

# Quantitative Analysis of Glycerol Accumulation, Glycolysis and Growth under Hyper Osmotic Stress

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## Abstract

We provide an integrated dynamic view on a eukaryotic osmolyte system, linking signaling with regulation of gene expression, metabolic control and growth. Adaptation to osmotic changes enables cells to adjust cellular activity and turgor pressure to an altered environment. The yeast *Saccharomyces cerevisiae* adapts to hyperosmotic stress by activating the HOG signaling cascade, which controls glycerol accumulation. The Hog1 kinase stimulates transcription of genes encoding enzymes required for glycerol production (Gpd1, Gpp2) and glycerol import (Stl1) and activates a regulatory enzyme in glycolysis (Pfk26/27). In addition, glycerol outflow is prevented by closure of the Fps1 glycerol facilitator. In order to better understand the contributions to glycerol accumulation of these different mechanisms and how redox and energy metabolism as well as biomass production are maintained under such conditions we collected an extensive dataset. Over a period of 180 min after hyperosmotic shock we monitored in wild type and different mutant cells the concentrations of key metabolites and proteins relevant for osmoadaptation. The dataset was used to parameterize an ODE model that reproduces the generated data very well. A detailed computational analysis using time-dependent response coefficients showed that Pfk26/27 contributes to rerouting glycolytic flux towards lower glycolysis. The transient growth arrest following hyperosmotic shock further adds to redirecting almost all glycolytic flux from biomass towards glycerol production. Osmoadaptation is robust to loss of individual adaptation pathways because of the existence and upregulation of alternative routes of glycerol accumulation. For instance, the Stl1 glycerol importer contributes to glycerol accumulation in a mutant with diminished glycerol production capacity. In addition, our observations suggest a role for trehalose accumulation in osmoadaptation and that Hog1 probably directly contributes to the regulation of the Fps1 glycerol facilitator. Taken together, we elucidated how different metabolic adaptation mechanisms cooperate and provide hypotheses for further experimental studies.

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## Introduction

Upon increase in external osmolarity, cells first shrink and subsequently recover volume by accumulating compatible solutes [1]. Different processes contribute to adaptation, establishing negative feedback loops for the regulation of osmotic pressure, cell volume, and turgor [2–5]. The yeast *Saccharomyces cerevisiae* employs glycerol as compatible solute for osmo-regulation. Upon hyperosmotic shock the High Osmolarity Glycerol (HOG) pathway is activated, resulting in phosphorylation of the stress-activated protein (SAP) kinase Hog1. Phosphorylated Hog1 stimulates expression of genes encoding enzymes involved in glycerol

production and uptake. Hyperosmotic stress also leads to rapid closure of the glycerol facilitator Fps1 preventing glycerol outflow. The contributions to glycerol accumulation of the different processes and regulatory mechanisms – including central metabolism – have not been systematically studied in a quantitative and time-resolved manner.

In the absence of hyperosmotic stress, glycerol production is required for maintaining the redox balance [1,6] and excess glycerol leaks out freely through the glycerol facilitator Fps1 [7,8]. The glycerol production pathway starts with the reduction of the glycolytic intermediate di-hydroxyl-acetone phosphate (DHAP) to glycerol-3-phosphate (G3P) catalyzed by the NAD<sup>+</sup>-dependent



mentary Information (SI) and Fig. S10). We included carbon flux to biomass, the influence of Hog1 on glycolytic flux via Pfk26/27, and Stl1-mediated glycerol uptake. This model comprises the following modules:

