A yeast chemogenomic screen identifies pathways that modulate adipic acid toxicity

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Highlights
- Deletion of the TCA cycle gene KGD1 improves tolerance to adipic acid and catechol
- Ergosterol and Pdr12 play non-overlapping roles protecting cell from adipic acid
- Adipic acid-induced plasma membrane localization of Pdr12 is independent of ergosterol
- Adipic acid disrupts the Membrane Compartment of Can1 (MCC) and induces endocytosis
SUMMARY
Adipic acid production by yeast fermentation is gaining attention as a renewable source of platform chemicals for making nylon products. However, adipic acid toxicity inhibits yeast growth and fermentation. Here, we performed a chemogenomic screen in <i>Saccharomyces cerevisiae</i> to understand the cellular basis of adipic acid toxicity. Our screen revealed that <i>KGD1</i> (a key gene in the tricarboxylic acid cycle) deletion improved tolerance to adipic acid and its toxic precursor, catechol. Conversely, disrupting ergosterol biosynthesis as well as protein trafficking and vacuolar transport resulted in adipic acid hypersensitivity. Notably, we show that adipic acid disrupts the Membrane Compartment of Can1 (MCC) on the plasma membrane and impacts endocytosis. This was evidenced by the rapid internalization of Can1 for vacuolar degradation. As ergosterol is an essential component of the MCC and protein trafficking mechanisms are required for endocytosis, we highlight the importance of these cellular processes in modulating adipic acid toxicity.

INTRODUCTION
Adipic acid has been described as the most important dicarboxylic acid as it is a valuable platform chemical for the production of nylon-6,6 (Polen et al., 2013). The global need for adipic acid has been projected to increase because it can be converted to other products including plasticizers, plastics, lubricants, and a food ingredient in gelatin (Musser, 2000; Polen et al., 2013; Weber et al., 2012). Nearly all of the world’s adipic acid is currently produced from a chemical process that involves the conversion of benzene to adipic acid through several chemical reactions requiring an oxidation reaction of nitric acid and air (Musser, 2000). These chemicals are costly and toxic to the environment and the nitric acid oxidation step results in the production of nitrous oxide gas, a greenhouse gas contributing to global warming and thinning of the ozone layer. Indeed, it is estimated that about 10% of the worldwide nitrous oxide gas released yearly into the atmosphere comes from adipic acid production (Alini et al., 2007).

Considering these challenges in the chemical synthesis of adipic acid, there has been a huge drive toward a more sustainable and environment friendly production of adipic acid by microbial fermentation. In this bioprocess, microorganisms such as bacteria and yeast are engineered to convert glucose to adipic acid. Yeasts, particularly <i>Saccharomyces cerevisiae</i>, are a preferred microbial host for adipic acid fermentation via the shikimate pathway, primarily because yeast metabolism has been extensively characterized and is easy to engineer (Averesch and Krömer, 2018; Nevoigt, 2008). Also, yeasts have previously been shown to be more tolerant to adipic acid toxicity than bacteria (Karlsson et al., 2017) and <i>S. cerevisiae</i> is generally robust to industrial stresses including low pH conditions (Chen et al., 2009; Fletcher et al., 2017), making it suitable for the production of organic acids including adipic acid. Although progress has steadily been made to produce adipic acid using engineered yeasts (Raj et al., 2018), the process is constrained by the toxicity of adipic acid to yeast. Hence, product toxicity is a key factor that should be considered during the design of engineered strains as this will greatly impact the final titer and productivity of the strain during the industrial production of adipic acid on a commercial scale.

Currently, the exact mechanism of adipic acid toxicity is unclear but is likely similar to the generic mechanism for organic acid toxicity. At a pH lower than the pKa of the acid (adipic acid pKa1 = 4.41, pKa2 = 5.41), organic acids freely diffuse into the cell where they then dissociate in the cytosol at a pH ~7.0 into the anionic form, which is unable to diffuse back out of the cell due to the hydrophilic nature of the anions (Jarboe et al., 2013). In order to prevent accumulation and toxicity of the anionic form of the acid, the cell...
exports the anion using specific transmembrane pumps. These transmembrane pumps have broad specificities to groups of organic acids. For example, the yeast ATP-binding cassette (ABC) superfamily transporter, Pdr12, has been reported to pump propionate, benzoate, and sorbate out of the cell but not octanoate (Kren et al., 2003; Piper et al., 1998). The transporter Qdr3, belonging to the major facilitator superfamily class, was identified as playing a role in muconic acid production and tolerance to three dicarboxylic acids including adipic acid (Pereira et al., 2019). Even though the study did not show efflux of adipate by Qdr3, it demonstrated that overexpressing Qdr3 resulted in yeast tolerance to adipic acid.

Although the effect of low pH in yeast has been well established (Piper et al., 2001; Ullah et al., 2013) the actual toxicity of organic acids, which stems from the rapid intracellular accumulation of the anion at low pH, also results in specific cellular responses. Currently, knowledge of these cellular responses to dicarboxylic acids such as adipic acid remains unclear. However, the biological mechanisms underlying the toxicity of monocarboxylic acids including sorbic acid, benzoic acid, propionic acid, and acetic acid have been thoroughly investigated (Mira et al., 2010b). Transcriptional analyses for monocarboxylic acids (acetic acid, propionic acid, benzoic acid, and sorbic acid) revealed similar transcriptional responses (Abbott et al., 2008) and also showed that four major transcriptional pathways regulated by Haa1, War1, Rim101, and Msn2/Msn4 modulate organic acid toxicity. This was observed across the different monocarboxylic acids. Functional genomics studies, on the other hand, indicated very specific toxicity mechanisms for each acid. For instance, genes associated with carbohydrate metabolism and ribosome biogenesis were specific to acetic acid, whereas genes involved in cell wall protein synthesis, tryptophan biosynthesis, and respiration were specific to benzoic acid, sorbic acid, and propionic acid, respectively (Kren et al., 2003; Mira et al., 2009, 2010b; Schüller et al., 2004). Furthermore, Trk1 involved in potassium influx was associated with propionic acid response (Xu et al., 2019). Owing to the very specific nature of the genetic mechanisms associated with different monocarboxylic acids it was not surprising that common genes were not identified for yeast evolved in parallel to three different dicarboxylic acids (glutaric acid, adipic acid, and pimelic acid) (Pereira et al., 2019). Therefore, monocarboxylic acid tolerance mechanisms in yeast cannot be used as an effective predictor of dicarboxylic acid tolerance. Toward improving adipic acid tolerance in S. cerevisiae, it is crucial to identify specific mechanisms underlying its toxicity and key genetic targets that can be engineered for tolerance.

Chemogenic screens have previously been used to identify mechanisms of toxicity not only for organic acids (Hazelwood et al., 2010; Kawahata et al., 2006; Mira et al., 2009; Piotrowski et al., 2015; Schüller et al., 2004) but also for several other toxic compounds including fermentation inhibitors (Fletcher et al., 2019; Skerker et al., 2013). This screen allows the identification of hypersensitive mutants that show significant growth defects in the presence of a toxic compound. It also allows the identification of hypertolerant mutants lacking a suppressor gene that when deleted results in improved growth in the presence of the inhibitor compound. Despite the promising potential of adipic acid production by yeast fermentation, no previous study has reported specific mechanisms underlying the acid’s toxicity to yeast. Here, we utilized a yeast chemogenic screen to identify unique biological processes that modulate adipic acid toxicity, which can be potentially targeted to engineer tolerance in yeast.

