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Industrial Systems Biology of *Saccharomyces cerevisiae* Enables Novel Succinic Acid Cell Factory

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Abstract

*Saccharomyces cerevisiae* is the most well characterized eukaryote, the preferred microbial cell factory for the largest industrial biotechnology product (bioethanol), and a robust commercially compatible scaffold to be exploited for diverse chemical production. Succinic acid is a highly sought after added-value chemical for which there is no native pre-disposition for production and accumulation in *S. cerevisiae*. The genome-scale metabolic network reconstruction of *S. cerevisiae* enabled *in silico* gene deletion predictions using an evolutionary programming method to couple biomass and succinate production. Glycine and serine, both essential amino acids required for biomass formation, are formed from both glycolytic and TCA cycle intermediates. Succinate formation results from the isocitrate lyase catalyzed conversion of isocitrate, and from the α-keto-glutarate dehydrogenase catalyzed conversion of α-keto-glutarate. Succinate is subsequently depleted by the succinate dehydrogenase complex. The metabolic engineering strategy identified included deletion of the primary succinate consuming reaction, Sdh3p, and interruption of glycolysis derived serine by deletion of 3-phosphoglycerate dehydrogenase, Ser3p/Ser33p. Pursuing these targets, a multi-gene deletion strain was constructed, and directed evolution with selection used to identify a succinate producing mutant. Physiological characterization coupled with integrated data analysis of transcriptome data in the metabolically engineered strain were used to identify 2nd-round metabolic engineering targets. The resulting strain represents a 30-fold improvement in succinate titer, and a 43-fold improvement in succinate yield on biomass, with only a 2.8-fold decrease in the specific growth rate compared to the reference strain. Intuitive genetic targets for either over-expression or interruption of succinate producing or consuming pathways, respectively, do not lead to increased succinate. Rather, we demonstrate how systems biology tools coupled with directed evolution and selection allows non-intuitive, rapid and substantial re-direction of carbon fluxes in *S. cerevisiae*, and hence show proof of concept that this is a potentially attractive cell factory for over-producing different platform chemicals.

Introduction

Industrial biotechnology is a promising alternative to traditional petrochemical production of chemicals focused on developing commercially sustainable and environmentally favorable processes [1]. Metabolic engineering, the directed genetic modification of cellular reactions, aims to change the metabolic architecture of microorganisms to efficiently produce target chemicals [2]. Although examples of metabolic engineering successes exist, there has yet to be developed a pipeline where preferred industrial hosts are rapidly engineered to produce a non-native accumulating target metabolite. Recent advances in systems biology enabled *in silico* genome-scale metabolic network reconstructions to guide metabolic engineering strategies [1], [3], [4]. Here we describe a pipeline where a microbial strain was constructed, physiologically characterized, and genomic tools were used to verify the predictions. An essential part of the pipeline is the use of genome-scale metabolic models for initial guiding of the metabolic engineering, which has been shown to be useful also in earlier studies [3], [5], [6]. This approach was repeated and complemented with traditional directed evolution and selection until a proof of concept microbial cell factory was reached. This pipeline resulted in the construction of a non-intuitive *Saccharomyces cerevisiae* cell factory over-producing succinic acid, a building block chemical.

*S. cerevisiae* is the most well characterized eukaryote and is unique in its broad application as an industrial production platform for a large portfolio of products including foods and beverages, bioethanol, vaccines, and therapeutic proteins [1].
Many systems biology tools, including high-throughput genome sequencing, transcriptional profiling, metabolomics, carbon flux estimations, proteomics, \textit{in silico} genome-scale modeling, and bioinformatics driven data integration were first applied to \textit{S. cerevisiae} \cite{4}. Metabolic engineering has benefited from each of these tools; however, relatively few examples exist where cumulative integration has resulted in a generalized pipeline, in particular for the production of a target compound that the organism does not accumulate significantly naturally.

Succinic acid, systematically identified as butanedioic acid (pKa1 4.21, pKa2 5.72), is a value-added chemical building block, with an estimated 15,000 t/year world-wide demand predicted to expand to commodity chemical status with 270,000 t/year \cite{7}, \cite{9} representing a potential >2 billion USD annual market. There are several elegant examples of bio-based production of succinate in \textit{Anaerobiospiillum succiniciproducens}, \textit{Acinetobacter succinogenes}, \textit{Succinobiostrbx terrosolvens}, \textit{Corynebacterium glutamicum}, \textit{Prevotella ruminocola}, a recently isolated bacterium from bovine rumen, \textit{Mannheimia succiniciproducens}, and a metabolically engineered succinic acid over-producing \textit{E. coli} \cite{7}, \cite{9}, \cite{10}, \cite{11}, \cite{12}. All of the hosts described are prokaryotic that grow at neutral pH, and consequently secrete the salt, succinate, requiring a cost-intensive acidification and precipitation to reach the desired succinic acid. This concern is not specific to succinic acid production, but rather universal when considering organic acid producing microbial cell factories \cite{13}. \textit{S. cerevisiae} represents a well-established, generally regarded as safe, robust, scalable (1L to 100,000L) industrial production host capable of growth on diverse carbon sources, chemically defined medium, both aerobic and anaerobic, and a wide pH operating range (3.0–6.0). Unlike the hosts described above, succinate is not naturally produced by \textit{S. cerevisiae}; but as there are many factors of importance for the choice of a microbial cell factory it is not uncommon that the chosen cell factory lacks predisposition to produce the target chemical of choice \cite{14}. As industrial biotechnology progresses forward, and the concept of biorefineries are gaining increased importance, platform technology enabled \textit{S. cerevisiae} are increasingly being considered for the production of succinate and other organic acids, with the yeast \textit{S. cerevisiae} being particularly investigated due to its versatility, well characterized genome, and recognized ability to produce succinate \cite{15}. \textit{S. cerevisiae} is uniquely positioned as a platform technology as it is already used widely for bioethanol production, but also because of the extensive library of genetic engineering tools, a very well annotated genome, many omics tools available, and well established complimentary approaches for directed evolution and selection. We therefore addressed the question whether it is possible to metabolically engineer \textit{S. cerevisiae} such that the carbon fluxes are redirected towards succinic acid, and hereby establish proof-of-concept of using this yeast as a general cell factory platform for chemical production.