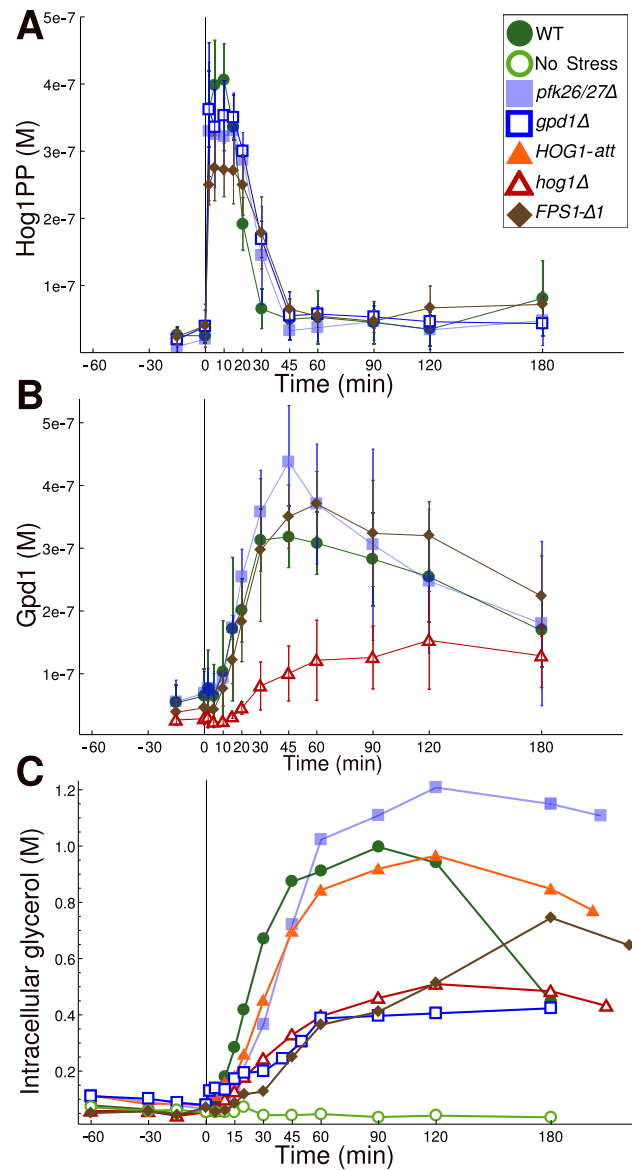
- **Biophysical changes:** changes in cell volume (basal solid volume,  $V_b$ , and osmotically active volume,  $V_{os}$  [23,24]), cell surface area, osmotic pressure, and turgor pressure (where changes depend on  $V_{os}$  [2] and external osmolarity).
- **Glycolysis module:** metabolic reactions from glucose uptake via phosphorylated intermediates to glycerol, trehalose, ethanol, and acetate. The module is based on models [2,25] where reactions have been lumped to reduce the number of parameters (see Text S1 for details). The  $V_{max}$  of the Pfk1 reaction is modulated by F26DP, which depends on the Hog1-regulated activity of Pfk26/27.
- **Transport module:** glucose, trehalose, ethanol, acetate transport, glycerol exchange via Fps1, and irreversible glycerol uptake via Stl1. We corrected rate laws of transport reactions for cell density increase in time-course experiments.
- **Biomass module:** biomass production is necessary for describing the interplay between osmotic regulation and glycolysis, but also to maintain correct carbon balance. Biomass is measured as cell number or cell density. Upon osmoshock, Hog1 arrests cell cycle progression [26,27] and hence proliferation. Hog1-activity affects metabolism by altering expression of metabolic genes [20,28–30]. A Hog1-dependent drop in growth rate is included in the model ( $v_{14}$ ).
- **Adaptation module:** Hog1 phosphorylation, *GPD1*, and *STL1* mRNA, translation to Gpd1 and Stl1, activation of Pfk26/27, and closure of Fps1. Hog1 activity is correlated with volume changes and turgor [24]. The Hog1 signaling system is described with only two reactions.

In order to ensure that the available data support the model, which covers different major processes, metabolism is highly condensed for this specific study, compared to metabolic reconstructions suitable for steady state analyses. For example, we refrained from including redox balance dynamics since they are affected by cellular changes beyond the scope of this model, such as detailed biosynthetic pathways (see Text S1). We do not assume a steady state for the entire model because we consider external metabolites and cell density. Although most intracellular concentrations can be considered constant before osmotic stress, the increasing external concentrations of ethanol, acetate and glycerol and the decreasing external glucose level together with the increase in OD slightly affect model dynamics. Therefore, the model is not in steady state in a strict sense.

Models for the different mutant strains are generated by implementing known effects of the respective genetic perturbation (see Text S1). Besides these modifications, all models use the same parameter set.