RESULTS

Chemogenic screen reveals a role for ergosterol and protein trafficking in adipic acid tolerance

In order to identify biological processes underlying adipic acid toxicity in yeast, we performed a chemogenic screen using the S. cerevisiae haploid deletion mutant array (DMA) collection to identify mutants that are hypersensitive or have increased tolerance to a sublethal concentration (80 mM) of adipic acid. The yeast DMA collection was pinned onto YPD agar plates containing 80 mM adipic acid (dissolved in DMSO), which lowered the pH of the agar plates to 4.0. To specifically identify mutants with modulated growth on adipic acid and not solely decreased pH, as a control the DMA collection was also pinned onto YPD plates containing DMSO (solvent control) and buffered at pH 4.0. We incubated the plates for 48 h after which the colony sizes of mutants grown on the adipic acid plates were compared with those of the pH 4.0 control plates and were analyzed using the synthetic genetic array (SGA) tools (Wagih et al., 2013). We identified a total of 77 adipic acid-sensitive mutants (SGA cutoff < −0.3) and one mutant that conferred improved growth on adipic acid (Table S1). To eliminate any false-positives, we quantified the growth rate of the 77 mutants in YPD medium supplemented with adipic acid at pH 4.0. The ratio of growth rate in the presence of adipic acid to that in the absence of adipic acid was obtained as the PRECOG score
which represented the level of sensitivity or tolerance of the mutants to adipic acid. Using a PRECOG score of $R^2 > 0.5$, signifying more than 50% growth inhibition in the presence of adipic acid, we confirmed 43 mutants to be sensitive to adipic acid. In contrast only one suppressor or deletion mutant was confirmed to grow better in adipic acid (PRECOG score $> 0.8$).

To further validate that our screen specifically selected mutants sensitive to adipic acid and not to the low pH (pH 4.0) conferred by adipic acid, we compared our data with a previously reported chemogenomic screen for pH 4.0 toxicity (Shin et al., 2016). We found no overlap between our data and the 129 genes identified in the low pH screen (Shin et al., 2016) (Table S1), giving a strong indication that the sensitive strains obtained in our screen have been selected for specific sensitivity to adipic acid. Previous work had reported that overexpression of $PDR12$ improved tolerance to adipic acid when overexpressed (Pereira et al., 2019), and our screen identified $pdr12$ cells as being hypersensitive to adipic acid, providing further evidence that the chemogenomic screen is specific to adipic acid.

Gene Ontology (GO) enrichment analysis (Robinson et al., 2002) identified numerous biological processes that were enriched in the screen including the ergosterol biosynthesis pathway ($p = 1.147 \times 10^{-3}$ (Table S1; Figure 1). Indeed, strains with an $ERG2$, $ERG3$, or $ERG4$ deletion were hypersensitive to adipic acid and showed no growth in the presence of 120 mM adipic acid, suggesting that ergosterol biosynthesis is required for growth in adipic acid (Figure S1). The most represented biological process that modulates
adipic acid toxicity in our screen is protein transport and trafficking (p = 3.80 × 10^{−11}; Table S1 and Figure 1). In particular, genes encoding proteins within the ESCRT pathway such as STP22-VPS29 (ESCRT-I), VPS25-SNF8-VPS36 (ESCRT-II), and SNF7 (ESCRT-III) were essential for growth in the presence of 120 mM adipic acid. Indeed, all these ESCRT pathway mutants failed to grow in the presence of 120 mM adipic acid. Deletion of Bro1, an auxiliary protein in the ESCRT pathway, which binds to SNF7 (Wemmer et al., 2011), also results in sensitivity to adipic acid. Furthermore, deleting genes encoding protein complexes such as VPS51-VPS53 (GARP complex) and COG5-COG6-COG7 (COG complex) results in significant growth inhibition in the presence of adipic acid (Figure 1). Taken together, ergosterol biosynthesis and protein trafficking are essential biological processes required for yeast growth in adipic acid as deletion of key genes in these processes results in hypersensitivity to adipic acid.

**KGD1 in the tricarboxylic acid cycle suppresses adipic acid toxicity**

The only adipic acid suppressor mutant identified was KGD1, which encodes an enzyme (α-ketoglutarate dehydrogenase) that converts α-ketoglutarate to oxaloacetate (Repetto and Tzagoloff, 1989) in the tricarboxylic acid (TCA) cycle (Figure 2A). We confirmed that the kgd1Δ strain grew better than the wild type during exposure to adipic acid in both liquid and solid YPD media (Figures 2B and S1A). Through liquid growth assays, we next investigated the involvement of other genes in the TCA cycle in modulating adipic acid tolerance. Apart from sdh1Δ, which is hypersensitive to adipic acid, all the other TCA cycle mutants grew similarly as the wild-type strain in adipic acid (Figure S2B).

Next, we asked if deleting KGD1 also improved tolerance to catechol and cis,cis-muconic acid, two precursors in the adipic acid biosynthetic pathway. Therefore, we calculated yeast growth rates in liquid cultures containing 12 mM catechol or 50 mM cis,cis-muconic acid and compared it with that grown in YPD without any chemicals added. Although the kgd1Δ strain grew better than the wild type in media containing catechol (p < 0.05), the mutant strain grew similarly as the wild-type strain in media containing cis,cis-muconic acid (Figure 2C), suggesting that KGD1 deletion confers cross-resistance to some precursors in the adipic acid biosynthesis pathway. In summary, KGD1, and not the TCA cycle, modulates adipic acid tolerance in yeast. Furthermore, deletion of KGD1 also improved tolerance to catechol, a toxic adipic acid precursor.

**Genes encoding enzymes in the last four steps of the ergosterol biosynthesis pathway are required for growth in adipic acid**

To determine whether the ergosterol biosynthesis pathway (Figure 3A) modulates adipic acid toxicity, using liquid growth analysis we tested the sensitivity of all nine non-essential ergosterol pathway deletion mutants to adipic acid (HMG1, HMG2, ERG24, ERG6, ERG28, ERG2, ERG3, ERG5, ERG4) (Bhattacharya et al., 2018). These included the last four genes in the ergosterol biosynthesis pathway that were identified in our chemogenomic screen (ERG2, ERG3, ERG5, and ERG4, Figure 1). Similar to our chemogenomic screen, whereas erg5Δ cells display a mild sensitivity to adipic acid, erg2Δ, erg3Δ, and erg4Δ cells showed a complete growth inhibition in media containing 120 mM adipic acid (Figure 3B). However, whereas hmg2Δ cells have minor sensitivity to adipic acid at higher OD, the other non-essential ergosterol pathway deletion mutants cells’ growth profile was similar to those of wild-type cells upon 120 mM adipic acid treatment (Figure 3B).