The final strain emerging from this study requires a significant further metabolic engineering and process development prior to commercialization, but the approach and integration of methods demonstrated supports the hypothesis that highly regulated central carbon metabolism in ideal production hosts can be reconfigured to produce target chemicals, relatively quickly and with minimal resources.

\textbf{Results}

In the present study, our objective was to evaluate the use of genome-scale metabolic models for \textit{in silico} identification of gene deletion targets. We therefore used results from a previous study where an evolutionary programming method, termed OptGene, was developed for identification of deletion targets to couple biomass formation (or other biological objective function that is applicable to the organism under question) with the design objective function, such as yield (or other linear/non-linear objective) \cite{16}. These results guided the metabolic engineering strategy described in Figure 1. Glycine, serine, and threonine, all representing essential amino acids required for biomass formation, may be formed from both glycolytic and tricarboxylic acid cycle intermediates. Succinate formation results from the isocitratelyase, Icl1p, catalyzed conversion of isocitrate to equimolar glyoxylate and succinate, and from the 2-keto-gluconate dehydrogenase complex, Kgd1p/Kgd2p/Lpd1p, catalyzed conversion of 2-keto-gluconate to equimolar succinate, with a net production of CO\textsubscript{2}, NADH, and ATP. Succinate is subsequently depleted by the succinate dehydrogenase complex, Sdh1p/Sdh2p/Sdh3p/Sdh4p to equimolar fumarate with the net production of protonated ubiquinone. The metabolic engineering strategy identified by OptGene included deletion of the primary succinate consuming reaction encoded by Sdh3 (cytochrome b subunit of the succinate dehydrogenase complex, essential for function), and interruption of glycolysis derived serine by deletion of 3-phosphoglycerate dehydrogenase, Ser3p/Ser33p (isoenzymes).

The remaining pathway for serine synthesis must originate from glycine, and glycine synthesis is largely derived from the alanine-pyruvate aminotransferase, Agx1p, converting glyoxylate and alanine to glycine and pyruvate. With this strategy, glycine and serine biomass requirements are directly coupled to succinate formation via the glyoxylate cycle. Substantial succinate accumulation (defined as >0.1 g \text{L}^{-1}) in the culture broth is not observed in wild-type \textit{S. cerevisiae}, and deletion of \textit{sdh3} has not resulted in appreciable succinate accumulation \cite{17}; a conclusion also found by chemical inhibition of the succinate dehydrogenase complex with titration of malonate (Figure S1), a chemical inhibitor of this complex \cite{10}.

The mutant resulting from the \textit{in silico} strategy, referred to as 8D (\textit{sdh3} \textit{ser3} \textit{sdh3}3), required supplementation with 500 mg \text{L}^{-1} glycine to be able to grow. When evaluated in well controlled, aerobic, batch stirred reactor cultures supplemented with 20 g \text{L}^{-1} glucose in chemically defined medium, it exhibited a 13-fold improvement in succinate secreted titer (0.03 v 0.40 g \text{L}^{-1}), 14-fold improvement in succinate biomass yield (0.01 v 0.14 g-succinate g-biomass \textsuperscript{-1}), and a modest 33% reduction in the specific growth rate. Thus, the \textit{in silico} guided metabolic engineering strategy was shown to work, representing a proof-of-concept of the use of model guided metabolic engineering. However, in order to obtain a prototrophic strain directed evolution was employed to screen and select for 8D mutants that did not require glycine supplementation. Specifically, six consecutive shake flask cultivations in media supplemented with decreasing concentrations of glycine, from an initial 500 mg \text{L}^{-1} to 0 mg \text{L}^{-1} (see Figure 2) were performed. The resulting strain demonstrated a 7.7-fold improvement in succinate yield on biomass (0.09 v 0.69 g-succinate g-biomass \textsuperscript{-1}), strongly suggesting the direct coupling of glycine formation from glyoxylate and succinate formation. The resulting strain had a relatively low specific growth rate, 0.03 h \textsuperscript{-1}, and was therefore subsequently cultivated in shake flasks and transferred across six shake flasks (only first three shake flasks shown in Figure 2) to improve the specific growth rate. Finally, a specific growth rate of 0.14 h \textsuperscript{-1} was reached, however, resulting in a decreased succinate yield on biomass (0.69 v 0.27 g-succinate g-biomass \textsuperscript{-1}). The final strain, referred to as 8D Evolved, was shown to exhibit a 60-fold improvement in biomass coupled succinate production (0.01 v 0.30 g-succinate g-biomass \textsuperscript{-1}), and 20-fold improvement in succinate titer (0.03 v 0.60 g \text{L}^{-1}) relative to the reference strain when grown in aerobic batch cultivations.
To investigate the apparent decoupling of succinate coupled biomass formation, and potentially identify second-round metabolic engineering strategies, the transcriptome was measured in aerobic, glucose-limited, mid-exponential phase grown batch cultivations of 8D Evolved and the reference strain. Continuous cultivations, both carbon-limited and nitrogen-limited chemostats were attempted with the 8D Evolved mutant; however, in both cases steady-state was not reached and wash-out occurred, even at relatively low dilution rates ($D = 0.03$ h$^{-1}$ compared to $\mu_{\text{max}} = 0.14$ h$^{-1}$). It was expected that 8D Evolved would not support cultivation in carbon-limited continuous culture due to the down-regulation of the TCA cycle ($sdh3$), and consequently, reduced capacity for respiratory metabolism and oxidative phosphorylation. Therefore, batch cultivations were employed acknowledging the significant differences in specific growth rate (0.33 v 0.13 h$^{-1}$), and glucose uptake rate (90 v 26 C-mmol g$^{-1}$DCW h$^{-1}$), while maintaining relatively similar biomass yields (0.18 v 0.19 C-mol biomass C-mol glucose$^{-1}$).

