Engineered yeast tolerance enables efficient production from toxified lignocellulosic feedstocks

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Lignocellulosic biomass remains unharnessed for the production of renewable fuels and chemicals due to challenges in deconstruction and the toxicity of hydrolysates pose to fermentation microorganisms. Here, we show in *Saccharomyces cerevisiae* that engineered aldehyde reduction and elevated extracellular potassium and pH are sufficient to enable near-parity production between inhibitor-laden and inhibitor-free feedstocks. By specifically targeting the universal hydrolysate inhibitors, a single strain is enhanced to tolerate a broad diversity of highly toxified genuine feedstocks and consistently achieve industrial-scale titers (cellulosic ethanol of >100 grams per liter when toxified). Furthermore, a functionally orthogonal, lightweight design enables seamless transferability to existing metabolically engineered chassis strains: We endow full, multifeedstock tolerance on a xylose-consuming strain and one producing the biodegradable plastics precursor lactic acid. The demonstration of “drop-in” hydrolysate competence enables the potential of cost-effective, at-scale biomass utilization for cellulosic fuel and nonfuel products alike.

INTRODUCTION

Meaningful displacement of greenhouse gas emissions from continued oil consumption requires a renewable feedstock that is transformable into products fungible with petrofuels and petrochemicals and is deployable on a similar scale. Despite the declining cost of carbon-free electricity and rise of emission-free vehicles, studies estimate that this segment will comprise at most 31% of the global transportation sector remains the largest generator of carbon dioxide emissions (1). As the transportation sector remains the largest generator of carbon dioxide, the sheer number of legacy vehicles necessitates that liquid biofuels play a dominant role in any future energy mix to minimize net emissions (2).

Lignocellulosic biomass, the largest renewable terrestrial resource, provides a realistic intermediate-term route to sustainable fuel and nonfuel commodities at enormous scale when paired with suitable fermentation infrastructure (3). In addition to quantities on the magnitude of fossil carbon, lignocellulose addresses issues such as food-fuel competition and arable land use that beset present usage, and pretreatment research has refocused on conversions that yield clarified, biocatalyst-friendly substrates (9, 10). Even then, the greater complexities required by these processes have generally increased costs (estimates as high as 30¢ per gallon ethanol) as well as eroded scalability and competitiveness (11). Engineering elevated microbial tolerance to the inhibitors released in simpler, less aggressive, hydrolytic methods would, therefore, address one of the major obstacles impeding greater utilization of cellulosic feedstocks (12–15). Here, we show that a targeted combination of genetic and feedstock modifications is sufficient to enhance a single strain to tolerate a wide variety of highly toxified biomass hydrolysates and deliver cellulosic ethanol with performance comparable to current clean sugar ethanol. Our rationally designed approach is, additionally, highly modular: With introduction of a single gene and no further engineering, feedstock-agnostic hydrolysate tolerance is conferred on previously engineered metabolic chassis strains (including one synthesizing a biodegradable plastic) to enable cellulosic products beyond ethanol.

RESULTS

Our prior research demonstrated that in yeast, *Saccharomyces cerevisiae*, responsible for the current global output of biofuel ethanol, increases in media potassium (K+) and pH were sufficient to strengthen membrane potential and enhance production universally across laboratory and industrial strains (16). We, therefore, sought to ascertain the impact of these extracellular adjustments on the fermentation of toxic lignocellulosic feedstocks. Unrefined hydrolysates of all plant biomass, particularly those pretreated under acidic conditions, contain a spectrum of inhibitory by-products but are dominated by the furan aldehydes furfural and 5-hydroxymethylfurfural (HMF) (from dehydration of pentoses and hexoses, respectively) and acetic acid (from deacetylation of hemicellulose) (17–23).

To systematically characterize the impact of these constituent toxicities, we assessed ethanol production in yeast synthetic complete medium (YSC) with the three inhibitors added individually at equimolar concentration. As a completely chemically defined formulation with trace vitamins, minerals, and amino acids, YSC lacks the
Elevated K⁺ and pH also conferred improvements to furfural and HMF but to titers substantially below that of equivalently elevated YSC. On the basis of our prior work that delineated a mechanism of multialcohol tolerance, these improvements, however small, were unlikely to have arisen from tolerance elicited to the 4 ± 0.3 or 34 ± 0.3 g/liter of ethanol produced under inhibition (16). Rather, given reports that S. cerevisiae has various dehydrogenases sufficiently promiscuous to reduce furfural and HMF, we surmised that these extracellular adjustments were conferring tolerance to their furan alcohol products (27–30). We detected 2-furanmethanol (FF-OH) and furan-2,5-dimethanol (HMF-OH) in significant amounts (P ≤ 3.35 × 10⁻⁵), along with the disappearance of furfural and HMF, after only 24 hours of fermentation [Fig. 1B, wild type (“WT”). When comparing the relative toxicity imposed by these alcohols versus their aldehyde equivalents, an ethanol output of 35 ± 0.4 g/liter demonstrated that FF-OH was 9× more tolerated than furfural at equimolar concentration (pink versus purple outline, gray); with extracellular adjustments (pink outline, dark blue), production was boosted an additional 69% to 59 ± 0.5 g/liter (Fig. 1A). The same trend was recapitulated with HMF where HMF-OH accorded 45% higher titer (beige versus brown outline, gray), and media modifications yielded a further 39% that corresponded to production of 69 ± 0.6 g/liter (beige outline, dark blue). Unlike acetic acid where K⁺-only supplementation showed no improvement,
the addition of KCl alone to FF-OH and HMF-OH (pink and beige outline, light blue) elicited improvements intermediate to those with KCl and NH₄OH together. This behavior was consistent with that observed previously for ethanol and underscored the specificity of the membrane permeabilization, and K⁺/pH countermeasures, to alcohol toxicity (16). Furthermore, across concentrations relevant to genuine hydrolysates, these adjustments consistently elicited an enhancement, one whose efficacy was sustained over a larger range with FF-OH and HMF-OH than with furfural and HMF (fig. S1, B and C). Given the gains attainable individually on the three dominant inhibitors, we surmised that augmenting in vivo conversion of furfural and HMF to alcohols, paired with the elevation of extracellular K⁺ and pH, could encapsulate a unified method for bestowing tolerance against the totality of toxicities present in genuine hydrolysates.

On the basis of literature describing reductases with detoxifying activity toward furfural and HMF, we constructed yeast strains overexpressing ADH6, ADH7, or GRE2 from *S. cerevisiae* or ADH4 from *Scheffersomyces stipitis* (29–33). Fermentation benchmarking over two repressive conditions combining the trio of inhibitors revealed that *S. cerevisiae* GRE2 evoked the greatest improvement among the candidates. When compared to the WT, these improvements amounted to as much as 32% (fig. S2). Furthermore, that the GRE2 strain sustained a smaller production drop than the WT under the harsher of the two conditions suggested that increased detoxification can enhance robustness over a wider range of toxicity. We quantified FF-OH and HMF-OH to corroborate the augmented reduction capacity and found that GRE2 overexpression produced 25 ± 1% higher concentrations of FF-OH and 56 ± 6% of HMF-OH ($P \leq 3.2 \times 10^{-2}$; fig. 1B "GRE2"). Despite nonstoichiometric conversion of furfural and HMF, that greater formation of FF-OH and HMF-OH could be engineered indicated that we could indeed productively mitigate the aggregate toxicity by converting the aldehydes into a form (alcohols) for which we have the means to counteract effectively.