In order to delve deeper into the role of the ergosterol pathway in modulating adipic acid toxicity, including the essential pathway proteins, we next asked if 2 h of adipic acid treatment impacted the protein levels and/or subcellular localization of ergosterol pathway enzymes tagged with a green fluorescent protein (GFP). We did not observe any significant changes in protein level of any of the GFP-tagged ergosterol pathway enzymes upon adipic acid treatment (Figures S3 and S4). This suggests that even though the ergosterol biosynthesis pathway, particularly the last four steps, is required for growth in adipic acid, the pathway is not induced in the presence of the compound. However, two of the enzymes in the pathway changed localization upon adipic acid treatment. The endoplasmic reticulum (ER) localized Erg24-GFP (Jordá and Puig, 2020) and Erg4-GFP (Srinivas et al., 2019) moved to the cytosol upon adipic acid treatment (Figure 4A). Erg4-GFP also changed localization to punctate structures (Figure 4A). As Erg4 is the last enzyme in the pathway leading to the formation of ergosterol and its deletion results in hypersensitivity to adipic acid, we did a time course experiment to track changes in localization of Erg4-GFP. To this end, we grew cells expressing Erg4-GFP to log phase in YPD before adding 120 mM adipic acid, and localization was assessed at 0, 15, 30, and 60 min. We found that Erg4-GFP changed localization to punctate structures as early as 15 min of adipic acid exposure (Figure S5). As ergosterol is a major component of
Figure 2. KGD1 deletion suppresses adipic acid toxicity in yeast

(A) Schematic of the tricarboxylic acid (TCA) cycle.

(B) Dot assays were performed by spotting 4-fold serial dilutions of mid-log phase wild-type and kgd1Δ (YKB4943) cultures onto YPD agar plates supplemented with increasing concentrations of adipic acid and incubated at 30°C for 48 h. The images are representative of three biological replicates.

(C) Wild-type (WT) and kgd1Δ (YKB4943) strains were grown to the mid-log phase; diluted to an OD600 of 0.1 in liquid YPD cultures with or without 12 mM catechol, 50 mM muconic acid, or 140 mM adipic acid; and automated growth curves were obtained in triplicates using the Bioscreen at 30°C. Growth rates of kgd1Δ were calculated for each treatment and normalized to that of the WT. Error bars represent 1 standard deviation. *p < 0.05.

See also Figure S2.
mitochondrial membranes (Krumpe et al., 2012) and decreases in pH can cause fragmentation of mitochondria (Pereira et al., 2010), using the mitochondrial marker Cit1-RFP, we asked if Erg4-GFP was relocalizing to mitochondria upon adipic acid treatment. As expected, in untreated cells, the ER-localized Erg4-GFP did not co-localize with Cit1-RFP (Figure 4B). Adipic acid treatment caused fragmentation of the mitochondria, and Erg4-GFP co-localized with Cit1-RFP. Together, these data suggest that the last four steps in the ergosterol biosynthesis pathway that convert fecosterol to ergosterol, although not induced during adipic acid treatment, are required for yeast growth during exposure to adipic acid. Indeed, key enzymes such as Hmg1, which catalyzes a rate-limiting step of the pathway (Bhattacharya et al., 2018), and Erg4, which catalyzes the final step in the pathway, change localization upon adipic acid treatment, suggesting a potential regulation of the pathway during growth in adipic acid.

**Ergosterol and Pdr12 efflux play non-overlapping roles protecting the cell from adipic acid toxicity**

Plasma membrane ergosterol has been implicated in the localization and activity of many plasma membrane transporters (Bagnat et al., 2000; Kodedová and Syčrová, 2015). Hence, we next sought to determine if the hypersensitivity of ergosterol biosynthetic pathway mutants to adipic acid reflected defects in mitochondrial membranes (Krumpe et al., 2012) and decreases in pH can cause fragmentation of mitochondria (Pereira et al., 2010), using the mitochondrial marker Cit1-RFP, we asked if Erg4-GFP was relocalizing to mitochondria upon adipic acid treatment. As expected, in untreated cells, the ER-localized Erg4-GFP did not co-localize with Cit1-RFP (Figure 4B). Adipic acid treatment caused fragmentation of the mitochondria, and Erg4-GFP co-localized with Cit1-RFP. Together, these data suggest that the last four steps in the ergosterol biosynthesis pathway that convert fecosterol to ergosterol, although not induced during adipic acid treatment, are required for yeast growth during exposure to adipic acid. Indeed, key enzymes such as Hmg1, which catalyzes a rate-limiting step of the pathway (Bhattacharya et al., 2018), and Erg4, which catalyzes the final step in the pathway, change localization upon adipic acid treatment, suggesting a potential regulation of the pathway during growth in adipic acid.

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in Pdr12, the only plasma membrane transporter identified in our chemogenomic screen (Figures 1 and S6). As the overexpression of PDR12 improves adipic acid tolerance (Pereira et al., 2019), we first asked if adipic acid treatment changed the protein levels or localization of Pdr12. We C terminally tagged endogenous PDR12 with GFP (Pdr12-GFP) and assessed the impact of adipic acid treatment on protein levels and localization. Upon 120 mM adipic acid exposure, there was a rapid induction of Pdr12-GFP, which localized to the plasma membrane (Figure 5A). The induction of the protein was observed as early as 15 min after exposure to adipic acid (Figure 5A). Using quantitative western blots we found that Pdr12-GFP protein

Figure 4. Ergosterol biosynthetic enzymes Hmg1, Erg4, and Erg24 change localization upon adipic acid treatment
(A) Cells containing GFP fusions to Hmg1 (YKB4947), Erg4 (YKB5088), and Erg24 (YKB4951) were grown to mid-log phase, centrifuged, and resuspended in YPD media with or without adipic acid and grown at 30°C for 2 h and imaged. (B) Adipic acid treatment causes mitochondrial fragmentation and Erg4-GFP relocalization to the mitochondrial marker Cit1-RFP. Cells endogenously expressing Erg4-GFP and Cit1-RFP (YKB5091) were grown to mid-log phase, centrifuged, and resuspended in YPD media with or without adipic acid and grown at 30°C for 2 h before being imaged. Scale bar, 10 μm. Images are representative of three biological replicates. See also Figures S3–S5.
Figure 5. Adipic acid induction and plasma membrane localization of Pdr12 is not dependent on ergosterol

(A and B) Pdr12-GFP is induced upon adipic acid treatment. (A) Wild-type cells expressing endogenously tagged Pdr12-GFP (YKB4889) were grown to mid-log phase, centrifuged and resuspended in YPD media with or without adipic acid, grown at 30°C, and images taken at the indicated time points. (B) Protein levels of Pdr12-GFP in YPD with and without adipic acid treatment for 30 min were quantitated using western blots. *Statistical significance at p < 0.05. Error bars represent 1 SD.

(C) Deletion of ERG2 and ERG4 does not impact the adipic acid induction and plasma membrane localization of Pdr12-GFP. Wild-type (WT), erg2Δ, and erg4Δ strains expressing endogenously tagged Pdr12-GFP were grown to mid-log phase, centrifuged and resuspended in YPD media with or without adipic acid, and grown at 30°C for 1 h before imaging. Scale bar, 10 μm.