Several studies have shown that significant differences in specific growth rate directly impact transcriptome interpretation, with anywhere between 268 and 2400 genes classified as potentially growth-related [19], [20], [21]. Previously generated continuous cultivation transcriptome data for both carbon-limited (glucose, respiratory growth) and nitrogen-limited (ammonium sulfate, respiro-fermentative growth) conditions at dilution rates of 0.03, 0.1, and 0.2 h$^{-1}$ were therefore used to identify statistically differentially expressed growth-related genes [21]. A total of 6 and 7 differentially expressed genes were identified within the carbon-limited and nitrogen-limited data sets as being growth-related ($p$-value$<0.1$, $n=3$ at each dilution rate), respectively, and a total of 66 differentially expressed genes were identified when comparing carbon-limited and nitrogen-limited data sets, paired at each dilution rate ($p$-value$<0.1$, $n=3$ at each dilution rate). Of the total 2406 differentially expressed genes between the 8D Evolved and reference strain ($p$-value$<0.01$, $|\text{log-fold change}|>0.5$, $n=3$ biological replicates, $n=2$ DNA microarray duplicates), 36 unique growth-related genes were identified suggesting that few of the genes with a significant change in transcription in 8D Evolved are due to changes in the specific growth rate (see Figure 3). However, a total of 8 of the top 20 $p$-value$<0.1$ ranked differentially expressed genes identified from pair-wise comparison of 8D Evolved and the reference strain, are growth-related genes ($ARO9$, $SER3$, $JLP1$, $HMALPHA1$, $ARO10$, $MFALPHA2$, and two uncharacterized genes, YPL033c and YLR267w).

The top 2000 (there were no metabolic genes in the remaining 406 genes nor were there any biological process annotations available as determined by gene ontology, and therefore they were not included in further analysis) differentially expressed genes were selected for further analysis, and after removal of the 36 growth-related genes, a list of 1964 genes was submitted for metabolic
pathway visualization and characterization to the Expression Viewer [22] available at the Yeast Genome Database [23] (see Figure 3). The log-fold change of the 8D Evolved:Reference expression ratio was mapped onto the metabolic map of S. cerevisiae strain S288c, version 12.0, composed of 140 pathways, 925 enzymatic reactions, and a total of 675 compounds (see Figure S2). A total of 315 genes mapped to a specific metabolic pathway on the expression viewer, with a mean log-fold expression ratio value of 0.3 ± 0.3 (n = 315, ± SD).

Three biological insights were immediately apparent (see Figure S2). First, SDH3, SER3, and SER33 had negative log-fold expression ratios (log-fold change, -8.0) confirming the gene deletions targeted in the 8D strain and the maintained low expression through the directed evolution. Second, when examining the glycine, serine, and threonine metabolism, AGX1 was 4.3 log-fold change upregulated in the 8D Evolved strain, confirming significant upregulation of glycine synthesis from glyoxylate pools, as predicted by the original metabolic engineering strategy. However, there was no upregulation of SHM2, SHM1, the genes encoding pathways for L-serine formation from L-glycine pools.

Most surprisingly GLY1, encoding threonine aldolase, was significantly up-regulated (log-fold change 1.6). In the genome-scale metabolic network reconstructions of S. cerevisiae iFF708, iMM904, and iND750, upon which the 8D metabolic engineering strategy is based, Gly1p encodes the irreversible conversion of glycine and acetaldehyde to threonine [24], [25], [26], leading to the prediction that threonine biosynthesis from glycolytic intermediates could be down-regulated, and provided for from glycine pools. This consequently leads to a greater biomass-coupled drive for glyoxylate synthesis from isocitrate, yielding equimolar succinate. Leveraging this over-all strategy, another S. cerevisiae mutant was constructed, referred to as 20G (Δsdh3, Δser3, Δthr1), where Thr1p, encoding homoserine kinase that is required for threonine biosynthesis, was deleted. However, this strain required threonine supplementation and after several extensive attempts at adaptive evolution, the threonine auxotrophy persisted, suggesting the irreversibility of Gly1 was incorrect and the aldolase strongly favors glycine formation (Figure S3). The significant up-regulation of Gly1 therefore provides a strong hypothesis for why 8D Evolved had an attenuation of succinate production, even under increasing specific growth rate, suggesting a decoupling of biomass coupled succinate production. It should be noted that in the most recent update of the genome-scale metabolic reconstruction of S. cerevisiae, iMM904, the directionality of Gly1 was corrected to now indicate threonine irreversible conversion to glycine and acetaldehyde [27].