## Data set

To estimate model parameters and determine time scales, we measured intracellular and extracellular concentrations of glycerol, glucose, ethanol, acetate, and trehalose over 240 min for wild type and mutants lacking particular adaptation mechanisms (*hog1Δ*, *gpd1Δ*, *pfk26Δ*, *pfk27Δ*, *pfk26/27Δ*) or with altered regulation nodes (*HOG1-att*, *FPS1-Δ1*) (Fig. 2, SI). While external glucose is consumed, glycerol, ethanol, and acetate as well OD<sub>600</sub> and cell numbers (SI, Fig. S7) increased over time. Internal glycerol (Fig. 2C) accumulates transiently, while trehalose shows two peaks at 45 min and 180 min (Fig. S7G). We also measured *GPD1*



**Figure 2.** Time courses of (A) phosphorylated Hog1, (B) Gpd1, and (C) intracellular glycerol following hyperosmotic stress of 0.4 M NaCl at time point 0. The full dataset is provided in Datasets S1, S2, S3, S4, S5, S6, S7.

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mRNA concentration, Gpd1 concentration, and Hog1 phosphorylation (Fig. 2A–B, Fig. S13B). The following characteristics were noted.

- In the *gpd1Δ* mutant, the extracellular glycerol decreases, while the *FPS1-Δ1* strain excretes large amounts of glycerol (Fig. S8).
- Intracellular glycerol (Fig. 2C) builds up with a delay in the *FPS1-Δ1* and *hog1Δ* mutants. In the *pfk26/27Δ* and *Hog1-att* [21] strains, intracellular glycerol remains high over the entire period.
- The *pfk26/27Δ* mutant displays slower glucose consumption (Fig. S7B).
- Trehalose dynamics at 180 min indicate that it may accumulate further in wild type; *pfk26/27Δ* and *HOG1-att* strains do not display a late increase of trehalose (Fig. S7G).

- Hog1 phosphorylation (Fig. 2A) is slightly prolonged in cells lacking Gpd1 or Pfk26/27 or expressing constitutively open Fps1.
- The lowest Gpd1 levels are observed for the *hog1Δ* mutant (Fig. 2B).

### Simulation of osmoadaptation

The model was fitted to experimental data (Figs. S12, S13, S14). In general, model fitting can follow two different paradigms: (i) either fit the model to part of the data and use the rest of the data to test model predictions or (ii) use all data for fitting to obtain the model best describing the observations. Since our quantitative time-resolved data of various compounds for wild type and different mutants comprehensively elucidates the potential network dynamics, we decided to eventually use all data from all yeast strains and conditions to parameterize the final model presented here. To reproduce the Gpd1-mediated increase of glycerol production, dimerization of Gpd1, as reported for human Gpd [21,31,32], was implemented. We further implemented a partial Hog1-independent increase of *GPD1* mRNA to reproduce *hog1Δ* data, which is in accordance with experimental observations [21,22]. To fit *gpd1Δ* mutant data, we incorporated a small osmo-dependent increase in *GPD2* transcription. Including regulation of biomass production [30] significantly improved model behavior (SI): A rapid drop in cell growth, dependent on stress and cell volume, is necessary to provide sufficient carbon for glycerol production and reproduce experimental data on cell density. To reproduce experimental data for intracellular glycerol in *hog1Δ* and *HOG1-att* strains, we included a negative regulation of open Fps1 by active Hog1.

### Model analysis and predictions

To characterize the contribution of different processes to osmoadaptation (Fig. 3), we plotted the absolute fluxes to and from glycerol for wild type and *gpd1Δ* over time (Fig. 3A and B) and relative contributions of fluxes in ternary plots (Fig. 3C). We distinguish Fps1-reliant, Gpd1-reliant, and other contributions ('Others') to net glycerol flux in each strain. For wild type, immediate adaptation is influenced mainly by changes in volume (part of 'Others') and closure of Fps1. Within approximately 15 min after stress, glycerol production becomes the predominant contribution. After 30 min, Fps1 reopens and glycerol levels decrease. Open Fps1 in *FPS1-Δ1* results in constitutive glycerol efflux. Although the model underestimates the glycerol production for this mutant, it clearly shows that the distribution of fluxes remains at a state of sustained glycerol production. Sustained glycerol production occurs also in the *hog1Δ* mutant, but due to reduced Hog1-dependent *GPD1* transcription its level remains low until 60 min after stress. Subsequently, the system moves to a state similar to that observed for the *FPS1-Δ1* strain.