To characterize productivity under combined furfural and HMF stress, as well as the impact from modulating reductase activity, we quantified the relationship between tolerance and ethanol production for the WT, the GRE2 overexpression strain, and one deleted for GRE2. Our prior work had demonstrated that the viable fraction in an actively fermenting population declined rapidly with accumulating ethanol; however, this mortality could be rescued by K⁺ and pH adjustments in a dose-dependent manner. Furthermore, time integrations of these viable population measurements (“integrated viable cell density”) from progressively higher adjustments, followed by correlation with ethanol titers, established the relationship between tolerance and production. Specifically, a readout of the time-averaged specific productivity exclusive to the differentially decaying viable fractions was revealed in the correlation slope (16). We, therefore, measured titers and viabilities from a series of fermentations containing furfural, HMF, and incrementally higher adjustments of K⁺ and/or pH. The slope from the GRE2 strain revealed a 59% improvement in per-cell performance over the WT, demonstrating that increased detoxification subdued the combined inhibition effectively to sustain metabolic activity (Fig. 1C). For higher extracellular adjustments (e.g., yellow to red data points), we quantified the GRE2 strain to actually have lower viabilities yet greater ethanol production, illustrating that WT cells, while alive, were stalled metabolically (fig. S3). Furthermore, the higher prevalence of alcohols from GRE2 overexpression (from HMF-OH and FF-OH formation as well as improved ethanol production) likely predisposed the strain to the increasing alcohol protective benefits of incremental K⁺ and pH, the outcome of which was reflected as higher per-cell performance. Deletion of GRE2 corroborated these trends: While the gre2Δ/gre2Δ strain retained the same specific productivity as the WT, the downshift in correlation indicated that any given viability or particular extracellular adjustment would result in lower ethanol output. Thus, GRE2 contributes directly to the resilience of the population by converting furfural and HMF aldehyde stress into alcohols that are subsequently ameliorated by K⁺ and pH treatment.

Given the efficacy exhibited by the GRE2 overexpression strain, we considered various adaptation approaches to further improving hydrolysate tolerance. While whole-strain laboratory adaptive evolution is well practiced for augmenting fitness, selective advantages from genome-wide drift have been shown to incur costs in robustness (34, 35). To minimize the risk of pleiotropic deficits undermining feedstock range and strain performance, we opted to hone the detoxification capabilities of GRE2 specifically via directed evolution. Therefore, we cultured a yeast library consisting of polymerase chain reaction (PCR)–mutagenized GRE2 variants under combined furfural, HMF, and acetic acid stress and challenged it to increasing loads over time (fig. S4) (36). Validated, postselection sequence isolates were subcloned and introduced anew into S288C, and individual strains screened for a fermentation advantage.

The allele exhibiting the greatest gain was a triple mutant containing a proline to serine substitution at amino acid 48, isoleucine to valine at amino acid 290, and a silent aspartate mutation at amino acid 133 (GRE2P48S + I290V + D133S; hereafter as GRE2evol). Across several toxicity combinations mimicking a range of pretreatment severities, GRE2evol consistently conferred improvements over unevolved GRE2 in ethanol production (fig. S5A). Other than K⁺ and pH requirements, the superior phenotype was not dependent on extracellular factors supplied by a favorable nutritional environment: In yeast nitrogen base (YNB) minimal medium, containing no amino acids and solely glucose, ammonium sulfate, salts, and trace vitamins, GRE2evol was capable of eliciting a percentage gain comparable to those observed under nutrient-replete conditions (fig. S5B). Furthermore, under nontoxic conditions where the superphysiological abundance and reductive capacity of GRE2evol could potentially cross-react with acetaldehyde to boost ethanol yield, we observed statistically unchanged levels of performance (fig. S5C). Along with the absence of a major negative impact, these data suggested that the highly transcribed GRE2evol imposed a low-expression burden and functioned largely in an orthogonal manner specific to the hydrolysate inhibitors.

As rates of detoxification are directly proportional to cell biomass (Fig. 1B demonstrated that even the WT could completely reduce furfural and HMF in under 24 hours at production cell densities), we lowered inocula more than 20× in a growth assay designed to emphasize fitness advantages enabled by GRE2evol under full toxicity. Here, the WT failed completely to expand, while GRE2evol shortened the lag phase by approximately 7 hours compared to overexpressed unevolved GRE2 (Fig. 1D). The exit from lag was preceded by the detoxification of furfural and HMF, where GRE2evol exhibited the highest rates of depletion. Incidentally, when juxtaposed with growth, the decreases in inhibitor concentration revealed further that, unlike furfural, HMF need not be fully detoxified in order for growth to commence and approximately 1.5 g/liter are tolerable by yeast.
Expression of $\textit{GRE2}^{\textit{evol}}$ was, moreover, capable of conferring near-parity ethanol production between inhibitor-free and fully toxified conditions. To first establish a reference upper bound for inhibitor-free productivity and titer, we fermented the WT in traditional high glucose synthetic medium supplemented with potassium bicarbonate ($\text{KHCO}_3$) and calcium carbonate ($\text{CaCO}_3$), selected for their widespread industrial and agricultural availability, to provide elevated $K^+$ and pH buffering. Under these optimal conditions, ethanol reached $109 \pm 1$ g/liter in under 32 hours (Fig. 2A, blue).

When we proceeded to toxify with 62 mM furfural, 48 mM HMF, and 100 mM acetic acid, a benchmark of above-average toxicity that we formulated to balance broad feedstock applicability with acceptable yeast performance, output was repressed by 69% to $34 \pm 1.2$ g/liter of ethanol. This was despite the adjustment to pH 5 per standard bioethanol practices that also neutralized the acetic acid component (Fig. 2A, red) (17, 18, 21, 37). Subsequent supplementation with $\text{KHCO}_3$ and $\text{CaCO}_3$ was sufficient to rescue stalled productivity in the WT and achieve production of $81 \pm 2$ g/liter (Fig. 2A, orange). However, substitution with the $\textit{GRE2}^{\textit{evol}}$ strain provided a further gain, boosting rate by an additional mean 39% and final product by 31% (Fig. 2A, gray solid). The ethanol titer of $106 \pm 1$ g/liter amounted to 97% of that from inhibitor-free medium, demonstrating near-unrestricted ethanol production under full toxicity conditions. Moreover, that the $\textit{GRE2}^{\textit{evol}}$ strain remained predominantly repressed under toxification at pH 5 (Fig. 2A, gray dotted) illustrated the necessity of both the genetic- and extracellular-derived enhancements to attain maximal tolerance.

The targeted detoxification provided by the combination of $\textit{GRE2}^{\textit{evol}}$ expression and feedstock adjustments was applicable beyond artificial formulations to a range of genuine lignocellulosic feedstocks. Influenced partly by their history of toxicity and limited utility, the hydrolysates currently available are produced largely at research scale and focused on maximally detoxified cellulose sugars (6–8, 10, 38, 39). We procured a collection of seven samples representing a diversity of plant sources (corn stover, sugar cane bagasse, wheat straw, giant miscanthus, and switchgrass) and various pretreatment methods and confirmed that inhibitors were present in the ranges of 0.1 to 21 mM furfural, 0.1 to 6 mM HMF, and 0 to 178 mM acetic acid (table S1). These quantities were sufficiently mild such that, aside from the standard supplementation with urea (to provide nitrogen) and adjustment to pH 5, the WT was capable of fermenting all glucose to completion under otherwise unmodified conditions (Fig. 2B, blue, and fig. S6).