The transcriptome not only provides for a global, rapid, and quantitative assessment of the predicted in silico metabolic engineering strategy and insight into the genetic engineering modifications that result from directed evolution and selection, but also provides a source for identification of second round metabolic engineering targets not previously predicted. Several targets were identified, but of particular interest was ICL1, encoding isocitrate lyase, converting isocitrate to glyoxylate and succinate in equimolar concentrations. All tricarboxylic acid cycle genes are up-regulated, with the exception of SDH3 (target gene deletion), and ICL1, providing a clear metabolic engineering target for up-regulation in the 8D Evolved strain. Therefore, native ICL1 was PCR amplified and cloned into the 2 μm ori plasmid pRS426CT containing the strong constitutive TEF1 promoter and CYC1 terminator [28], and then transformed into the reference, 8D, and 8D Evolved strain (strains transformed with the constructed plasmid pRS426T-ICL1-C are referred to as “with pICL1”).
strains were evaluated in aerobic, glucose-supplemented batch fermentations, and only 8D Evolved with pICL1 exhibited a change in succinate production (see Figure 4). Specifically, the succinate titer, biomass yield, and glucose yield were 0.90 g L$^{-1}$, 0.43 g-succinate g-biomass$^{-1}$, and 0.05 g-succinate g-glucose$^{-1}$, respectively, representing a 1.5-fold, 1.4-fold, 1.7-fold improvement over 8D, respectively (see Figure 4).

**Discussion**

A *S. cerevisiae* strain capable of succinate production, requiring redirection of carbon flux from typically produced C$_2$ (ethanol, acetate) and C$_3$ (glycerol, pyruvate) overflow metabolites to the target C$_4$ succinic acid was achieved through metabolic engineering, requiring integration of systems biology methods and directed evolution. Clearly, the resulting strain (8D Evolved with pICL1), while being a successful demonstration of a multi-round metabolic engineering strategy, still requires significant process development and further enhancement to compete commercially with existing bacterial platforms.

The resulting strain, 8D Evolved with pICL1, represents a 30-fold improvement in succinate titer, and a 43-fold improvement in succinate yield on biomass, with only a 2.8-fold decrease in the specific growth rate compared to the reference strain. Despite success of using simple stoichiometric-based calculations for driving metabolic engineering, it is interesting to note that regulatory mechanisms not captured in these models are likely playing a significant role in the succinate production observed. The biomass requirements for glycine and serine are 0.290 and 0.185 mmol g-DCW$^{-1}$ [24]. Assuming that all glycine, and all glycine and serine combined demands are supplied from the glyoxylate pool, then the theoretical production of succinate would amount to 0.034 and 0.056 g-succinate g-DCW$^{-1}$, respectively. The 8D and 8D Evolved strains are producing 0.30 and 0.43 g-succinate g-biomass$^{-1}$, respectively, suggesting a nearly 8-fold higher succinate production than required to meet biomass amino acid demands. A potentially 3rd metabolic engineering target would be deletion of *GLY1* to further minimize alternative biosynthetic routes of glycine production, thereby isolating all glycine production to be dependent on glyoxylate formation, and consequently succinate formation. Yet, it’s clear that any increase in succinate formation would not be due to biomass requirements, but rather regulatory (e.g., non-stoichiometric driven) mechanisms. Therefore, while the strategy presented and demonstrated here is likely to be a major component of an overall succinate production cell factory, complimentary strategies focusing on the other major succinate production pathway, TCA cycle, will be required. Examples of malic acid production, that included
engineering of pyruvate carboxylation (overexpression of \textit{PYC2}), oxaloacetate reduction (overexpression of \textit{MDH3}), and malate export (functional expression of the non-native \textit{SpMAE1}), resulted in a \textit{S. cerevisiae} strain capable of producing 59 g-malate L\(^{-1}\) and 0.42 mol malate mol-glucose\(^{-1}\) [29]. A similar approach, requiring yet further engineering and understanding of the reductive TCA cycle to convert malate to succinate is likely required, but a major hurdle with this strategy is the conversion of fumarate to succinate by fumarate reductase which is thermodynamically favoured in the direction of fumarate.

The transcriptome analysis performed, specifically consideration of continuous culture data sets at different dilution rates to filter growth-related genes was an integral part of identifying the 2\(^{nd}\) round of metabolic engineering targets. Although a relatively small number of growth related genes were identified, they were of high-value. For example, \textit{ARO9} and \textit{ARO10}, encoding key enzymatic conversion steps in aromatic amino acid metabolism may have incorrectly pointed towards tryptophan, tyrosine, and phenylalanine catabolism or phosphoenolpyruvate decarboxylase activity as metabolic areas of interest for understanding physiological differences between 8D Evolved and the reference strains. This approach may be extended to future efforts and other organisms, where continuous cultivation of engineered strains may not be possible, as in this case, or applied industrially where the dominant and preferred processing mode is batch.