(D) Synergistic effects of erg4Δ and pdr12Δ on adipic acid toxicity. Wild-type (WT), erg4Δ (YKB4876), pdr12Δ (YKB4383), and erg4Δpdr12Δ (YKB5036) were grown to mid-log phase in YPD medium before being diluted into fresh medium to an OD₆₀₀ of 0.1. Four 10-fold serial dilutions were spotted onto YPD agar plates with or without adipic acid (60 mM) and incubated at 30°C for 48 h. The image is representative of three biological replicates.

See also Figure S6.
As ergosterol has been reported to play a role as lipid raft in transporting newly synthesized proteins from the Golgi to the plasma membrane (Bastos et al., 2012; Mollinedo, 2012), we asked if ergosterol biosynthesis genes ERG2 and ERG4 were required for either the induction or localization of Pdr12-GFP to the plasma membrane upon adipic acid treatment. Wild-type, erg2Δ, and erg4Δ cells expressing endogenously tagged Pdr12-GFP were grown to midlog phase before being treated with 120 mM adipic acid for 1 h. We found that Pdr12-GFP in the ergosterol mutants was induced and localized to the plasma membrane similar to wild type, suggesting that Pdr12 localization to the plasma membrane is independent of ergosterol lipid rafts (Figure 5C). Although ergosterol may not be needed to localize Pdr12 to the plasma membrane, it may still be required for Pdr12 activity. If this was the case we would hypothesize that an erg4Δpdr12Δ sensitivity to adipic acid would be epistatic or similar to the single mutants. If, however, these two pathways were working in parallel, the double mutant would have increased sensitivity to adipic acid compared with the single mutants. Wild-type, erg4Δ, pdr12Δ, and erg4Δpdr12Δ cells were tested for growth on YPD agar plates with and without a sublethal dosage of adipic acid (Figure 5D). Although 60 mM adipic acid treatment causes slow growth of erg4Δ, it has limited impact on pdr12Δ cells. In contrast, the ability of erg4Δpdr12Δ cells to grow on adipic acid is dramatically slower than either erg4Δ or pdr12Δ cells. The synergistic sensitivity to adipic acid suggests that ergosterol and Pdr12 play independent roles in protecting the cells from adipic acid toxicity (Figure 5D). Taken together, these data suggest that although Pdr12 is rapidly induced upon adipic acid treatment, its localization to the plasma membrane is not mediated by ergosterol lipid rafts. Furthermore, our genetics suggests that Erg4, and presumably the ergosterol biosynthesis pathway, and Pdr12 play non-overlapping roles in buffering adipic acid toxicity.

Adipic acid disrupts the Membrane Compartment of Can1 (MCC) and induces endocytosis

To further probe the potential roles of ergosterol in adipic acid protection, we asked if the ergosterol-rich MCC was affected by adipic acid. MCCs are membrane microdomains and occur as one of several large membrane compartments on the plasma membrane (Malinsky et al., 2010). They also serve as membrane reservoirs of specific proteins such as the proton symports Can1, Tat2, and Fur4 as well as other proteins with unknown functions including Sur7, Fmp45, and Yn1194c (Douglas et al., 2011). Can1 is an arginine transporter that interacts with ergosterol in the MCC and also serves as an MCC marker (Grossmann et al., 2008). Not only is ergosterol required to target some of these membrane proteins including Can1 and Tat2 to the MCC but also ergosterol accumulates in the MCC, which is visualized as patches when cells are stained with the filipin dye (Grossmann et al., 2007; Malínska et al., 2003). Indeed, deletion of ergosterol biosynthesis genes has been shown to impair MCC formation (Grossmann et al., 2006). To assess the effect of adipic acid on the MCC, wild-type and erg4Δ cells were stained with filipin before treatment with adipic acid. As expected in untreated wild-type cells, filipin staining of the plasma membrane appeared as punctate structures or MCC patches, whereas in erg4Δ cells punctate filipin staining is disrupted (Figure 6A). Upon adipic acid treatment, the filipin-stained MCC patches on the plasma membrane were lost in wild-type cells (Figure 6A), suggesting that adipic acid impacts the plasma membrane MCC. As a secondary method to assess the impact of adipic acid on the MCC, we utilized the MCC marker Can1-GFP, and as expected during growth in adipic acid, Can1-GFP is lost from the plasma membrane and is enriched in punctate internal structures (Figures 6B and S7). Previous studies have reported that Can1-GFP undergoes endocytosis to the vacuole when it is lost from the plasma membrane (Ghaddar et al., 2014; Gournas et al., 2017). Using two vacuolar markers, CMAC and FM4-64, which stain the lumen of the vacuole and the vacuolar membrane, respectively, we determined that Can1-GFP localized to the vacuoles upon adipic acid treatment (Figures 6B and S7). Furthermore, upon adipic acid treatment there is a significant decrease in full-length Can1-GFP protein levels and an increase in free GFP (Figure 6C). Together this suggests that adipic acid exposure disperses Can1 from the MCC, which is subsequently internalized by endocytosis to the vacuole for degradation.

As the MCCs are sites for endocytosis on the plasma membrane, we hypothesized that adipic acid treatment induced endocytosis as a mechanism to potentially downregulate membrane transporters that import adipic acid into the cell, which results in accumulation and toxicity. To test this hypothesis, we utilized the vacuole dye FM4-64, which is also commonly used to measure endocytosis (Brach et al., 2014). We performed a time course experiment and determined that adipic acid-treated cells internalized FM4-64 levels were induced 12-fold after 30 min of adipic acid treatment (Figure 5B). These data indicate that Pdr12 expression is increased during growth in adipic acid potentially to facilitate export of cytosolic adipic acid anions (adipate) in order to reduce intracellular toxicity.
Figure 6. Adipic acid disrupts the MCC and induces endocytosis

(A) Adipic acid treatment disrupts ergosterol plasma membrane patches. Mid-log phase cultures of wild-type (WT, YKB1079) and erg4Δ (YKB4876) cells were stained with the filipin dye and treated with or without 120 mM adipic acid for 1 h, before imaging. Scale bar, 10 μm; images are representative of three biological replicates.

(B) Adipic acid treatment results in Can1-GFP relocalization to the vacuole. Wild-type cells endogenously expressing Can1-GFP (YKB5097) were grown to the mid-log phase in YPD media. Cells were then stained with the vacuole lumen dye CMAC for 15 min before being treated with 120 mM adipic acid for 1 h at 30°C after which they were imaged. Scale bar, 10 μm; images are representative of three biological replicates.

(C) Can1-GFP is degraded upon adipic acid treatment. Quantitative western blot analysis were performed on whole-cell protein extracts, which had been collected by TCA extraction from Can1-GFP yeast cultures that were untreated (0 mM) or treated with adipic acid (120 mM) for 1 h. The anti-GFP blot is shown with the corresponding total protein for each lane as assessed using the Bio-Rad TGX system (left panel). The abundance of full-length Can1-GFP protein and free GFP was quantified relative to total protein and normalized to untreated levels (fold expression) (right panels). n = 3, *p < 0.05, error bars represent 1 SD.