Because inhibitor and glucose loads inherent in most of the samples fell below that of our formulated benchmark, we toxified all samples to impinge on our engineered tolerance limits (table S2). Furfural was raised to an average of 45 mM, HMF to 35 mM, and acetic acid to 100 mM to more closely align with published inhibitory combinations, and glucose was raised to an average of 220 g/liter for increased osmotic, and ensuing ethanol, stress (17–23). Supplementation with pure chemical forms of these components was necessary as prehydrolyzed solids (and hydrolysis methods) were unavailable to us to boost toxicity using raw cellulosic material. Under these conditions, the WT became repressed even with adjustments to pH 5 (Fig. 2B, red).

In deacetylated and mechanically refined (DMR) corn stover, for example, this repression amounted to an output 39% of its minimally modified control. (Glucose supplementation in the remaining feedstocks concomitantly raised ethanol ceilings, rendering comparisons with their minimally modified, lower-glucose controls invalid). The subsequent addition of $K^+$ and pH buffering elicited increases of 1.8 to 10.7×, enabling 57 to 91% of the previously residual glucose to be consumed (Fig. 2B, orange, and fig. S6). However, when these additions were combined with the $\textit{GRE2}^{\textit{evol}}$ strain, over 91% of the final remaining substrate was consumed, increasing production by 10 to
61% over the WT to titers of 81 ± 1 to 113 ± 0.4 g/liter (Fig. 2B, gray, and fig. S6). This performance corresponded to theoretical conversion yields of 78 to 91% and were all obtained from a single strain derived from a historically underperforming laboratory lineage; yet, they exceeded currently published values of cellulosic ethanol produced from undetoxified industrial feedstocks (40–42). Thus, despite the compositional and pretreatment–by-product complexity across this diversity of material (e.g., miscanthus contains ferulic and p-coumaric acids derived from lignin degradation; table S3), the combination of GRE2<sup>evol</sup> and alcohol protective adjustments exhibited robustness and sufficiency to efficiently ferment highly toxified genuine feedstocks.

Because genuine hydrolysates are highly undefined mixtures with unknown levels of salts, uninformed additions of K<sup>+</sup> and particular buffer counterions could potentially exceed osmotic shock thresholds in yeast (43–45). As we had previously established fermentation-beneficial effects of K<sup>+</sup> to at least 110 mM and Ca<sup>2+</sup> to be fermentation neutral, we deemed it important to maintain total K<sup>+</sup> within this limit and modulate pH via Ca<sup>2+</sup>-based buffers or bases (16). Thus, cation concentrations in each sample were determined by mass spectrometry and used to determine the specific mixes of KHCO<sub>3</sub>, CaCO<sub>3</sub>, and calcium hydroxide to provide as supplementation (tables S1 and S2). The atypically high K<sup>+</sup> concentration in wheat straw provided an opportunity to validate these constraints: When 50 mM KHCO<sub>3</sub> and 100 mM CaCO<sub>3</sub> were used in lieu of 150 mM sodium bicarbonate, the oversupply of K<sup>+</sup> resulted in decreases to performance despite the same buffering capacity (fig. S7). Therefore, supplementation customized to each feedstock conforming to salt-specific limits is necessary to achieve maximal efficacy.

These tolerance capabilities were fully transferrable to the fermentation of inhibitor-laden xylene, the pentose comprising a substantial portion of lignocellulosic sugars that unmodified <i>S. cerevisiae</i> cannot consume (6, 17). Our laboratory had previously engineered a strain (XYL<sup>+</sup>) that efficiently fermented xylose to ethanol but preferentially used glucose if present (46). Therefore, we formulated a YSC-based hydrolysate to favor xylene metabolism, yet mimic genuine cellulose proportions, by combining xylene with starch whose glucan polymers were slowly digested to glucose via incrementally dosed amylases. In the absence of inhibitors, the XYL<sup>+</sup> chassis completely consumed available xylene and glucose in this medium and produced 62 ± 0.9 g/liter of ethanol (Fig. 3A, bold blue, and fig. S8). Toxification with the benchmark suite of furfural, HMF, and acetic acid repressed production to 8 ± 1 g/liter, and subsequent treatment with elevated K<sup>+</sup> and pH buffering provided recovery to 22 ± 0.9 g/liter with improved proportional usage of sugar (Fig. 3A, bold red and orange, and fig. S8). However, when we introduced GRE2<sup>evol</sup> into XYL<sup>+</sup> with no further modifications, the chassis was functionalized to ferment all xylose and glucose, restoring production essentially 100% to 66 ± 0.6 g/liter (Fig. 3A, bold gray, and fig. S8). Our combination of tolerance enhancements thus liberated the prior-engineered metabolism to achieve full production capacity under complete toxicity conditions.

Moreover, the GRE2<sup>evol</sup>-enhanced XYL<sup>+</sup> strain maintained its tolerance capabilities on multiple toxified genuine hydrolysates. Because of catabolite repression, xylene went unconsumed by the XYL<sup>+</sup> chassis strain in minimally modified hydrolysates of sugarcane bagasse and high-acid corn stover (Fig. 3A, nonbold blue, and fig. S8). Miscanthus proved, fortuitously, to be an exception: As glucose was present in amounts lower than xylose and depleted rapidly, xylene metabolism remained sufficiently active such that both monomeric sugars were near-entirely consumed. Toxification from increased glucose, furfural, HMF, and acetic acid repressed production in all three hydrolysates to below 15 g/liter of ethanol. While these inhibitory conditions were relieved by elevated K<sup>+</sup> and pH buffering, the recoveries exhibited much greater variability than in S288C (Fig. 3A, nonbold red and orange, and table S2). In high-acid corn stover, for example, an improvement was statistically questionable (<i>P</i> = 0.76), while that for miscanthus was a significant 689 ± 1% (<i>P</i> = 1.00 × 10<sup>−10</sup>). However, these inconsistent recoveries were subsequently remedied in a robust manner by the introduction of GRE2<sup>evol</sup> where gains averaged a further 260% (Fig. 3A, nonbold gray). In miscanthus, this additionally enabled partial fermentation of xylene despite higher catabolite repression from supplemented

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**Fig. 3. Elevated K<sup>+</sup>, pH buffering, and GRE2<sup>evol</sup> overexpression encapsulate a lightweight cellulosic tolerance platform integrable with heterologous engineered pathways.** Bars follow the conditions and color scheme described in Fig. 2B. (A) Cellulosic ethanol titers from a xylose-consuming strain (XYL<sup>+</sup>) fermenting synthetic medium (bold) containing xylene and starch (slowly hydrolyzed to glucose via amylases), or the indicated biomass hydrolysate (nonbold). (B) Cellulosic lactic acid titers from an ethanol-handicapped strain expressing lactate dehydrogenase from <i>L. mesenteroides</i> (D-LA<sup>+</sup>) fermenting synthetic medium (bold) or the indicated biomass hydrolysate (nonbold). Data are means ± SD from three biological replicates.
glucose (fig. S8). Again, our combination genetic and extracellular tolerance enhancements proved effective across disparate toxified genuine feedstocks.