Furthermore, this work clearly demonstrated that obvious genetic targets did not result in increased succinate formation. Specifically, deletion of the primary succinate consuming pathway (\textit{Dsdh2}) [17], or constitutive over-expression of one of two of the primary succinate formation pathways (\textit{ICL}) did not result in any increased succinate production (See Figure S1). It is further interesting to note that the 8D with pICL1 strain also did not

**Figure 4. Summary of succinate microbial cell factory construction.** The specific growth rate (1/h), maximum succinate titer (g/L), maximum succinate yield on biomass (g/g-biomass), and maximum yield on glucose (g/g-glucose) are reported for the reference strain, 8D, 8D evolved, and 8D evolved with pICL1. A 43-fold improvement in succinate yield on biomass was observed across the full cycle of metabolic engineering that included \textit{in silico} guided approaches, directed evolution, and transcriptome based identification of a 2\(^{nd}\) round of metabolic engineering targets.

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result in any increased succinate production, but rather only in the 8D Evolved with pICL1 strain. The ability to measure transcriptome on a strain that underwent targeted genetic engineering and directed evolution was critical to identifying pICL1 as a 2nd metabolic engineering target, which would have been discarded if selected based on intuition.

The approach employed represents an integration of diverse methods for rapid metabolic engineering proof-of-concept. The strain selection process thus need not be limited to considering organisms showing a predisposition to the production of the metabolite of interest, but rather, should include hosts most suitable for large-scale, robust, and biorefinery processing. With such hosts, carbon and redox flux redistribution requiring multi-gene approaches can be predicted, tested, and supplemented with directed evolution, screening, and selection. These strains are then transcriptionally characterized and optimized until commercially viable titers, productivities, and yields are reached. It is only through whole-process optimization and elimination of severe constraints such as forced use of non-industrially favorable strains, that the promise of a bio-based economy may be fully realized.

Materials and Methods

Strain Construction

The reference strain Saccharomyces cerevisiae CEN.PK113-5D (Mat a MAL2-8C SUC2 UR3-32) [30] was used for the construction of the Adsh3 Asn3 Aso33 knockout strain, referred to as the 8D mutant, and for the construction of the Adsh3 Asn3 Abi1 knockout strain, referred to as the 20G mutant, through the cloning-free PCR-based allele replacement method previously described [31].

The upstream SDH3 fragment was amplified by PCR from genomic DNA using the primers SDH3_Up_Fw (sequence 5'-GACCGTTAACGG-3') and SDH3_Up_Rv (sequence 5'-TGTTATTATTCTTAGAGC-3'). Similarly, the downstream SDH3 fragment was amplified by PCR from genomic DNA using the upstream primers SDH3_Dw_Fw (sequence 5'-CAGGGATGGCGGCCGCTGACGACATCGTTTATTTATCTTCTTAGGAC-3') and SDH3_Dw_Rv (sequence 5'-GTAATCTGTTATCGATAATCTGCC-3').

The upstream ICL1 fragment was amplified by PCR from genomic DNA using the primers ICL1_Up_Fw (sequence 5'-GACCGTTAACGG-3') and ICL1_Up_Rv (sequence 5'-TGTTATTATTCTTAGAGC-3'). Similarly, the downstream ICL1 fragment was amplified using the upstream primers ICL1_Dw_Fw (sequence 5'-CAGGGATGGCGGCCGCTGACGACATCGTTTATTTATCTTCTTAGGAC-3') and ICL1_Dw_Rv (sequence 5'-GTAATCTGTTATCGATAATCTGCC-3').

The upstream THR1 fragment was amplified by PCR from genomic DNA using the primers THR1_Up_Fw (sequence 5'-GACCGTTAACGG-3') and THR1_Up_Rv (sequence 5'-TGTTATTATTCTTAGAGC-3'). Similarly, the downstream THR1 fragment was amplified using the primers THR1_Dw_Fw (sequence 5'-CAGGGATGGCGGCCGCTGACGACATCGTTTATTTATCTTCTTAGGAC-3') and THR1_Dw_Rv (sequence 5'-GTAATCTGTTATCGATAATCTGCC-3').

The upstream CATGTTAGG-3' and SER33_Dw_Rv (sequence 5'-CTTTTATTTCTTAGGACGATCGCTTGACCAGAAGGTGCTTCGG-3') and SER33_Dw_Rv (sequence 5'-CTTTTATTTCTTAGGACGATCGCTTGACCAGAAGGTGCTTCGG-3'). The lithium acetate transformation method was used [32]. As described previously, URA3 from Kluyveromyces lactis was used as the selection marker in the transformation process [31]. With this approach transformants are easily selected on uracil depleted media supplemented with 5-fluoroorotic acid. The knockout was confirmed by restriction analysis followed by sequencing (MWG Biotech AG, Ebersberg, Germany).

The plasmid pRS426T-ICL1-C was constructed and transformed into 8D Evolved, described earlier and used for constitutive S. cerevisiae ICL1 overexpression. The parent plasmid, pRS426CT (6347 bp), was previously constructed in our laboratory by inserting the strong constitutive TEF1 promoter (gene encoding S. cerevisiae translation-elongation factor 1α) and the CYC1 transcription terminator into pRS426 [28]. This original backbone plasmid is a 5726 bp yeast episomal plasmid (YEp-type) that includes the TEF1 promoter and the CYC1 transcription terminator sequence, verified by sequencing (MWG Biotech AG, Ebersberg, Germany).