(D) Adipic acid treatment induces endocytosis. Mid-log phase cultures of the wild-type yeast (YKB1079) were stained with the FM4-64 and treated with or without adipic acid. Samples from the cultures were taken at 5, 30, 60, and 120 min after adipic acid treatment. The cells were centrifuged, resuspended in SC medium, and imaged. Scale bar, 10 μm; images are representative of three biological replicates.

See also Figure S7.
faster than untreated cells (Figure 6D). Remarkably, instead of diffused plasma membrane staining at time point 0 min, FM4-64 is already being internalized in adipic acid-treated cells, and vacuolar staining of cells by FM4-64 was visible after 15 min of incubation with adipic acid, whereas it took up to 120 min to observe clear vacuolar staining in the untreated cells. Together, these data indicate that adipic acid disrupts the membrane component of Can1 potentially by an increased rate of endocytosis.

DISCUSSION

Overcoming the challenge of adipic acid toxicity in yeast is imperative to achieve high titers of the acid in production through yeast fermentation. Therefore, in this study we sought to identify the mechanisms underlying the toxic effect of the adipic acid in yeast and identify genes that could be targeted to improve tolerance in yeast. Toward this goal, we performed a chemogenomic screen and systematically assessed the impact of each non-essential gene on adipic acid tolerance. Our screen revealed that a disruption in protein trafficking and the ergosterol biosynthesis pathway resulted in significant increase in sensitivity to adipic acid. On the contrary, deleting *KGD1*, a gene in the TCA cycle, resulted in improved tolerance to adipic acid.

How is *KGD1* associated with enhanced tolerance to adipic acid?

In our screen, we identified one suppressor gene whose deletion resulted in tolerance to adipic acid. To the best of our knowledge, this is the first study reporting the role of the TCA cycle gene, *KGD1*, in increasing tolerance to adipic acid and its precursor catechol (Figure 2). Surprisingly, only the deletion of *KGD1*, but not other genes in the TCA cycle, improved adipic acid tolerance (Figures 2 and S2). How is this occurring? One possibility is that deleting *KGD1* results in the redirected metabolic flux toward acetyl-CoA production and this resulted in high amounts of acetate in the cell (Gruchatka and Kayser, 2015). Although we did not quantify intracellular levels of acetate in our *kgd1Δ* strain, future work should investigate if priming yeast cells with acetic acid before adipic acid exposure results in increased tolerance. This investigation will be worthwhile because acetic acid has also been shown in earlier reports to activate the plasma membrane transporter Pdr12 (Kim et al., 2019; Mira et al., 2010a). Given that Pdr12 overexpression improved tolerance to adipic acid (Pereira et al., 2019), intracellular levels of acetate in *kgd1Δ* strains may also result in Pdr12 induction and increased tolerance to adipic acid. If this was the case, however, one would anticipate that other mutants in the TCA cycle that increase cellular acetate may also confer adipic acid tolerance. However, these were not identified in the screen, potentially due to functional redundancy. Alternatively, Kgd1 may have biological functions outside of the TCA cycle. This is potentially the case because although *KGD1* is a component of the mitochondrial alpha-ketoglutarate dehydrogenase complex together with *KGD2* (Heublein et al., 2014), only *KGD1* is associated with adipic acid tolerance. It will be interesting to further investigate why deleting *KGD1* suppresses adipic acid toxicity.

How does ergosterol modulate adipic acid toxicity?

Ergosterol and other yeast sterols have been reported to modulate different stresses in yeast (Caspeta et al., 2014; Fletcher et al., 2017; Liu et al., 2017). Interestingly, in the present study we establish that cellular ergosterol also plays a role in adipic acid tolerance. Here, we show that disrupting the final steps of the ergosterol biosynthetic pathway results in severe growth defects in yeast upon adipic acid treatment (Figures 3B and S1). It is likely that ergosterol mutants lacking genes in the final steps of the pathway have variations in their sterol composition. Previously, a study reported that yeast mutants lacking enzymes coded for by these ergosterol pathway genes accumulate a mixture of sterols because Erg2, Erg3, Erg4, and Erg5, which catalyze the last steps of the pathway, are promiscuous in their choice of substrate (Abe and Hiraki, 2009). Indeed, erg2Δ has been shown to accumulate fecosterol (Erg2 precursor) as well as ergosta-5,8,12-trienol and ergosta-7-enol, whereas erg3Δ accumulates a mixture of episterol, ergosta-7,22-dienol, and ergosta-7-enol (Abe and Hiraki, 2009; Johnston et al., 2020). On the other hand, erg5Δ accumulates ergosta-5,7,24(28)-tri-enol and ergosta-5,7-dienol, whereas erg4Δ mutant accumulates ergosta-5,7,22,24(28)-tetraen-3-ol (ergosterol precursor) (Abe and Hiraki, 2009). These sterol precursors will impact the physical properties of the cell membrane in a way different from the wild-type membrane, which is made up of ergosterol as the predominant sterol. Ergosterol is a flat sterol, whereas sterol precursors such as fecosterol and episterol are bent (Caspeta et al., 2014; Shahedi et al., 2006) and will alter the cell membrane by interfering with the van der Waals’s interactions within the membrane’s lipid bilayer (Grossmann et al., 2007). In the context of protecting cells from adipic acid toxicity, it is possible that the membrane structure maintained by ergosterol limits the rate at which the undissociated adipic acid diffuses across the plasma membrane into the cytosol from the extracellular environment.
As mentioned earlier, maintaining the right amount and composition of sterol is essential for growth in adipic acid. Changes in subcellular localization of sterol enzymes occur in cells as a way of regulating the activity of these enzymes (Leber et al., 1998). This might be the case for Erg24 and Erg4 during growth in adipic acid (Figure 4A) particularly because Erg4 catalyzes the final step of the pathway required for ergosterol biosynthesis. Alternatively, and not mutually exclusive, membrane sterol composition may be playing a critical role in the activity and/or localization of membrane-bound proteins. For example the activity of Pdr5, an ABC transporter reported to increase yeast tolerance to certain stresses, is impaired in ergosterol biosynthesis mutants (Kodedová and Sychrová, 2015). Similarly, ergosterol has been shown to be required for Pdr12 function, which is essential for tolerance to organic acids (Xu and Li, 2020). Yeast mutants that lack genes in the ergosterol biosynthesis pathway are sensitive not only to adipic acid but also to other organic acids such as sorbic acid (Mollapour et al., 2004). Although we did not investigate whether the efflux activity of Pdr12 is also significantly impaired in ergosterol mutants, we showed that deleting Pdr12 in an ergosterol mutant resulted in a synergistic effect on growth in adipic acid. The erg4Δpdr12Δ mutant was more sensitive to adipic acid than the erg4Δ or pdr12Δ alone (Figure 5D). This observation suggested that ergosterol plays other roles in modulating adipic acid toxicity beyond a potential role in regulating Pdr12. Regarding its role in the localization of membrane-bound proteins, ergosterol transports and targets membrane proteins such as Can1 and Tat2 to the MCC via ergosterol lipid rafts (Grossmann et al., 2008). Furthermore, ergosterol along with several proteins forms the MCC, which are sites of endocytosis on the plasma membrane, and defects in MCC formation results in impaired endocytosis (Heese-Peck et al., 2002). From our data, adipic acid impacts endocytosis by disrupting MCC formation, which was evidenced in the rapid internalization of Can1-GFP and FM4-64 (Figure 6D).