Although the relative composition of fluxes approaches pre-stress values in wild-type, we do not observe perfect adaptation [33] (Fig. 4). For a more comprehensive analysis, the complete state of cells has to be considered, including changes in reactions or pathways indirectly affecting glycerol concentrations, e.g. changes in biomass production. To do so, we employed scaled time-dependent response coefficients (RCs),  $R_q^s(t) = \partial \ln s(t) / \partial \ln q$  for a compound concentration  $s(t)$  and a parameter  $q$  [34]. RCs express the relative change in  $s(t)$  given a small change in  $q$ , serving as quantitative measure for the effect of a parameter perturbation on a time course taking all direct and indirect effects into account. Positive or negative values indicate that the time course increases or decreases upon parameter

increase, respectively. Scaled RCs (Fig. 4B–C) show that regulation of Pfk26/27 does not have a pronounced effect on intracellular glycerol (Fig. 4B), which is in accordance with experimental data for *pfk26/27Δ* (Fig. 2C). Additionally, other reactions downstream of Pfk1 are positively affected while glycolytic reactions upstream of Pfk1 (trehalose and biomass production) are reduced. This indicates that Pfk26/27 might be part of a rerouting mechanism of metabolic flux ensuring that the influx to each branch of glycolysis is adjusted to demands and that ATP-production downstream of pyruvate is maintained during osmoadaptation.

We use RCs to compare the role of Stl1 in different experiments (Fig. 4C): apparently the contribution of Stl1, though negligible in most strains until 90 min after stress, is significant in *gpd1Δ*. *In silico* predictions show an early decrease in intracellular glycerol accumulation in the *gpd1Δ stl1Δ* double mutant compared with *gpd1Δ* but a late decrease in the *stl1Δ* mutant compared with wild type (Fig. 4D). The different roles of Stl1 in wild type and *gpd1Δ* cells highlight context specificity in osmoadaptation.

The regulation of biomass production significantly contributes to glycerol accumulation (Fig. 5, Fig. S15B). Rerouting of glycolytic flux can be assumed to result from Hog1-mediated cell-cycle arrest and glycolytic regulation. The cost of maintaining a certain cell volume by producing more glycerol is compensated by a decrease in growth rate. Changes in doubling times before and after stress are plotted in Fig. 5A. The observed decrease in growth rate is similar for wild type, *pfk26/27Δ*, and *HOG1-att* strains. This indicates that the main contribution of Hog1 activity to osmoadaptation is not the transcriptional activation of *GPD1*, which is absent in the *HOG1-att* strain. Instead, control of cytosolic or membrane-bound targets constitutes the main contributions of Hog1. A strong growth rate drop is observed in *FPS1-Δ1* and *hog1Δ*, while this drop is relatively low in the *gpd1Δ* mutant. This supports the idea that a Hog1-effect on Fps1 is mainly responsible for the prominent role of Hog1 in osmoadaptation. Fig. 5B shows model simulations of the relative carbon fluxes from glycolysis to glycerol or biomass production, respectively, for wild type and different mutants before 20 min and 90 min after stress induction, indicating a trade-off of cellular adaptation *versus* growth.