The recapitulation of phenotype and substrate robustness on a preexisting metabolic chassis suggested that GRE2<sup>evol</sup> expression and K<sup>+</sup>/pH feedstock adjustment could constitute a functionally orthogonal tolerance platform integrable with other engineered end products. Given that our alcohol-focused countermeasures may have conferred a bias toward ethanol production, we opted to endow hydrolysate tolerance on a glucose-consuming strain synthesizing the biodegradable plastics precursor lactic acid (d-LA), a chemical commodity projected to reach U.S. $9 billion by 2025 (47). Given the evolutionary predilection of S. cerevisiae for alcoholic fermentation, elimination of ethanol has been a major goal in all efforts to reengineer yeast for nonethanol products. Here, we opted to minimize, rather than eliminate, glycolytic flux toward ethanol to maintain subsistence adenosine 5’-triphosphate generation for cell growth and active lactate export (48). As lactate and ethanol share a common precursor in pyruvate, we curtailed pyruvate decarboxylase activity and losses to ethanol by creating a 

\[ \text{pdc1Δ/pdc1Δ pdc5Δ/pdc5Δ:pTEF1m4-PDC5} \]

strain where a sole chromosomal pyruvate decarboxylase gene was transcribed using a handicapped variant of the TEF1 promoter (49–51). Reductive conversion of pyruvate to lactate was fulfilled via expression of d-lactate dehydrogenase from Leuconostoc mesenteroides (52).

In synthetic medium supplemented with KHCO<sub>3</sub> and sufficient CaCO<sub>3</sub> to buffer lactic acid accumulation, and likewise in DMR corn stover minimally supplemented with CaCO<sub>3</sub>, our D-LA<sup>+</sup> chasis generated inhibitor-free reference titers of 66 ± 0.3 and 65 ± 0.4 g/liter, respectively (Fig. 3B, blue). Under complete toxification, elevated K<sup>+</sup> and pH buffering, combined with GRE2<sup>evol</sup> expression and no additional modifications, succeeded in conferring tolerance such that cellulosic lactic acid reached 90 and 96% of these clean sugar benchmarks (Fig. 3B, gray). Similar to XYL<sup>+</sup>, the engineered D-LA<sup>+</sup> metabolism was largely liberated from inhibition to elicit near-unrestricted production. Furthermore, we tested for strain robustness to additional hydrolysates using low-acid corn stover and wheat straw. As before, these feedstocks (unlike DMR corn stover) contained the stress from supplemented glucose and toxification from the trio of inhibitors. Nevertheless, GRE2<sup>evol</sup> expression was capable of eliciting mean gains of 19% over the unengineered chassis to attain cellulosic product titers greater than 50 g/liter.

Although higher lactic acid production from yeast has been reported, all prior studies were conducted using traditional, clean sugar feedstocks; titers here, furthermore, were restrained by our chassis’ inherently limited synthesis capabilities (52–54). The single-transformation tolerance phenotype minimally interfered with the engineered lactic acid metabolism (fig. S5C); likewise, GRE2<sup>evol</sup> remained predominantly orthogonal and unperturbed to alleviate toxicity efficiently. To our knowledge, this represented the first demonstration of a nonethanol cellulosic product delivered with industrially relevant performance from multiple highly toxified genuine feedstocks using a single strain.

**DISCUSSION**

Our results describe a functionally independent, lightweight platform that both endows yeast with general lignocellulosic hydrolysate tolerance and integrates harmoniously with preexisting metabolically engineered chassis strains. Through systematic characterization of the three dominant toxicities released from biomass pretreatment, we have demonstrated that tolerance to each inhibitor can be realized through standard neutralization (for acetic acid) or conversion of the aldehydes to alcohols, which are subsequently ameliorated by the prior-identified elevated K<sup>+</sup> and pH treatment (for furfural and HMF). The general practice of hydrolysate tolerance can, therefore, be reduced to two specific and readily modifiable parameters: In a genetic background enhanced by GRE2<sup>evol</sup> for the accelerated reduction of furfural and HMF, a large diversity of feedstocks, regardless of plant source and/or pretreatment process, can be accommodated via tailored adjustment of K<sup>+</sup> and pH. This substrate robustness indicates that the spectrum of hydrolytic by-products other than furfural, HMF, and acetic acid (for example, the various acidic and phenolic inhibitors shown in table S3) may be qualitatively immaterial.

These benefits, collectively, renew and boost the value proposition of cellulosic fermentation. Wide feedstock compatibility can reduce the dependence on specific crop types or pretreatments and, consequently, ameliorate the supply variability (e.g., from seasonality, storage stability) and cost uncertainties surrounding biomass (13). Similarly, heightened tolerance, in addition to harnessing toxic sugar streams or transport-friendlier concentrates, enables production conditions of minimized contaminant growth that would otherwise require the standard-practice, but public health-concerning, use of antibiotics (55). Last, the targeted specificity of our detoxification approach underlies the high decoupling with metabolism and straightforward integration with previously engineered pathways such as those for xylose consumption and lactic acid synthesis. This underscores the notion of a “drop-in” tolerance phenotype extensible to even more non-native capabilities and high-volume biofuels and biochemicals.

**MATERIALS AND METHODS**

**Plasmid construction**

All plasmids were assembled using the Gibson method from segments generated via PCR. Amplification of plasmid backbone, yeast promoter, protein coding, and transcription termination fragments (see below) was carried out using the Phusion High-Fidelity DNA Polymerase (New England Biolabs, #M0530; www.neb.com) in 50 μl of reaction following the manufacturer’s directions. Primers were designed with 25- to 30–base pair (bp) 5’ overhangs to serve as assembly junctions and annealing temperatures of primer pairs optimized using the vendor-provided calculator (tmcalculator.neb.com). The suggested extension times of 30 s/kb were often inadequate empirically and extended to 60 s/kb for problematic amplicons (fragments of >2 to 3 kb tended to require the higher extension rate). For templates containing a bacterial origin of replication (e.g., plasmid backbone segment), PCR products were further digested with 20 U of Dpn I (New England Biolabs, #R0176) added directly to the reaction sample after thermocycling (i.e., no additional restriction enzyme buffer), incubated for 90 min at 37°C, and heat-inactivated for 20 min at 80°C. All fragments were purified and concentrated (up to three pooled PCR reactions per column) using the QIAquick PCR Purification Kit (QIAGEN, #28106; www.qiagen.com), and DNA concentrations were quantified with a NanoDrop Microvolume ultraviolet-visible spectrophotometer. Gibson reactions were prepared from these eluates using 50 to 100 ng of vector and molar ratios of one part vector to four to eight parts of each insert.
in the lowest volume possible (i.e., no additional H2O to meet the instructed minimum of 10 μl). Assembly enzymes were supplied through a cocktail (New England Biolabs, #E2611), and reactions were incubated for 30 min at 50°C, followed by an additional 1 hour to overnight at room temperature. Chemically competent Escherichia coli (New England Biolabs, #C2992) were transformed and cultured per the manufacturer’s instructions, and ampicillin-resistant isolates were screened by PCR using vector- and insert-specific primers. Plasmids derived from positively scoring transformants were extracted using the QIAprep Spin Miniprep Kit (QIAGEN; #27106) and validated by Sanger sequencing (Quintarabio; www.quintarabio.com/).

Minimal backbone segments, containing solely the bacterial replication origin, ampicillin marker, yeast replication origin, and yeast selection marker, were sourced from the p415 and p426 expression series developed by Mumberg et al. (56). For strong transcription in yeast coupled to glycolytic activity, we either retained the TDH3 (“GPD”) or TEF1 promoters from the Mumberg vectors or cloned the ~703 to ~1 fragment of the PDC1 promoter from FY4/5 genomic DNA. Protein coding sequences for ADH6, ADH7, and GRE2 were likewise amplified directly from FY4/5 genomic DNA. For ADH4 from S. stipitis, we contracted with Bio Basic (www.biobasic.com) to synthesize a S. cerevisiae codon-optimized sequence from the publicly available protein translation (GenBank accession no. XM_001387085). Similarly, ldha from L. mesenteroides subsp. mesenteroides was produced by Eurofins Genomics (www.eurofinsgenomics.com) from the amino acid sequence available from UniProt (gene entry LEUM_1756). For transcription termination, we either retained the CYC1 element from Mumberg or, from FY4/5 genomic DNA, cloned the 166 bp immediately following the stop codon of ADH1 or the 296 bp following ACT1.