**Why is protein trafficking and vacuolar transport essential for yeast growth in adipic acid?**

Nearly half of the genes in the adipic acid chemogenomic profile are associated with protein trafficking and vacuolar transport, suggesting that protein degradation and recycling in the vacuole is an essential biological process required for survival during yeast growth in adipic acid. Interestingly, most of these genes code for proteins that are components of the complexes that form the Endosomal Sorting Complexes Required for Transport (ESCRT) machinery. The ESCRT pathway is used by cells to degrade specific proteins on the plasma membrane by endocytosis (Wenzel et al., 2018). Intriguingly, our study shows that adipic acid impacts and disrupts the MCC (Figure 6A), which serves as a site for endocytosis and also provides a protective surrounding for several plasma membrane proteins and transporters including Can1 permease. A previous study has shown that when the protective area offered by the MCC is disrupted, proteins that reside in this compartment are released and become susceptible to cellular internalization and endocytosis (Grossmann et al., 2007, 2008). This explains why when adipic acid disrupts the MCC, Can1 rapidly changes localization to the vacuole for degradation (Figures 6B and 6C). This explanation was independently strengthened by the fact that adipic acid induced endocytosis (Figure 6D).

Endocytosis is essential for degrading and recycling of plasma membrane proteins. Specific membrane proteins are ubiquitinylated, internalized by endocytosis, and transported to the endosomes via the ESCRT pathway for vacuolar degradation (Feyder et al., 2015). Although organic acids are generally known to enter cells by passive diffusion, it has been shown that some plasma membrane transporters such as Jen1 and Acy1 are able to transport lactate, acetate, and other organic acids into the cell (Casal et al., 2008; McDermott et al., 2010). We speculate that endocytosis is induced in adipic acid-treated cells as a way of degrading non-specific membrane transporters that import extracellular adipic acid into the cell in order to prevent intracellular accumulation of the acid. It will be interesting to screen and identify non-specific transporters that import adipic acid into the cell and determine how adipic acid treatment impacts their degradation by endocytosis. Furthermore, as endocytosis is induced in adipic acid-treated cells as a survival mechanism, it is imperative that all the proteins involved in this process such as the ESCRT pathway proteins are maintained. Hence, when any of these key proteins are disrupted by deleting the genes that encode them, the cells become hypersensitive to adipic acid as shown in the adipic acid chemogenomic profile in this study.

**Conclusion**

In conclusion, our study provides an outlook into the basic cellular mechanisms underlying adipic acid toxicity in yeast and provides potential genetic candidates that can be engineered for tolerance to boost adipic acid production in yeast. The results presented in this study highlight an induction of endocytosis...
in cells treated with adipic acid. It is therefore essential to maintain ergosterol biosynthesis because ergosterol is required to maintain the integrity of the plasma membrane as well as the MCC needed for endocytosis. Furthermore, KGD1 deletion suppresses the toxicity of adipic acid and its toxic catechol precursor, making this gene an important target for engineering robust adipic acid production strains.

**Limitations of the study**

The present study demonstrated an important role for the last four steps of the ergosterol biosynthesis pathway in protecting yeast cells from the toxic effects of adipic acid. Further studies will be required to determine if modulating the ergosterol pathway, such as overexpression, can improve growth upon adipic acid exposure. In addition, as this work was conducted in the laboratory yeast strain background S288c further studies are required to determine if the findings presented here are transferrable to industrial yeast strains.

**Resource availability**

**Lead contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Kristin Baetz (kbaetz@uottawa.ca).

**Materials availability**

This study did not generate new unique reagents. Yeast strains generated in this study that are not available commercially are available from the lead contact without restriction with the requestor paying for shipping costs.

**Data and code availability**

The published article includes all datasets generated or analyzed during this study.

**METHODS**

All methods can be found in the accompanying transparent methods supplemental file.

**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.102327.

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**AUTHOR CONTRIBUTIONS**

E.F. and K.B. conceptualized the project. E.F., K.M., and E.A.W. performed the experiments. E.F. and K.B. prepared the manuscript. K.B. acquired the funds. All authors approved the final version of the manuscript.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

**INCLUSION AND DIVERSITY**

One or more of the authors of this paper self-identifies as an underrepresented ethnic minority in science. One or more of the authors of this paper self-identifies as living with a disability.

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Supplemental information

A yeast chemogenomic screen identifies pathways that modulate adipic acid toxicity