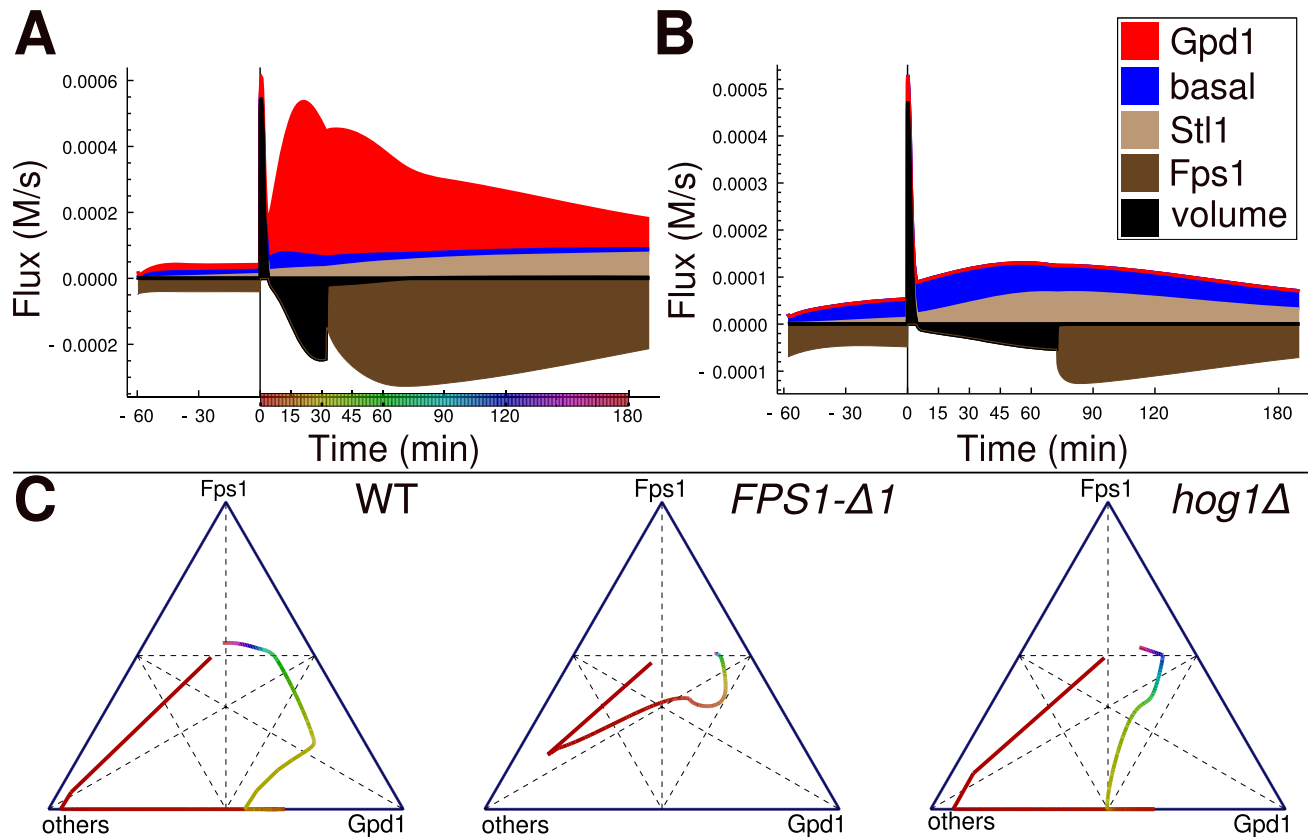
The model is fitted to data covering a period until 180 min after stimulation. However, cellular processes beyond the scope of the model may interfere with the adaptation process at later stages, hence model results and hypotheses derived from those may be less reliable beyond 120 min.

### Discussion

For the first time we present a data-driven analysis of the dynamic regulation of a eukaryotic osmolyte system mediating adaptation to hyperosmotic stress that integrates different regulatory layers. Our mathematical model, supported by a detailed dedicated dataset, provides novel insight into the quantitative contributions of regulatory processes underlying glycerol accumulation during the yeast osmostress response.

The main conclusions are:

- The Hog1 SAPK mediates regulation of glycolysis via indirect activation of the enzyme Pfk1 and increased expression of glycerol-producing enzymes.
- Pfk1 activation serves stabilization of glycolytic flux as much as increased glycerol accumulation.
- Glycerol accumulation occurs at the expense of biomass production.



**Figure 3. Contribution of glycerol accumulation mechanisms in different strains.** (A, B) Absolute fluxes towards glycerol as well as relative contributions of specific mechanisms differ between wild type WT (A) and *gpd1Δ* (B). (C) Changes in relative contributions of Fps1, Gpd1, and other effects (basal glycerol production, uptake through Stl1, effects of volume change) over time are depicted for WT, *fps1-Δ1* and *hog1Δ*. Colors in (C) indicate time as shown on the x-axis in (A).  
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- (iv) Glycerol accumulation is achieved by different fluxes comprising basal (Gpd2-dependent) and Hog1-regulated (Gpd1-dependent) production, glycerol exchange over the plasma membrane as well as concentration increase through volume loss; these fluxes contribute to varying extent during different periods of adaptation.

The parameter set was fitted to reproduce data from seven different strains and experimental observations on 14 of 29 model variables, providing the model with an unprecedented coverage [35]. This coverage was achieved employing coordinated model reduction and experimentation with the intention to maintain all regulatory nodes while lumping metabolites and proteins for which no experimental data was gathered.