To clone the GRE2 mutants emerging from toxicity selection, the plasmid-based coding sequences were distinguished from chromosomal GRE2 via an initial PCR using primers binding to library plasmid elements. Specifically, the collection of mutated genes, including their nonmutagenized promoters and terminators, was originally subcloned via Gateway recombination and positioned between attB1 and attB2 sequences (36). Thus, from total DNA isolated from the final inhibitor-tolerant culture (fig. S4), we performed an initial amplification using primers annealing to these unique attB elements. To further subclone the coding sequences of hypertolerant GRE2 mutants, the plasmid-derived amplicons were used to template a second-pass PCR excluding the GRE2 promoter and terminator. These amplified protein-coding fragments were subsequently Gibson-assembled into our final expression constructs. See table S4 for a complete list of overexpression plasmids used in this study.

Yeast strain construction

Recombinant strains were created following the lithium acetate chemical transformation method of Gietz et al. (57). For single plasmid introduction, 50 ng of p(RS)415-based DNA was used with one OD600 unit of cells and selection carried out on solid YSC-Leu dropout medium. For strains originating from two simultaneous plasmids, 150 to 300 ng each of p426TEF- and p(RS)415-based DNA was used with three to four OD600 units, and selection was performed on YSC-Ura-Leu solid medium. For chromosomal integrations, 800 ng to 1 μg of linear DNA was used with 7 to 10 OD600 units. Low transformation efficiencies (e.g., from variability in strain, locus, and DNA secondary structure) were typically resolved by increasing the DNA to cell ratio, amount of salmon sperm carrier DNA, or heat shock incubation time (up to 40 to 50 min at 42°C). In addition to laboratory standard BY4743, the gre2Δ::kanMX4/gre2Δ::kanMX4 diploid used for LAMY629 preexisted this study and was obtained from the Saccharomyces Genome Deletion Project collection (www-sequence.stanford.edu/group/yeast_deletion_project/).

To create the diploid xylose-consuming chassis (XYL\(^+\)), a xylose-enabled MATa leu2-3 haploid (internal strain F258) available from the development efforts of Zhou et al. (46) was transformed with plasmid pH727 (GAL::HO LEU2; gift from J. Haber of Brandeis University) to generate a MATa equivalent (46, 58). Induction of HO in Leu\(^+\) transformants was conducted in liquid medium containing 20 g/liter of galactose for 6 hours at 30°C (complete mating type switching protocol, including preinduction, is available from the Haber Lab website: www.bio.brandeis.edu/haberlab/jehsite/protocol.html). Individual colonies, recovered from growth in glucose medium additionally containing leucine to discard pH727, were screened for MATa haploids by α- and α-factor sensitive mating type tester strains (59). A validated MATa leu2-3 haploid was subsequently mated with F258 to create the homozygous XYL\(^+\) leu2-3 chassis strain LAMY435 that preceded LAMY419 and LAMY665.

To create the S288C leu2-3 predecessor of LAMY660, LAMY661, and LAMY663, the defective his3Δ1 and ura3Δ0 alleles in BY4743 were complemented sequentially by targeted chromosomal replacements. Briefly, a PCR product encompassing the full-length coding sequence of HIS3 was amplified from FY4/5 genomic DNA, introduced into BY4743, and transformants selected for histidine prototrophy. To repair ura3Δ0, which spans a segment larger than the open reading frame of URA3, a PCR product including 320 bp of the URA3 promoter and 194 bp beyond the stop codon was amplified from FY4/5 genomic DNA, introduced into the His\(^+\) intermediate, and transformants selected on minimal YNB medium supplemented solely with leucine to yield chassis strain LAMY651.

To create the diploid ethanol-handicapped chassis for lactic acid production, we first generated a fully ethanol-deficient pdc\(^-\) haploid by creating a markerless deletion of PDC5 in a MATa pdc1Δ::kanMX4 strain sourced from the Saccharomyces Genome Deletion Project collection (PDC6, while intact, is functionally inert). Briefly, plasmid pCRSPR, PDC1+5 expressing a Candida albicans/S. cerevisiae codon-optimized version of Cas9, as well as a guide RNA targeting the protospacer adjacent motif (PAM)–proximal sequence TGCTAAAGAACCGTATTCT common to PDC1 and PDC5, was coelectroporated with the double-stranded linear repair template CATATCAATCTCAGAGAAACAACATAATACAAAA- CAAAGAAGAAACAGCTAATACACGTAA into the MATa pdc1Δ::kanMX4 haploid according to the protocol of Vyas et al. (60) (plasmid and template are gifts of B. Urankul). Transformants were selected on solid YP (yeast bacto-peptone) medium containing 3% ethanol, 3% glycerol, and nourseothricin (YPEG + NAT) (100 μg/ml), and isolates comparatively grown on YPEG + NAT and YPD (YP with 2% glucose) solid media to identify the pdc\(^-\) phenotype (development strain LAMY399).

Separately, a haploid containing PDC5 transcribed by the low-strength TEF1m4 promoter mutant was generated in a MATa pdc1Δ::kanMX4 strain sourced from the Saccharomyces Genome Deletion Project collection. Here, the in-locus markerless edit was likewise accomplished through a similar CRISPR protocol with plasmid LAMB66 (featuring uracil selection in yeast and improved Cas9 expression) encoding a guide RNA targeting sequence TTCTCGATCAATATACTGTA in the PDC5 promoter and the double-stranded
replica template CAAAGGTCGCCCTTTTCTTAGAAAAACTCATGCTACTGTTAACTTTTACCTTCTGCTAGGCAAGACGACGTGTCGCTACGGCTCCCTTCTGCAGATTCCTGCTAGTCTTCACTTTTTTTTACTTCTCTCGCTGTCAGTAGAGAAGACGACGTGACACTCACGTTTTTTCTCTTTCGATGACCTCCCATTGATATTTACACGTCTTTTTTTCTTTTTGAGTTcTTTcTtGGTTTCTTTcTCTTCGTCGAgAgAGGCAATAAAAATTTTTATGAAATAACCTTAGGATATTTGAGATTGAGC -
CCCTCTTTCTTCCTCTAGGGTGTCGcTAATTACCCG-
CAGATTTTCTCGGACTCCGCGCATCGCCGTACCACTTCAA-
ATACGTAAACCTGCATTAAGGGAACAAAAGCTGGAGCT-
ATAGTTTTTTCTCTTCTCGGACTCCGCGCATCGCCGTACCACTTCAA-
ATACGTAAACCTGCATTAAGGGAACAAAAGCTGGAGCT-

Media and fermentation conditions

To provide a consistent but modifiable medium to accommodate our entire collection of strains, all baseline culturing was performed in YSC composed of 1.5 g/liter of YNB without amino acids and ammonium sulfate (BD Difco, #233520; www.bd.com), 5 g/liter of ammonium sulfate, 0.2 mM inositol, 0.1 g/liter of each of the 20 amino acids, and 0.1 g/liter each of adenine and uracil (all from Sigma-Aldrich; www.sigmaaldrich.com). Strains containing a p(RS)415 plasmid were maintained in medium lacking leucine and those with a p426 plasmid lacking uracil. Unless indicated otherwise, individual strains were expanded and acclimated to high cell density and high sugar conditions in singlicate YSC-based cultures and divided into triplicate biological samples upon inoculation into fermentation (Fig. 1A to C, 2, and 3) or growth (Fig. 1D) media. All yeast culturing and fermentations were conducted at 30°C in Erlenmeyer flasks (≥25 ml) shaken at 200 rpm or glass tubes (≤12 ml) rotated at 4°C until analysis. Similarly, preinoculation fermentation media were syringe-filtered and diluted 5−1 in water for HPLC verification of starting glucose and inhibitor concentrations.

