
- 12. Lab, Boeke (2003) The Met15 Resource: Update
- Nickerson, W. J. (1953) Reduction of inorganic substances by yeasts. I. Extracellular reduction of sulfite by species of Candida. J. Infect. Dis. 93, 43–56
- 14. Choi, K. M., Kim, S., Kim, S., Lee, H. M., Kaya, A., Chun, B. H., et al. (2019) Sulfate assimilation regulates hydrogen sulfide production independent of lifespan and reactive oxygen species under methionine restriction condition in yeast. Aging (Albany NY) 11, 4254–4273
- Boer, V. M., Amini, S., and Botstein, D. (2008) Influence of genotype and nutrition on survival and metabolism of starving yeast. *Proc. Natl. Acad. Sci. U. S. A.* 105, 6930–6935
- Lee, T. A., Jorgensen, P., Bognar, A. L., Peyraud, C., Thomas, D., and Tyers, M. (2010) Dissection of combinatorial control by the Met4 transcriptional complex. *Mol. Biol. Cell* 21, 456–469
- Cherry, J. M., Hong, E. L., Amundsen, C., Balakrishnan, R., Binkley, G., Chan, E. T., *et al.* (2012) Saccharomyces Genome Database: the genomics resource of budding yeast. *Nucleic Acids Res.* 40, D700–D705
- Wilson, D., Pethica, R., Zhou, Y., Talbot, C., Vogel, C., Madera, M., et al. (2009) SUPERFAMILY–sophisticated comparative genomics, data mining, visualization and phylogeny. *Nucl. Acids Res.* 37, D380–D386
- Eder, M., Sanchez, I., Camarasa, C., Daran, J.-M., Legras, J.-L., and Dequin, S. (2022) Genetic bases for the metabolism of the DMS precursor S-methylmethionine by Saccharomyces cerevisiae. *Food Microbiol.* 106, 104041
- Wang, H., Robinson, J. L., Kocabas, P., Gustafsson, J., Anton, M., Cholley, P. E., *et al.* (2021) Genome-scale metabolic network reconstruction of model animals as a platform for translational research. *Proc. Natl. Acad. Sci. U. S. A.* **118**, e2102344118
- Byrne, K. P., and Wolfe, K. H. (2005) The yeast gene order browser: Combining curated homology and syntenic context reveals gene fate in polyploid species. *Genome Res.* 15, 1456–1461
- Jumper, J., Evans, R., Pritzel, A., Green, T., Figurnov, M., Ronneberger, O., et al. (2021) Highly accurate protein structure prediction with AlphaFold. *Nature* 596, 583–589
- Brewster, J. L., Pachl, P., McKellar, J. L. O., Selmer, M., Squire, C. J., and Patrick, W. M. (2021) Structures and kinetics of Thermotoga maritima MetY reveal new insights into the predominant sulfurylation enzyme of bacterial methionine biosynthesis. *J. Biol. Chem.* 296, 100797
- Yamagata, S. (1976) O-Acetylserine and O-acetylhomoserine sulfhydrylase of yeast. Subunit structure. J. Biochem. 80, 787–797
- Lu, H., Li, F., Sánchez, B. J., Zhu, Z., Li, G., Domenzain, I., et al. (2019) A consensus S. cerevisiae metabolic model Yeast8 and its ecosystem for comprehensively probing cellular metabolism. Nat. Commun. 10, 3586
- Linderholm, A. L., Findleton, C. L., Kumar, G., Hong, Y., and Bisson, L. F. (2008) Identification of genes affecting hydrogen sulfide formation in Saccharomyces cerevisiae. *Appl. Environ. Microbiol.* 74, 1418–1427
- Blackstone, E., Morrison, M., and Roth, M. B. (2005) H2S induces a suspended animation-like state in mice. *Science* 308, 518
- Kamoun, P. (2001) Mental retardation in down syndrome: a hydrogen sulfide hpothesis. *Med. Hypotheses* 57, 389–392
- Zhang, P., Luo, Q., Wang, R., and Xu, J. (2017) Hydrogen sulfide toxicity inhibits primary root growth through the ROS-NO pathway. *Sci. Rep.* 7, 868

- Fu, L. H., Hu, K. D., Hu, L. Y., Li, Y. H., Hu, L. B., Yan, H., et al. (2014) An antifungal role of hydrogen sulfide on the postharvest pathogens Aspergillus Niger and Penicillium italicum. PLoS One 9, e104206
- Philip, C. V., and Brooks, D. W. (1974) Iron(III) chelate complexes of hydrogen sulfide and mercaptans in aqueous solution. *Inorg. Chem.* 13, 384–386