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Supplemental Information

Figure S1: Deletion of genes in the late stage of the ergosterol biosynthesis pathway result in hypersensitivity to adipic acid, related to Figures 1 and 3. Wild type (WT), \( \text{erg2}\Delta \), \( \text{erg3}\Delta \) and \( \text{erg4}\Delta \) cultures were grown to the mid-log phase, diluted to an \( \text{OD}_{600} \) of 0.1 in liquid cultures with and without adipic acid and grown at 30 °C using a BioScreen. The automated growth curves were performed in triplicates.
Figure S2: KGD1 deletion suppresses adipic acid toxicity while other genes encoding TCA cycle enzymes do not modulate adipic acid toxicity, related to Figure 2. (A) WT and kgd1Δ cultures were grown to the mid-log phase, diluted to an OD$_{600}$ of 0.1 in liquid cultures with or without adipic acid and grown at 30°C using a BioScreen. (B) WT, aco1Δ (YKB5099), aco2Δ (YKB5100), cit1Δ (YKB5101), fum1Δ (YKB5102), idh1Δ (YKB5103), idh2Δ (YKB5104), kgd2Δ (YKB5105), lsc1Δ (YKB5106), lsc2Δ (YKB5107), mdh1Δ (YKB5108), pdc1Δ (YKB5109), sdh1Δ (YKB5110), sdh2Δ (YKB5111) and sdh4Δ (YKB5112) cultures were grown to the mid-log phase, diluted to an OD$_{600}$ of 0.1 in liquid cultures with or without adipic acid and grown at 30°C using a BioScreen. The automated growth curves were performed in triplicates.
Figure S3: Ergosterol biosynthesis enzymes are not induced upon adipic acid treatment, related to Figure 4. Wild type (WT) cells expressing GFP-tagged ergosterol biosynthesis enzymes were grown to mid-log phase, centrifuged and resuspended in YPD media with or without adipic acid and grown at 30 °C. Green fluorescence of the cultures was measured after incubation for 2 hours.
Figure S4: Ergosterol biosynthesis enzymes are not induced upon adipic acid treatment, related to Figure 4. Wild type (WT) cells expressing GFP-tagged ergosterol biosynthesis enzymes were grown to mid-log phase, centrifuged and resuspended in YPD media with or without adipic acid and grown at 30 °C. Green fluorescence of the cultures was measured after incubation for 2 hours. A minimum of 50 cells were captured for each image. Fluorescence intensity from cells were quantified using the ImageJ software and this was normalized to the untreated cultures for each strain tested. *n.s.* represents “not significant”. Error bars represent 1 STD.
Figure S5: Ergosterol biosynthesis enzyme Erg4 changes localization over time upon adipic acid treatment, related to Figure 4. Cells expressing Erg4-GFP (YKB5088) were grown to mid-log phase, centrifuged and resuspended in YPD media with adipic acid and incubated at 30 °C. Images were taken at the start of the experiment (0 minutes) and after 15, 30 and 60 minutes of incubation with 120 mM adipic acid. The scale bar represents 10 µm. Images are representative of three biological replicates.
Figure S6: *PDR12* deletion increases yeast sensitivity to adipic acid, related to Figure 5. Wild type (WT) and *pdr12Δ* (YKB4383) cultures were grown to the mid-log phase, diluted to an OD$_{600}$ of 0.1 in liquid cultures with and without adipic acid and grown at 30 °C using a BioScreen. The automated growth curves were performed in triplicates.
Figure S7: Can1-GFP changes localization from the plasma membrane to the vacuole upon adipic acid treatment, related to Figure 6. (A) Can1-GFP cells were grown to mid-log phase, centrifuged and resuspended in YPD media with or without adipic acid and grown at 30 °C. (B) The cells were then stained with the FM4-64 dye to visualize co-localization of Can1-GFP with the vacuole. FM4-64 (red) stains the vacuole membrane. Green and red fluorescence of the cultures were measured after incubation for 1 hour.
| Strains              | Auxotrophies                                      | Ref. or source |
|----------------------|--------------------------------------------------|----------------|
| BY4741, YKB1079      | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0                | Open Biosystems |
| YKB4943              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 kgd1Δ::KANMX    | Open Biosystems |
| YKB5099              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 aco1Δ::KANMX    | Open Biosystems |
| YKB5100              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 aco2Δ::KANMX    | Open Biosystems |
| YKB5101              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 cit1Δ::KANMX    | Open Biosystems |
| YKB5102              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 fum1Δ::KANMX    | Open Biosystems |
| YKB5103              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 idh1Δ::KANMX    | Open Biosystems |
| YKB5104              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 idh2Δ::KANMX    | Open Biosystems |
| YKB5105              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 kgd2Δ::KANMX    | Open Biosystems |
| YKB5106              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 lsc1Δ::KANMX    | Open Biosystems |
| YKB5107              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 lsc2Δ::KANMX    | Open Biosystems |
| YKB5108              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 mdh1Δ::KANMX    | Open Biosystems |
| YKB5109              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 pdc1Δ::KANMX    | Open Biosystems |
| YKB5110              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 sdh1Δ::KANMX    | Open Biosystems |
| YKB5111              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 sdh2Δ::KANMX    | Open Biosystems |
| YKB5112              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 sdh4Δ::KANMX    | Open Biosystems |
| YKB4956              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 hmg1Δ::KANMX    | Open Biosystems |
| YKB4957              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 hmg2Δ::KANMX    | Open Biosystems |
| YKB4958              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 erg2Δ::KANMX    | Open Biosystems |
| YKB4959              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 erg6Δ::KANMX    | Open Biosystems |
| YKB4960              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 erg28Δ::KANMX   | Open Biosystems |
| YKB4874              | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 erg2Δ::KANMX    | Open Biosystems |
| Strain   | Phenotype                  | Source                        |
|----------|----------------------------|-------------------------------|
| YKB4875  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 erg3Δ::KANMX | Open Biosystems               |
| YKB4877  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 erg5Δ::KANMX | Open Biosystems               |
| YKB4876  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 erg4Δ::KANMX | Open Biosystems               |
| YKB5079  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG10-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB4947  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 HMG1-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB4948  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 HMG2-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5080  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG12-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB4950  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG8-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5081  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 IDI1-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5082  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG9-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5098  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG19-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB4949  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG1-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5083  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG7-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5084  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG11-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB4951  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG24-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5113  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG25-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5085  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG26-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5086  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG27-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB4952  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG6-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5087  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG3-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5088  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG4-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5089  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG5-GFP::HISMX | (Ghaemmaghami et al., 2003)   |
| YKB5091  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0 ERG4-GFP::HISMX CIT1-RFP ::URA | This study                    |
| Strain   | Genetic modifications                                                                 | Source               |
|----------|---------------------------------------------------------------------------------------|----------------------|
| YKB4383  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0  
  pdr12Δ::KANMX                           | Open Biosystems      |
| YKB5036  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0  
  erg4Δ::KANMX  
  pdr12Δ::NATMX                         | This study           |
| YKB4889  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0  
  erg4Δ::KANMX  
  PDR12-GFP::HISMX                      | (Ghaemmaghami et al., 2003) |
| YKB4954  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0  
  erg2Δ::KANMX  
  PDR12-GFP::HISMX                      | This study           |
| YKB4955  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0  
  erg4Δ::KANMX  
  PDR12-GFP::HISMX                      | This study           |
| YKB5097  | MATa his3Δ1 leu2Δ0 met15Δ0 ura3Δ0  
  Can1::GFP::HISMX                      | (Ghaemmaghami et al., 2003) |

**Transparent Methods**

**Strains and plasmids**

The yeast strains used in this study are listed in Table S2. These yeast strains are derivatives of the haploid BY4741 strain. Strains with GFP tagged proteins were obtained from the GFP collection (Ghaemmaghami et al., 2003) and deletion mutants were obtained from the deletion mutant array collection (Open Biosystems). Strains that were in neither collections were generated using established methods (Longtine et al., 1998). All strains used in this study were PCR confirmed.

**Chemogenomic screen and analysis**

The chemogenomic screen was performed as previously described (Fletcher et al., 2019) but with a few modifications. Briefly, MATa yeast deletion mutant array (DMA) collection (~4200 mutants) was arrayed in duplicate and condensed at density of 1536 colonies per plate on YPD agar (1% w/v Yeast Extract, 2% w/v Bacto peptone, 2% w/v agar, 0.03% w/v Tryptophan and 2% w/v Glucose) plates supplemented with G418 using a Singer RoToR HDA (Singer Instruments). The condensed arrays were pinned onto YPD agar plates containing 80 mM adipic acid (Sigma-Aldrich; cat. #09582-250G) dissolved in DMSO. The resulting pH of the plates due to the adipic acid was 4.0. A control screen was performed at the same time where the condensed arrays were pinned onto YPD agar plates containing DMSO and the pH of the plates buffered at pH 4.0. Both the adipic acid and the control screens were performed in triplicate and the plates were incubated at 30 °C for 48 h after which images of each plate were taken for growth assessment of each colony using the SGAtools (http://sgatools.ccbr.utoronto.ca/) as previously described (Wagih et al., 2013). An average growth score of less than −0.3 and greater than 0.3 was used as
the cut-off to identify mutants that were sensitive and resistant, respectively, to adipic acid.