To screen the panel of overexpressed reductases on fermentation (Fig. S2), strains LAMy312, LAMy553, LAMy579, LAMy580, and LAMy589 were started in YSC-Leu containing 180 g/liter of glucose and diluted for further acclimation overnight to higher glucose in YSC-Leu containing 240 g/liter of glucose. Upon reaching OD600 = 2.5 to 3, 100 OD600 units of cells were harvested, and cell pellets were resuspended in 4 ml of YSC-Leu containing 260 g/liter of glucose, 62/48/100 mM furfural/HMF/acetic acid, and 60/140 mM KHCO3/CaCO3. A second set of cell pellets was resuspended in 4 ml of YSC-Leu containing 240 g/liter of glucose, 84/64/133 mM furfural/HMF/acetic acid, and 60/200 mM KHCO3/CaCO3. Cell-free samples of the fermentation medium were harvested after 46 hours per procedure described above for HPLC analysis.

To prepare samples for mass spectrometric quantification of furfural, HMF, FF-OH, and HMF-OH (Fig. 1B), strains LAMy312 and LAMy579 were started in YSC-Leu containing 180 g/liter of glucose and diluted for further acclimation overnight to higher glucose in YSC-Leu containing 240 g/liter of glucose. Upon reaching OD600 = 2.5 to 3, 100 OD600 units of cells were harvested, and cell pellets were resuspended in 4 ml of YSC-Leu containing 260 g/liter of glucose, 62/48/100 mM furfural/HMF/acetic acid, and 60/140 mM KHCO3/CaCO3. After 24 hours of fermentation, 10 μl of supernatant from cell-pelleted

samples was combined with 100 μl of 100% ultra-HPLC (UHPLC) grade methanol containing isotopically labeled amino acids (provided by the Whitehead Institute Metabolite Profiling Core Facility), and mixtures were stored at −80°C until analysis. Similarly, 100 μl of syringe-filtered preinoculation fermentation medium was extracted to determine starting concentrations of inhibitors.

To assess the impact of GRE2 and increasing K+ and pH conditions on cell viability under combined furan aldehyde stress (Fig. 1C and fig. S3), strains LAMy312, LAMy579, and LAMy629 were started in YSC-Leu containing 180 g/liter of glucose, diluted, and further acclimated overnight in YSC-Leu containing 260 g/liter of glucose. Upon reaching OD600 = 2.5 to 3, 100 OD600 units of cells were harvested, and cell pellets were resuspended in 4 ml of YSC-Leu containing 260 g/liter of glucose, 62/48 mM furfural/HMF, and supplemented with (from blue to red) 25 mM KCl, 5 mM NH4OH, 10 mM NH4OH, 15 mM NH4OH, 20 mM NH4OH, 24 mM NH4OH, 24/25 mM NH4OH/KCl, 24 + 5 mM NH4OH + NH4OH, 24 + 10 mM NH4OH + NH4OH, 24 + 15 mM NH4OH + NH4OH, or 24 + 20 mM NH4OH + NH4OH. As above, "/" indicates addition of components during preparation, while the "+" here indicates addition after 22 hours of fermentation. At 0, 24, 32, 52, and 71 hours after inoculation, cell densities were measured, and 20 μl was taken for immediate methylene blue viability staining and microscopy (fig. S3). Preinoculation media and cell-free fermentation samples at 71 hours were harvested for HPLC analysis.

To screen for a fermentation advantage conferred by overexpression of GRE2rd versus GRE2 (fig. SSA), prototrophic strains LAMy660, LAMy661, and LAMy663 were started in minimal YNB medium (i.e., no amino acids) containing 100 g/liter of glucose and diluted for further acclimation overnight to higher glucose in 1.3× YNB containing 240 g/liter of glucose. Upon reaching OD600 = 2.5 to 3, 100 OD600 units of cells were harvested, and cell pellets were resuspended in 4 ml of YSC-Leu containing 260 g/liter of glucose, 62/48/100 mM furfural/HMF/acetic acid, and 60/140 mM KHCO3/CaCO3. A second set of cell pellets was resuspended in 4 ml of YSC-Leu containing 280 g/liter of glucose, 62/48/100 mM furfural/HMF/acetic acid, and 60/140 mM KHCO3/CaCO3. A third set of cell pellets was resuspended in 4 ml of YSC-Leu containing 260 g/liter of glucose, 62/48/100 mM furfural/HMF/acetic acid, and 60/200 mM KCl/CaCO3. Here, because of the reduced acidity from the lack of amino acids in YNB, equimolar KCl was used in lieu of KHCO3, and CaCO3 consequently increased to 200 mM, to achieve a pH within range of that in equivalent YSC-Leu. For fig. S5C, LAMy660, LAMy663, and LAMy692 were started in YSC–Ura-Leu–His–Trp–Ade–Lys (“YSC–6 AA”) dropout medium containing 100 g/liter of glucose, diluted, and further expanded overnight in YSC–6 AA containing 250 g/liter of glucose. Upon reaching OD600 = 2.5 to 3, 100 OD600 units of LAMy660, 663 were harvested, and cell pellets were resuspended in 4 ml of YSC-Leu with 260 g/liter of glucose and 60/140 mM KHCO3/CaCO3. For LAMy690 and LAMy692, 90 OD600 units were harvested and resuspended in 4 ml of YSC-Ura-Leu with 165 g/liter of glucose and 60/250 mM KHCO3/CaCO3. Cell-free samples of the fermentation medium were harvested after 40 hours (fig. SSA), 48 hours (fig. S5B), or 24 hours (fig. S5C) for HPLC analysis.

For assaying a fitness advantage conferred by GRE2rd versus GRE2 overexpression (Fig. 1D), strains LAMy660, LAMy661, and LAMy663 were started in YSC–6 AA medium containing 100 g/liter of glucose, diluted, and further expanded overnight in YSC–6 AA containing 250 g/liter of glucose. Upon reaching OD600 = 2.5 to 3, nine OD600 units of cells were harvested, and cell pellets were resuspended in 13.5 ml of YSC–6 AA containing 50 g/liter of glucose, 40/28/55 mM furfural/HMF/acetic acid, supplemented with 50 mM KCl, and adjusted to pH 6 with NH4OH. Cell densities were measured at 0, 24, 48.5, 51.5, 56, 65.5, 69, 72, 75.5, 79.5, 91, and 96.5 hours after inoculation. Cell-free samples of the medium were collected at 0, 24, 48.5, 56, 65.5, 72, 79.5, and 96.5 hours for determination of furfural and HMF concentrations by HPLC.

For the fermentation time courses of Fig. 2A, strains LAMy660 and LAMy663 were started in YSC–6 AA containing 100 g/liter of glucose, diluted, and further expanded overnight in YSC–6 AA containing 250 g/liter of glucose. Upon reaching OD600 = 2.5 to 3, 100 OD600 units of LAMy660 were harvested, and cell pellets were resuspended in 4 ml of YSC-Leu with 260 g/liter of glucose and 60 or 140 mM KHCO3/CaCO3 (uninhibited control, blue), 62/48/100 mM furfural/HMF/acetic acid and adjusted to pH 5 with NH4OH per bioethanol practices (inhibited control, red), or 62/48/100 mM furfural/HMF/acetic acid and 60/140 mM KHCO3/CaCO3 (orange). This final condition, as well as that of the inhibited control (inhibitor trio adjusted to pH 5 with NH4OH), was repeated with 100 OD600 units of strain LAMy663 for direct comparison with LAMy660 (gray solid and gray dotted, respectively). At 0, 16, 32, and 48 hours after inoculation, cell-free samples of the fermentation medium were harvested for HPLC analysis.