- 32. Gaensly, F., Picheth, G., Brand, D., and Bonfim, T. M. (2014) The uptake of different iron salts by the yeast Saccharomyces cerevisiae. *Braz. J. Microbiol.* 45, 491–494
- Paul, B. D., Snyder, S. H., and Kashfi, K. (2021) Effects of hydrogen sulfide on mitochondrial function and cellular bioenergetics. *Redox Biol.* 38, 101772
- 34. Ferguson, L. R., and von Borstel, R. C. (1992) Induction of the cytoplasmic 'petite' mutation by chemical and physical agents in Saccharomyces cerevisiae. *Mutat. Res.* 265, 103–148
- 35. [preprint] Yu, J. S. L., Heineike, B. M., Hartl, J., Correia-Melo, C., Aulakh, S. K., Lehmann, A., et al. (2022) Inorganic sulfur fixation via a new homocysteine synthase allows yeast cells to cooperatively compensate for methionine auxotrophy. *bioRxiv*. https://doi.org/10.1101/2022.03.14.484209
- Hou, J., van Leeuwen, J., Andrews, B. J., and Boone, C. (2018) Genetic network complexity shapes background-dependent phenotypic expression. *Trends Genet.* 34, 578–586
- 37. Zhang, Z., and Ren, Q. (2015) Why are essential genes essential? the essentiality of Saccharomyces genes. *Microb. Cell* 2, 280–287
- 38. Laxman, S., Sutter, B. M., Wu, X., Kumar, S., Guo, X., Trudgian, D. C., et al. (2013) Sulfur amino acids regulate translational capacity and metabolic homeostasis through modulation of tRNA thiolation. Cell 154, 416–429
- 39. Sutter, B. M., Wu, X., Laxman, S., and Tu, B. P. (2013) Methionine inhibits autophagy and promotes growth by inducing the SAM-responsive methylation of PP2A. *Cell* 154, 403–415
- Green, C. L., Lamming, D. W., and Fontana, L. (2021) Molecular mechanisms of dietary restriction promoting health and longevity. *Nat. Rev. Mol. Cell Biol.* https://doi.org/10.1038/s41580-021-00411-4
- Endicott, M., Jones, M., and Hull, J. (2021) Amino acid metabolism as a therapeutic target in cancer: a review. *Amino Acids* 53, 1169–1179
- 42. Ruckenstuhl, C., Netzberger, C., Entfellner, I., Carmona-Gutierrez, D., Kickenweiz, T., Stekovic, S., *et al.* (2014) Lifespan extension by methionine restriction requires autophagy-dependent vacuolar acidification. *PLoS Genet.* 10, e1004347
- Hine, C., Harputlugil, E., Zhang, Y., Ruckenstuhl, C., Lee, B. C., Brace, L., et al. (2015) Endogenous hydrogen sulfide production is essential for dietary restriction benefits. *Cell* 160, 132–144
- 44. Parikh, S. B., Castilho Coelho, N., and Carvunis, A. R. (2021) LI detector: a framework for sensitive colony-based screens regardless of the distribution of fitness effects. G3 (Bethesda) 11
- 45. Sprouffske, K., and Wagner, A. (2016) Growthcurver: an R package for obtaining interpretable metrics from microbial growth curves. BMC Bioinform. 17, 172
- 46. Team, R. C. (2021) R: A Language and Environment for Statistical Computing, R Foundation for Statistical Computing, Vienna, Austria
- 47. Shen, X. X., Opulente, D. A., Kominek, J., Zhou, X., Steenwyk, J. L., Buh, K. V., *et al.* (2018) Tempo and mode of genome evolution in the budding yeast subphylum. *Cell* 175, 1533–1545.e1520
- Eddy, S. R. (2011) Accelerated profile HMM searches. *PLoS Comput. Biol.* 7, e1002195
- 49. Madeira, F., Park, Y. M., Lee, J., Buso, N., Gur, T., Madhusoodanan, N., et al. (2019) The EMBL-EBI search and sequence analysis tools APIs in 2019. Nucl. Acids Res. 47, W636–W641
- 50. Ebrahim, A., Lerman, J. A., Palsson, B. O., and Hyduke, D. R. (2013) COBRApy: COnstraints-based reconstruction and analysis for python. *BMC Syst. Biol.* 7, 74
- 51. Sanchez, B. J., Zhang, C., Nilsson, A., Lahtvee, P. J., Kerkhoven, E. J., and Nielsen, J. (2017) Improving the phenotype predictions of a yeast genome-scale metabolic model by incorporating enzymatic constraints. *Mol. Syst. Biol.* 13, 935
- Walvekar, A. S., and Laxman, S. (2019) Methionine at the heart of anabolism and signaling: perspectives from budding yeast. *Front. Microbiol.* 10, 2624