Deleting individual nodes of the glycerol accumulation network (Figs. 1, S10) results in slightly prolonged HOG pathway activation. For *FPS1-Δ1*, this prolongation is less pronounced than in previous data [2], which is probably due to different copy numbers of the expression construct. In contrast to Dihazi et al. [17], we found that only the double deletion mutant, *pfk26Δpfk27Δ* shows prolonged Hog1 phosphorylation. The mutation causes a transient delay of glycerol accumulation (Fig. 2C).

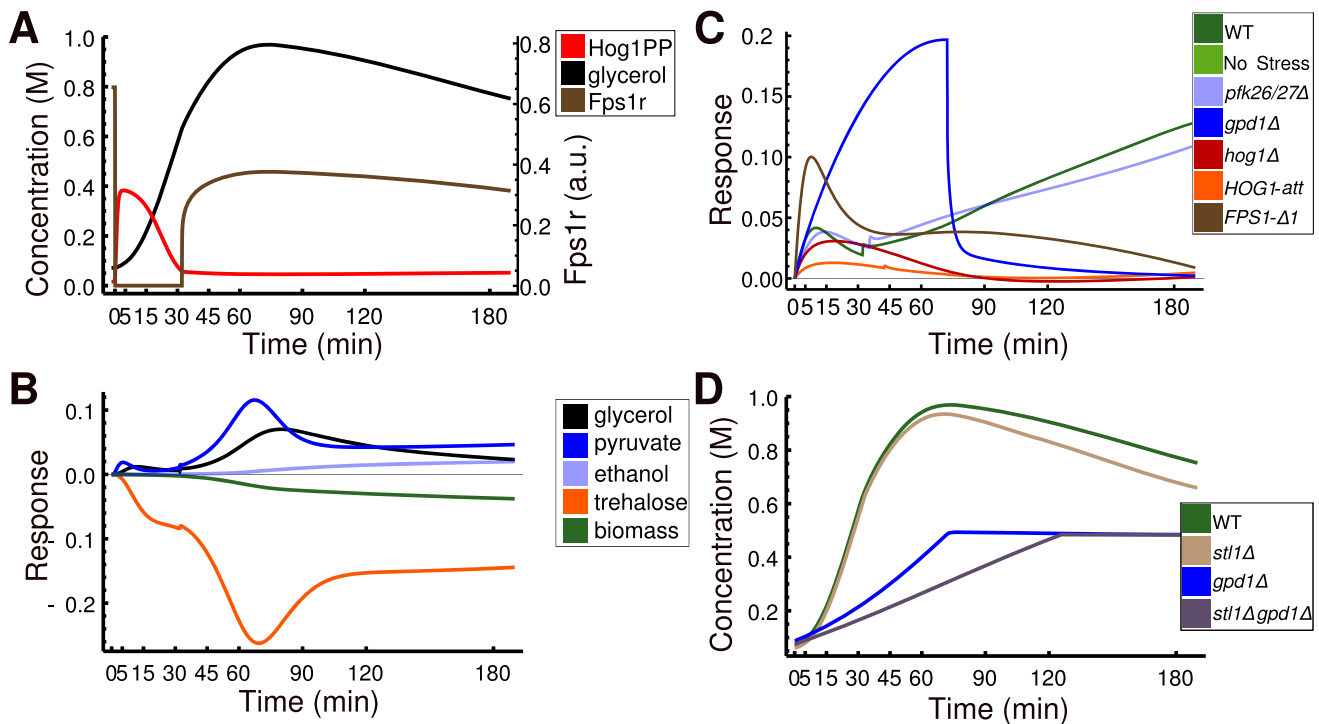
We hypothesize that, like human Gpd1 [32], also yeast Gpd1 dimerizes since this assumption results in significantly better fits to experimental data.

It has been shown that Hog1 interacts with and controls Fps1 under certain conditions [36–38], although the exact mechanism

of Hog1-dependent control of Fps1 remains elusive. Experimental observations (e.g. [8]), as well as the simulations shown here and elsewhere [5], are consistent with a Hog1-dependent down-regulation of glycerol flux through Fps1 under osmstress. In *hog1Δ* cells the glycerol efflux is elevated while membrane-attached Hog1 results in a very strict regulation.