To assess performance in genuine biomass feedstocks (Fig. 2B), strains LAMy660 and LAMy663 were started in YSC-Leu containing 100 g/liter of glucose, diluted, and further expanded overnight in YSC-Leu containing 250 g/liter of glucose. Upon reaching OD600 = 2.5 to 3, 90 OD600 units were harvested, and cell pellets were resuspended in 4 ml of the seven indicated lignocellulosic hydrolysates supplemented with 20 mM urea and modified as follows: In the minimally altered control (blue), Ca(OH)2 was also added, if necessary, to achieve pH 5; in the inhibited control (red), feedstocks were toxified to the final concentrations of glucose, furfural, HMF, and acetic acid listed in table S2 (white columns) and adjusted to pH 5 with Ca(OH)2 if needed; and in the toxification and K+∕pH-amerilator condition (orange), feedstocks were toxified identically but adjusted instead with KHCO3, CaCO3, and Ca(OH)2 as listed in table S2 (white and gray columns). These three conditions were fermented with LAMy660, and the final was repeated with LAMy663 (gray). The ordering and color scheme of bars in Fig. 2B follow that of Fig. 2A. The “high concentration sugar syrup” nature of DMR corn stover as described by the provider was found in pilot experiments to be inhibitory to yeast (above-average concentrations of glucose and xylose were confirmed; see table S1); therefore, dilutions of 65 to 75% were necessary to enable fermentation (S1). All supplements were added in the maximally concentrated forms available to minimize dilution of the original hydrolysate. The minimally altered controls (blue) for bagasse, wheat straw, miscanthus, and switchgrass hydrolysate were observed in preliminary experiments to be sufficiently low in available sugar and ethanol product inhibition such that ethanol was metabolized following the consumption of glucose. To maximize accuracy of titer, cell-free fermentation samples for HPLC analysis were harvested after 24 hours under these four control conditions and 52 hours in all remaining.
For demonstrating extensibility of hydrolysate tolerance to xylose-consuming strains (Fig. 3A), prototrophic strains LAMY419 and LAMY665 were started in YSC–6 AA containing 40 g/liter of xylose, diluted, and further expanded overnight in YSC–6 AA containing 100 g/liter of xylose. Upon reaching OD$_{600}$ = 2.5 to 3, 90 OD$_{600}$ units were harvested, and cell pellets were resuspended in 4 ml of bagasse, high-acid corn stover, or miscanthus hydrolysate (nonbold) supplemented with 20 mM urea and modified in the same order and manner as those described above for Fig. 2B: Blue contained the Ca(OH)$_2$ needed to achieve pH 5 and was fermented with LAMY419; red toxified to the conditions listed in table S2 (white columns) and fermented with LAMY419; orange was toxified, K$^+$ and pH-adjusted to the conditions in table S2 (white and gray columns) and fermented with LAMY419; and gray was the same as orange but fermented with LAMY665.

Formulation and fermentation of synthetic medium (bold) was performed in a manner to reflect genuine lignocellulosic hexosepentose proportions but maintain the xylose metabolism that would be suppressed from catabolite repression. Specifically, LAMY419 was fermented in YSC–6 AA prepared with 50 g/liter of xylose, 150 g/liter of potato starch (Sigma-Aldrich, #S2630), and 60/140 mM KHCO$_3$/CaCO$_3$ (uninhibited control, blue), 62/48/100 mM furfural/HMF/acetic acid and adjusted to pH 5 with NH$_4$OH (inhibited control, red), or 62/48/100 mM furfural/HMF/acetic acid and 60/140 mM KHCO$_3$/CaCO$_3$ (orange). The final condition was repeated with LAMY665 (gray). For these four conditions, the harvested 90 OD$_{600}$ units were resuspended in 2.5 ml of medium and glucoamylase (Sigma-Aldrich, #A7095) added in the following amounts and times: 333 μl at 0 hour (+25 U/ml), 333 μl at 18 hours, 666 μl at 24 hours (+50 U/ml), and 666 μl at 36.5 hours. Enzyme amounts and times of addition were predetermined in pilot experiments to support the full consumption of xylose in the uninhibited control (see also fig. S8). Fermentation media were prepared in the most concentrated form possible to minimize dilution from glucoamylase addition. Cell-free fermentation samples for HPLC analysis were harvested after 24 hours for the uninhibited controls (blue) to minimize ethanol loss to consumption and 72 hours for the remaining conditions.

For demonstrating extensibility of hydrolysate tolerance to lactic acid–producing strains (Fig. 3B), prototrophic strains LAMY690 and LAMY692 were started in YSC–6 AA containing 100 g/liter of glucose, diluted, and further expanded overnight in YSC–6 AA containing 200 g/liter of glucose. Upon reaching OD$_{600}$ = 2.5 to 3, 90 OD$_{600}$ units of LAMY690 were harvested, and cell pellets were resuspended in 4 ml of YSC–Ura-Leu (bold) with 160 g/liter of corn stover, high-acid corn stover, and after 72 hours for toxified conditions. For prototrophic strains LAMY665 (gray). For these four conditions, the harvested 90 OD$_{600}$ units were resuspended in 2.5 ml of medium and glucoamylase (Sigma-Aldrich, #A7095) added in the following amounts and times: 333 μl at 0 hour (+25 U/ml), 333 μl at 18 hours, 666 μl at 24 hours (+50 U/ml), and 666 μl at 36.5 hours. Enzyme amounts and times of addition were predetermined in pilot experiments to support the full consumption of xylose in the uninhibited control (see also fig. S8). Fermentation media were prepared in the most concentrated form possible to minimize dilution from glucoamylase addition. Cell-free fermentation samples for HPLC analysis were harvested after 48 hours for the uninhibited controls (blue) to minimize ethanol loss to consumption and 72 hours for the remaining conditions.

Directed evolution of GRE2

We revived the PCR-mutagenized GRE2 yeast library from the functional variomics collection (gift of X. Pan of Baylor University) in YSC-Ura containing 30 g/liter of glucose and supplemented with 10 mM KH$_2$PO$_4$ (36). To maintain the >2 x 10$^5$ diversity, 20 μl of thawed cells were expanded to saturation, diluted, and cultured overnight in medium to approximately OD$_{600}$ = 2.5. Cell biomass 3.5 OD$_{600}$ units was harvested and resuspended in 4 ml of YSC-Ura containing 30 g/liter of glucose, 32/25/52 mM furfural/HMF/acetic acid, supplemented with 10 mM KH$_2$PO$_4$, and adjusted to pH 6 with NH$_4$OH. Following the time course of fig. S4, cell densities reaching OD$_{600}$ values of 5 to 8 were subcultured in identically formulated YSC-Ura medium containing the indicated combinations of inhibitors. A 1-ml aliquot of the final culture was harvested for isolation of bulk DNA and the remaining mixed to 15% glycerol for preservation at −80°C.