A total of eight primers were required for amplification of the native ICL1 gene from the reference strain, sequencing of the constructed plasmid pRS426-ICL1-C, and PCR to verify plasmid presence in the transformed reference and 8D Evolved strains (referred to as 8D Evolved with pICL1). The PCR amplification of ICL1 was carried out using the Phusion™ High-Fidelity DNA Polymerase (Finzymes Oy, Espoo, Finland) according to the manufacturer’s protocol. The native ICL1 was amplified from genomic DNA using the up- and downstream primers ICL1_Sp1 (sequence 5'-GCTCCGCCTCAAATGCAAGGCAATTCTTG-11-29) and ICL1_Sp2 (sequence 5'-GCTCCGCCTCAAATGCAAGGCAATTCTTG-11-29), respectively. The amplicon length was 1915bp. The fragment was cut with restriction endonucleases (REN) SpeI, the restriction site of which was de novo introduced on primer ICL1_Sp1, and NgoMIV, and then ligated with pRS426CT cut with SpeI and XmaI. By using the non-compatible RENs in either end of the insertion, the direction of the insert is secured and furthermore the sole parent plasmid Xma site is lost. This allowed for an in vitro pre-selection for the correct pRS426-ICL1-C construct prior to transformation. The four sequencing primers for construct verification included M13_rev-29 (sequence 5'-GAGGAAACAGCTATTAGCAGC-3'), ICL1_In_1f (sequence 5'-GGTTTCTGGCATGTCATC-3'), ICL1_In_2f (sequence 5'-CATCCGGAGAGAGGGGACAG-3'), and M13_uni_-21 (sequence 5'-TCTGATCAGTGGATGTCAGTGGATGTCAGTGGATGTC-3'). The two primers used for plasmid verification via PCR (Taq DNA Polymerase of Thermus aquaticus from Sigma, St. Louis, MO, were ICL1_partSense (sequence 5'-TTCCGTTTCTGAGATTTCTGAAATCGGC-3') and ICL1_CYC_AntiSense (sequence 5'-AAATTAAAGCCGCTGAGGCTGG-3') and these were used for analytical PCRs according to the instruction manual’s recommendations). Plasmid transformation of electroporment E. coli DH5α were completed as described previously, as was plasmid transformation of the S. cerevisiae reference strain and 8D Evolved using the lithium acetate method [28], [31], [32].

Medium Formulation

A chemically defined minimal medium of composition 5.0 g L⁻¹ (NH₄)₂SO₄, 3.0 g L⁻¹ KH₂PO₄, 0.5 g L⁻¹ MgSO₄•7H₂O,
1.0 mL L\(^{-1}\) trace metal solution, 300 mg mL\(^{-1}\) uracil, 0.05 g L\(^{-1}\) antifoam 204 (Sigma-Aldrich A-8311), and 1.0 mL L\(^{-1}\) vitamin solution was used for all shake flask and 2L well-controlled fermentations [34]. The trace element solution included 15 g L\(^{-1}\) EDTA, 0.45 g L\(^{-1}\) CaCl\(_2 \cdot 2\)H\(_2\)O, 0.45 g L\(^{-1}\) ZnSO\(_4 \cdot 7\)H\(_2\)O, 0.3 g L\(^{-1}\) FeSO\(_4 \cdot 7\)H\(_2\)O, 100 mg L\(^{-1}\) H\(_2\)BO\(_3\), 1 g L\(^{-1}\) MnCl\(_2 \cdot 2\)H\(_2\)O, 0.3 g L\(^{-1}\) CoCl\(_2 \cdot 6\)H\(_2\)O, 0.3 g L\(^{-1}\) CuSO\(_4 \cdot 5\)H\(_2\)O, 0.4 g L\(^{-1}\) NaMoO\(_4 \cdot 2\)H\(_2\)O. The pH of the trace metal solution was adjusted to 4.00 with 2 M NaOH and heat sterilized. The vitamin solution included 50 mg L\(^{-1}\) d-biotin, 200 mg L\(^{-1}\) pama-mino benzoic acid, 1 g L\(^{-1}\) nicotinic acid, 1 g L\(^{-1}\) Carpantho- nate, 1 g L\(^{-1}\) pyridoxine HCl, 1 g L\(^{-1}\) thiamine HCl, and 25 mg L\(^{-1}\) 2-minositol. The pH of the vitamin solution was adjusted to 6.5 with 2 M NaOH, sterile-filtered and the solution was stored at 4°C. The final formulated medium, excluding glucose and vitamin solution supplementation, is adjusted to pH 5.0 with 2 M NaOH and heat sterilized. For carbon-limited cultivations the sterilized medium is supplemented with 20 g L\(^{-1}\) glucose, heat sterilized separately, and 1.0 mL L\(^{-1}\) vitamin solution is added by sterile filtration [0.20 µm pore size Ministar®-Plus Sartorius AG, Goettingen, Germany]. For cultures where glycerol or threonine auxotrophic strains were cultivated the final culture medium was supplemented with glycerine 500 mg L\(^{-1}\) or 100 mg L\(^{-1}\) threonine added by sterile filtration.