Agar-based chemogenomic screen was confirmed by measuring the growth rate of the identified mutants in liquid YPD medium supplemented with 120 mM adipic acid. Briefly, overnight cultures of the mutant strains were inoculated into fresh YPD medium and strains were incubated at 30 °C until they reached the mid log phase (OD_{600} 0.5–0.6), prior to being diluted to a final OD_{600} of 0.1 in a BioScreen C™ Honeycomb microplate. The growth assays were performed for each mutant in triplicate in YPD control media or YPD media supplemented with 120 mM adipic acid. Using a BioScreen C™ plate reader, the cultures were incubated at 30 °C for 72 h and OD_{600} readings were taken every 15 min to plot growth curves. Data from the BioScreen C™ were analyzed with the PRECOG software (Fernandez-Ricaud et al., 2016) to obtain the doubling time of all the strains. A PRECOG score for each of the strains tested was calculated as the ratio of the doubling time in the presence of adipic acid to the doubling time in the absence of adipic acid. Using a cut-off PRECOG score of > 2 and < 0.8 we identified strains that were sensitive or tolerant, respectively, to the adipic acid. The network plot was performed using the statnet package in the R programming software (Handcock et al., 2008).

Gene Ontology (GO) enrichment analysis was performed using the web-based tool (Robinson et al., 2002). The p-value cutoff of 0.01 was used to determine clusters that were enriched.

**Dot assay and growth rate measurements**

Starting with overnight cultures, the wild type and deletion mutant strains were used to inoculate fresh YPD medium to a final OD_{600} of 0.1. The diluted cultures were grown at 30 °C until they reached the mid log growth phase. The cultures at the mid log phase were diluted to an OD_{600} of 0.1 with fresh YPD and four 10-fold serial dilutions of the cultures were spotted on YPD agar plates supplemented with 80 mM adipic acid. Control plates contained no chemical. The YPD agar plates were incubated for 48 hours at 30 °C after which images of the plates were taken with the ChemiDoc XRS Molecular Imaging system (Biorad).

**Fluorescence microscopy**

Overnight cultures grown at 30°C were used to inoculate fresh YPD to a final OD_{600} of 0.1 and allowed to reach mid-log phase prior to imaging. To perform live cell imaging, the cells were briefly centrifuged (800xg for 2 min), followed by re-suspending in a minimal volume of SC medium (67% w/v yeast nitrogen base without amino acids, 0.2% w/v amino acid drop out mix, 2% w/v glucose). An aliquot of 5 µl was spotted onto glass slides, covered with a cover slip prior to imaging. Images were acquired using either a Leica DMI 6000 florescent microscope (Leica Microsystems GmbH, Wetzler Germany), equipped with a Sutter DG4 light source (Sutter Instruments, California, USA), Ludl emission filter wheel with Chroma band
pass emission filters (Ludl Electronic Products Ltd., NY, USA) and Hamamatsu Orca AG camera (Hamamatsu Photonics, Herrsching am Ammersee, Germany) or an Axio Observer 7 fluorescent microscope was also used to take some images. Images were acquired at 0.2 µm steps across 6 µm using a 63 x oil-immersion objective with a 1.4 numerical aperture.

Image J software was used to quantify fluorescence signal in cells as described by (Schneider et al., 2012) with a slight modification. Briefly, background fluorescence was subtracted from the images and the total fluorescence of all the cells in the image was measured and divided by the number of cells in the image. The images contained a minimum of 50 cells. Normalization was performed by calculating the ratio of fluorescence per cell of treated cultures divided by that of untreated cultures.

Quantitative western blot analysis

Overnight cultures of cells endogenously expressing Pdr12-GFP or Can1-GFP were diluted in fresh YPD medium to a final OD<sub>600</sub> of 0.1. These were grown at 30°C until they reached the early log phase (4 hours). Subsequently, the cells were centrifuged and resuspended in YPD with or without 120 mM adipic acid and were further incubated for 1 hour. To obtain whole cell extracts (WCE), cells were harvested by centrifugation at 3000 rpm for 3 minutes (4 °C). The cell pellets were then washed with chilled sterile water and flash frozen in liquid nitrogen before storage at -80°C. Cell lysis was achieved using the Trichloroacetic acid (TCA) protocol (Kao and Osley, 2003). For the quantitative western blot analysis, cell abundance was equalized using OD<sub>600</sub> prior to TCA lysis. Equal volumes of WCE in SDS loading dye were boiled for 10 min and proteins were separated using a TGX stain-free FastCast acrylamide gel (BioRad cat. # 161–0181). After the gel electrophoresis, the TGX gel was activated using UV (Stain-Free Gel program on the Bio-Rad ChemiDoc XRS Molecular Imaging system) and the proteins were transferred onto a nitrocellulose membrane. The total protein on the blot was imaged using the UV (Stain-Free Blot program on the Bio-Rad ChemiDoc XRS Molecular Imaging system) and the blot was blocked with 5% w/v non-fat milk powder in Tris-buffered saline solution with Tween 20 (TBS-T) for 2 h. After the blocking step, the nitrocellulose blots were incubated overnight with Anti-GFP primary antibody (Sigma-Aldrich # 11814460001) in 5% w/v milk TBS-T after which the nitrocellulose membrane was washed (3 x 10 min) with TBS-T. Next, the nitrocellulose blots were incubated for 1 hour with an HRP conjugated anti-mouse secondary antibody (Bio-Rad) diluted in 5% w/v milk TBS-T and then washed (3 x 10 min) again with TBS-T. In a final step, the nitrocellulose blot was developed with chemiluminiscence reagents (Biorad) and the protein bands were visualized with a ChemiDoc XRS Molecular Imaging system (Biorad). Protein abundance of Pdr12-GFP, Can1-GFP, and Free GFP in untreated versus adipic acid treated samples was quantified using the Image Lab software. This was calculated as the ratio of the Pdr12-GFP, Can1-GFP, or Free GFP band intensity to the intensity of bands in the entire lane signal using the TGX Stain-Free Fast Cast (total proteins). The amount of Pdr12-GFP, Can1-GFP, or Free GFP in the adipic acid treated cultures was then normalized to that of the untreated cultures.
Cellular staining

Starting with overnight cultures, the wild type and deletion mutant strains were used to inoculate fresh YPD medium to a final OD600 of 0.1. The diluted cultures were grown at 30°C until they reached the mid log growth phase. The cultures at the mid log phase were stained with the filipin dye (Sigma-Aldrich) for the visualization of sterols in the plasma membrane. To visualize the vacuolar membrane and lumen the FM4-64FX (Invitrogen) and CMAC dyes were used respectively to stain the cells. For the CMAC experiments, the midlog phase cells were stained with the dye at a final concentration of 25 µM for 15 minutes after which the cells were treated with adipic acid. For the FM4-64 experiments, the dye was added at a final concentration of 0.04 mM at the same time the midlog phase cells were grown with or without adipic acid. The stained cells were incubated for 1-2 hours prior to visualization under a fluorescence microscope.

Statistics

Student t-tests and standard deviations were calculated for at least three experimental replicates using the GraphPad Prism 6.05 software (GraphPad Software Inc., La Jolla, California).

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