Several attempts were required to converge on the conditions used in the successful iteration depicted in fig. S4. For example, the 62/48/100 mM furfural/HMF/acetic acid combination used as our fermentation benchmark was determined to be growth suppressive even when pH-adjusted to neutralize acetic acid. A reduction to 30/24/50 mM furfural/HMF/acetic acid provided baseline growth-permissive conditions; however, increments of 5 to 10% of each inhibitor at the first subculturing ended in suppression. Given these responses, we hypothesized that prolonging stress combinations over multiple expansion cycles was needed to allow stronger mutants to entrench within the population and seed further advantageous trajectories. Last, that the hypertolerant GRE$_{2}^{evol}$ mutant was capable of conferring an improvement to fermentation under the benchmark suite of inhibitors (Fig. 2A), yet still unable to demonstrate growth under the same conditions (fig. S9), underscored the divergence in tolerance thresholds between metabolic and biomass production.

The coding sequence for the GRE$_{2}^{evol}$ allele is as follows: AT-GTCAGTTCTTCAATGTTAGTATCATGCATTACGCGCTCAATGTTA- CCAACACATTGTCGATCTCCTGTGAAAGAGACATTATAAGGTCATCTGGTTCTCGCAGAAGCTAAGAAAGGCGGAGAAATTAAACCGGGAC- CCGAGATTATTGAACCGGGCTTTGGTAACACATCTCAATATTCTCATTGGAAGGAAATTTGCCAAGAGAATTGTTTTTCCAAAGACAGCGGCAAAGAT- TAAAGATAGTCTACATACGCGCTTCCCCATCTGCTGTGTAATCAGCAGTGAAGCGGATTTATATTATCTCTGTGTTGAGACGTGTGTTAGTTTC- CACCTCTTTTTATAGCACTGTGGTTGGAACTCACTAACTTAAGAAATACCGCCGTGATTGTGAGAAGCTGTAGTTTCACACCTTCTTTATAGCACTGTG- GGTTAGAAATACGCGGACGAGCAGGCAAAGATATCGGTAATGAGGAGTGGTAATGAGGAGTGGTAATGAGGAGTGGTAATGAGGAGTGGTAATGAGGAGT

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Chromatography
Quantification of ethanol, lactic acid, glucose, xylose, glycerol, furfural, HMF, and acetic acid was performed on cell-free, 0.45-μm filtered samples using an Agilent 1200 Infinity Series HPLC configured with G1362A Refractive Index Detector and Aminex HPX-87H carbohydrate analysis column (Bio-Rad, #125-0140; www.bio-rad.com). Analytes were separated isocratically in 5 mM sulfuric acid at 65°C using a flow rate of 0.6 m/min. Under these conditions, retention times were approximately as follows: glucose, 9.2 min; xylose, 9.9 min; lactic acid, 13.1 min; glycerol, 13.8 min; acetic acid, 15.3 min; ethanol, 22.3 min; HMF, 30.2 min; furfural, 45.0 min. Chromatogram peaks autointegrated by the Agilent OpenLab CDS ChemStation software were converted to concentrations through interpolation off standard curves calibrated over the ranges of 0 to 150 g/liter of glucose, 0 to 50 g/liter of xylose, 0 to 100 g/liter of lactic acid, 0 to 8 g/liter of glycerol, 0 to 60 g/liter of acetic acid, 0 to 8 g/liter of HMF, and 0 to 8 g/liter of furfural defined from chemically pure dilution series. To compensate for the minor overlap between the peaks for glucose and xylose, we used standard concentrations incorporating the two sugars at a ratio of 3:1 g/liter of glucose:xylose to reflect typical proportions. Likewise, lactic acid and glycerol standards incorporated 10:1 g/liter of lactic acid:glycerol.

Given the likelihood of calcium salts precipitating from the low pH in the running solvent and obstructing instrument fluidic lines, all samples derived from calcium-containing fermentations were acidified with 1% sulfuric acid (vol/vol), rotated for ≥1 hour at 4°C, precipitates removed via centrifugation, and supernatants 0.45-μm filtered before HPLC analysis.

Mass spectrometry
For targeted quantification of furfural, FF-OH, HMF, and HMF-OH, cell-free samples collected from fermentation were extracted 1:1 with 100% UHPLC-grade methanol containing seventeen 13C-labeled amino acids (Cambridge Isotope Laboratories; www.isotope.com) added as internal standards and stored at −80°C for further processing. Amino acid–methanol extraction buffer, downstream method development, and analysis were provided by the Metabolite Profiling Core Facility at the Whitehead Institute (metabolomics.wi.mit.edu).

In brief, additional dilutions to 20-1 and 80-1 (final) in 50% methanol were required to reach the linear range of the spectrometer. Samples of 1 μl were injected into a Dionex UltiMate 3000 ultra-high performance liquid chromatography (UPLC) unit equipped with an Ascentis Express C18 (2.1 mm by 150 mm, 27 μm particle size) column (Sigma-Aldrich, #53825-U) maintained at 35°C. Analytes were reverse phase separated at a flow rate of 0.25 ml/min using buffers A [0.1% formic acid in liquid chromatography–mass spectrometry (LC–MS) grade water] and B (0.1% formic acid in LCMS grade acetonitrile) under the following gradient conditions: 0 to 2 min (5% B); 2 to 20 min (5 to 75% B, linear gradient); 20.1 to 24 min (95% B); 24.1 to 28 min (5% B). Mass analysis was performed on a Thermo Fisher Scientific Q Exactive Orbitrap operating with a spray voltage of 3.0 kV, capillary temperature 275°C, HESI probe temperature 350°C, sheath gas flow 40 units, auxiliary gas flow 15 units, and sweep gas flow 1 unit. For targeted isolation of furfural, HMF, FF-OH, and HMF-OH, positive ionization mode was used with resolution set to 70,000, automatic gain control to 1×105 with maximum injection time of 250 ms, and isolation window to 4.0 m/z. Fragmentation patterns from MS/MS were matched with reference spectra available in the METLIN online database (metlin.scripps.edu). Quantification was performed using the Thermo Fisher Scientific Xcalibur Software calibrated against chemically pure dilution series of 3 μM to 3 mM furfural, 30 μM to 3 mM FF-OH, 0.1 μM to 3 mM HMF, and 30 μM to 3 mM HMF-OH.

For measurement of salt concentrations in hydrolysates, 10 ml of each sample was centrifuged (3500 rpm for 5 min) to remove large particulates, and the supernatants submitted to Environmental Testing and Research Laboratories (etrlabs.com) for quantification of K⁺ and Ca²⁺ (available as components of their water testing suite). Acid-digested samples were assayed in three replicate reads by inductively coupled plasma mass spectrometry.

Viability measurements
Yeast population viabilities measured via methylene blue staining and subsequent procedures to calculate correlation with ethanol titers were described previously (16).

Statistical analysis
Calculation of SD, propagation of error, hypothesis testing (two-sample, two-tailed t test, α = 0.05), and value determination were performed using MATLAB (The MathWorks; www.mathworks.com) on independent biological triplicates following standard procedures.

SUPPLEMENTARY MATERIALS
Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/7/26/eabf7613/DC1

View/request a protocol for this paper from Bio-protocol.

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Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Corn stover–derived hydrolysates can be provided by the U.S. National Renewable Energy Laboratory pending scientific review and a completed material transfer agreement. Requests should be directed to M.G.R. Wheat straw hydrolysate can be provided by Versalis (Eni) pending a similar review and agreement. Inquiries should be submitted to info.licensing@versalis.eni.com. Correspondence and requests for strains, plasmids, and other materials can be addressed to G.S. or G.R.F. Additional data related to this paper may be requested from the authors.

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Engineered yeast tolerance enables efficient production from toxified lignocellulosic feedstocks
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