**Shake Flask Cultivations and Stirred Tank Fermentations**

Shake flask cultivations were completed in 500 mL Erlenmeyer flasks with two diametrically opposed baffles and two side-necks with septums for sampling by syringe. Flasks were heat sterilized with 100 mL of medium, inoculated with a single colony, and incubated at 30°C with orbital shaking at 150 RPM. Stirred tank fermentations were completed in well-controlled, aerobic, 2.2L Braun Biotech Biostat B fermentation systems with a working volume of 2L (Sartorius AG, Goettingen, Germany). The temperature was controlled at 30°C. The fermenters were outfitted with two disk-turbine impellers rotating at 600 RPM. Dissolved oxygen was monitored with an autoclavable polarographic oxygen electrode (Mettler-Toledo, Columbus, OH). During aerobic cultivation the air sparging flow rate was 2 vvm. The pH was kept constant at 5.0 by automatic addition of 2 M KOH. Off-gas passed through a condenser to minimize the evaporation from the fermenter. The fermenters were inoculated from shake flask precultures to an initial OD\(_{600}\) 0.01.

**Fermentation Analysis**

**Off-gas Analysis.** The effluent fermentation gas was measured every 30 seconds for the determination of O\(_2\)(g) and CO\(_2\)(g) concentrations by the off-gas analyzer Bruel and Kjær 1308 (Bruel & Kjær, Nørrum, Denmark).

**Biomass Determination.** The optical density (OD) was determined at 600 nm using a Shimadzu UV mini 1240 spectrophotometer (Shidamazu Europe GmbH, Duisberg, Germany). Duplicate samples were diluted with deionized water to obtain OD\(_{600}\) measurements in the linear range of 0-0.4 OD\(_{600}\). Samples were always maintained at 4°C post-sampling until OD\(_{600}\) and dry cell weight (DCW) measurements were performed. DCW measurements were determined through the exponential phase, until stationary phase was confirmed according to OD\(_{600}\) and off-gas analysis. Nitrocellulose filters (0.45 µm Sartorius AG, Goettingen, Germany) were used. The filters were pre-dried in a microwave oven at 150W for 10 min., and cooled in a desiccator for 10 min. 5.0 mL of fermentation broth were filtered, followed by 10 mL DI water. Filters were then dried in a microwave oven for 20 min. at 150W, cooled for 15 min. in a desiccator, and the mass was determined.

**Metabolite Concentration Determination.** All fermentation samples were immediately filtered using a 0.45 µm syringe-filter (Sartorius AG, Goettingen, Germany) and stored at −20°C until further analysis. Glucose, ethanol, glycerol, acetate, succinate, pyruvate, fumarate, citrate, oxalate, and malate were determined by HPLC analysis using an Aminex HPX-87H ion-exclusion column (Bio-Rad Laboratories, Hercules, CA). The column was maintained at 65°C and elution performed using 5 mM H\(_2\)SO\(_4\) as the mobile phase at a flow rate of 0.6 mL min\(^{-1}\). Glucose, ethanol, glycerol, acetate, succinate, citrate, fumarate, malate, oxalate were detected on a Waters 410 differential refractometer detector (Shodex, Kawasaki, Japan), and acetate and pyruvate were detected on a Waters 468 absorbance detector set at 210 nm.

**Transcriptomics**

**RNA Sampling and Isolation.** Samples for RNA isolation from the late-exponential phase of glucose-limited batch cultivations were taken by rapidly sampling 25 mL of culture into a 50 mL sterile Falcon tube with 40 mL of crushed ice in order to decrease the sample temperature to below 2°C in less than 10 seconds. Cells were immediately centrifuged (4000 RPM at 0°C for 2.5 min.), the supernatant discarded, and the pellet frozen in liquid nitrogen and it was stored at −80°C until total RNA extraction. Total RNA was extracted using the FastRNA Pro RED kit (QiBiogene, Carlsbad, USA) according to manufacturer’s instructions after partially thawing the samples on ice. RNA sample integrity and quality was determined prior to hybridization with an Agilent 2100 Bioanalyzer and RNA 6000 Nano LabChip kit according to the manufacturer’s instruction (Agilent, Santa Clara, CA).

**Probe Preparation and Hybridization to DNA Microarrays.** Messenger RNA (mRNA) extraction, cDNA synthesis, labeling, and array hybridization to Affymetrix Yeast Genome Y2.0 arrays were performed according to the manufacturer’s recommendations (Affymetrix GeneChip® Expression Analysis Technical Manual, 2005–2006 Rev. 2.0). Washing and staining of arrays were performed using the GeneChip Fluidics Station 450 and scanning with the Affymetrix GeneArray Scanner (Affymetrix, Santa Clara, CA).

**Microarray Gene Transcription Analysis.** Affymetrix Microarray Suite v5.0 was used to generate CEL files of the scanned DNA microarrays. These CEL files were then processed using the statistical language and environment R v5.3 (R Development Core Team, 2007, www.r-project.org), supplemented with Bioconductor v2.3 (Bioconductor Development Core Team, 2008, www.bioconductor.org) packages Biobase, affy, germa, and limma [35], [36]. The probe intensities were normalized for background using the robust multivariate average (RMA) method only using perfect match (PM) probes after the raw image file of the DNA microarray was visually inspected for acceptable quality. Normalization was performed using the loess method and gene expression values were calculated from PM probes with the median polish summary. Statistical analysis was applied to determine differentially expressed genes using the limma statistical package. Moderated t-tests between the sets of experiments were used for pair-wise comparisons. Empirical Bayesian statistics were used to moderate the standard errors within each gene and Benjamini-Hochberg’s method was used to adjust for multi-testing. A cut-off value of adjusted p<0.03 was used for statistical significance. Furthermore, principal component analysis (PCA) was performed in order to elucidate the relative
adjusted the denominator, as described elsewhere [35]. The cut-off value of the moderated statistic from linear model theory but with the posterior variance substituted for the sample variance in the denominator, as described elsewhere [33]. The cut-off value of adjusted $p<0.1$ was used for statistical significance.

All microarray data is MIAME compliant and the raw data has been deposited in ArrayExpress (http://www.ebi.ac.uk/microarray-as/ae/).

Supporting Information

Figure S1 Inhibition of the succinate dehydrogenase complex with malonate supplementation in shake flask cultures was evaluated. The reference and $A_{sdh3}$ strain, previously described [17], were cultured in minimal media supplemented with 10 g L$^{-1}$ glucose and no succinate accumulation was detected (Panel a). The reference strain was cultured with 0.1, 1.0, 5.0, 10.0, and 50.0 mM malonate supplementation. Under no supplementation conditions succinate accumulation was observed (Panel b). In order to confirm that the concentration of malonate in the culture was effectively inhibiting succinate dehydrogenase activity, residual ethanol in the culture broth was monitored. Succinate dehydrogenase activity, as previously described, catalyzes the conversion of succinate to fumarate with net production of protonated ubiquinone. Ethanol is a carbon source readily catabolized by $S$. cerevisiae using respiro-fermentative pathways and requiring succinate dehydrogenase activity. Panel c shows the residual glucose concentration in the culture broth at 0, 17, 22, and 37h post-inoculation for no supplementation of malonate (reference) and then 0.1, 1.0, 5.0, 10.0, and 50.0 mM malonate supplementation. These growth profiles were generated using the reference strain (CEN.PK113-7D). As expected, full catabolism of glucose was observed at all malonate concentrations with the exception of 50.0 mM, thereby considered an upper limit. Similarly, in panel d, is the ethanol concentration in the culture broth for the same malonate concentrations and sample times. At 37 h, as expected, the reference strain had consumed nearly all ethanol produced during the glucose consumption phase. Malonate concentrations of 1.0, 5.0, and 10.0 mM malonate resulted in significant ethanol respiration inhibition compared to no supplementation and 0.1 mM malonate, confirming that respiro-fermentative catabolism was inhibited. Under no circumstances was succinate accumulation observed. Furthermore, the $A_{sdh3}$ strain was supplemented with 50.0 mM malonate to ensure no unexpected interaction between the genetic modification and malonate supplementation (panel e).

Figure S2 A total of 1964 genes were submitted to the Saccharomyces Genome Database tool, Pathway Expression Viewer. The resulting Pathway Expression map shows the relative log-fold change of all $S$. cerevisiae metabolic reactions (Evolved 8D vs. Reference). Three key results are highlighted from the transcriptome. First, isocitrate lyase ($ICL1$) was amongst the few genes not up-regulated in the Evolved 8D strain, thereby becoming a 2nd round metabolic engineering target. Second, alanine-glyoxylate aminotransferase ($AGXT$) was 4.3 log-fold higher in the Evolved 8D strain, confirming the predicted model-guided strategy of up-regulated glycine formation from glyoxylate pools. Third, threonine aldolase ($GLY1$) was 1.6 log-fold higher in the Evolved 8D strain. The genome-scale model reconstruction used for predictions annotated Gly1p as catalyzing the reversible conversion of threonine to glycine. This reaction has since been shown to be irreversible, converting threonine to glycine, consuming equimolar acetaldheyde. The transcriptome data suggests that the Evolved 8D strain demonstrated de-coupling of succinate and biomass production because alternative reactions (e.g., Gly1p) were supplying glycine pools.

Figure S3 Panel a briefly describes the mutant construction of 20G, $A_{sdh3}A_{ser3}A_{thr1}$, from the reference strain and initially supplemented with 100 mg L$^{-1}$ threonine and 500 mg L$^{-1}$ glycine to satisfy the resulting auxotrophies. All growth challenges were evaluated in shake flasks supplemented with minimal medium, 300 mg L$^{-1}$ uracil, 10 g L$^{-1}$ glucose, and either threonine and/or glycine added, as indicated. The mutant 20G was not capable of sustaining growth in the absence of threonine, and therefore a working cell bank was prepared. Panel b describes the shake flask experiments and progression followed to evaluate the strain’s ability to be evolved from threonine supplementation to glycine supplementation. When 20G culture was inoculated from threonine supplemented medium to glycine only supplemented medium, no growth was observed up to 14d post-inoculation (2 samples per day measuring OD$_{600}$). On day 14, a shake flask culture of 20G only supplemented with glycine, was then supplemented with 100 mg L$^{-1}$ threonine, and growth was immediately restored. It was therefore concluded that the mutant 20G was incapable of catalyzing glycine to threonine to satisfy threonine cellular demands, given that threonine synthesis was interrupted with the deletion of $thr1$. This experimental conclusion further supports that Gly1p encoding threonine aldolase, originally believed to catalyze the conversion of glycine to threonine, catalyzes the reverse reaction and thus cannot meet threonine cellular demands from glycine pools.

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Author Contributions

Conceived and designed the experiments: JMO DC KRP LO JN. Performed the experiments: JMO DC SGP. Analyzed the data: JMO DC JN. Wrote the paper: JMO JN.

References