In the *gpd1Δ* mutant the glycerol production rate is diminished. Our experimental data show that in this strain the extracellular glycerol concentration decreases over time, indicating glycerol uptake through the Stl1 glycerol-proton symporter. We determined the role of Stl1 in glycerol accumulation using time varying response coefficient analysis [34]. Our simulations indicate that the *stl1Δgpd1Δ* mutant adapts even worse than the *gpd1Δ* mutant and does not show a decrease in the extracellular glycerol concentration. This indicates that osmstress response can overrule glucose repression of Stl1, which is supported by experimental data from Ferreira et al [19] showing strongly stimulated *STL1* gene expression under osmstress in mutants unable to produce glycerol as well as growth phenotypes of such mutants lacking in addition Stl1.

The presented model underestimates the production of glycerol for *FPS1-Δ1*. This may have two reasons. First, *FPS1-Δ1* cells adapt to the imposed genetic perturbation prior to stress, which can, for example, lead to an up-regulation of glycerol production. Second, the proposed interaction of the modified protein with Hog1 or other potential regulators [38] may be perturbed. Thus, the model indicates that *FPS1-Δ1* has global effects beyond the



**Figure 4. Model analysis with time-dependent response coefficients.** **A:** Model simulation for phosphorylated Hog1, intracellular glycerol, and abundance of open Fps1. **B:** Effect of small changes in Pfk26/27 activation on different model variables as expressed by normalized response coefficient indicates that Pfk26/27 contributes to a rerouting of flux towards lower glycolysis. **C:** Response of intracellular glycerol concentration to perturbations in *Stl1* gene expression as measured by normalized response coefficients in different strains indicates a specific time- and context-dependent role of *Stl1* in osmoadaptation. **D:** Simulation of genetic perturbations of *Stl1* results in time courses as expected from C: in wild type, deletion of *STL1* affects intracellular glycerol levels only at later time points. In *gpd1Δ* background, the effect of additional deletion of *STL1* is early and transient.

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expected de-regulation of glycerol efflux that need to be investigated further.

In adapted yeast cells the intracellular glycerol level diminishes. We observed an increase in intracellular trehalose concentrations coinciding with the decline of intracellular glycerol (Fig. S7G). In the *FPS1-Δ1* strain, which is unable to accumulate intracellular glycerol, the intracellular trehalose concentration is permanently high. Although the combination of high trehalose and glycerol concentrations has a synergistic osmoprotective effect [39], the observed trehalose concentrations are too small to significantly contribute to intracellular osmotic pressure. Hence, trehalose may have more specific roles in osmoadaptation by protecting proteins and membranes [40,41].

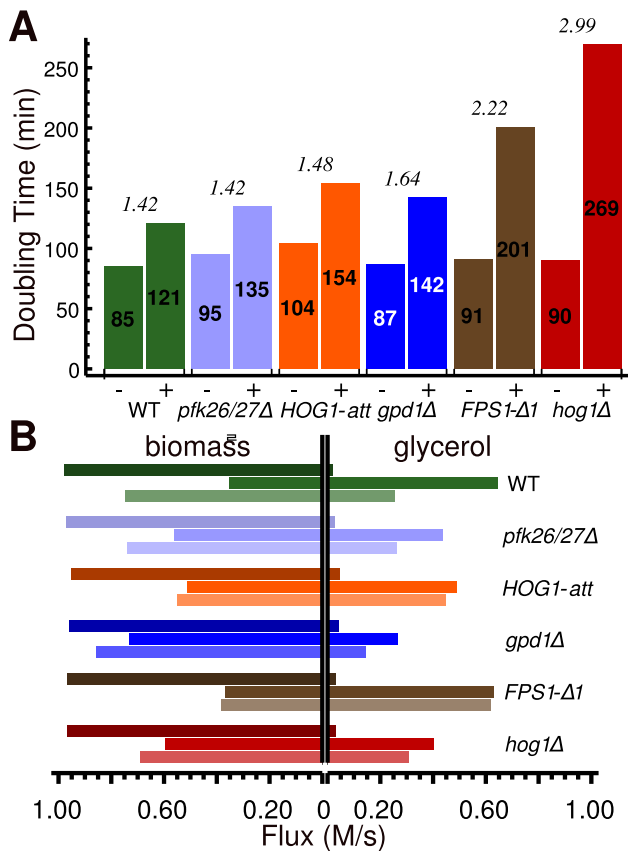
We do not observe perfect adaptation in the sense defined previously [33]. Although cell volume returns to its original values, other variables remain perturbed in adapted cells at constant high external osmolarity, such as *Gpd1*, trehalose or glycerol concentration, and especially growth rate.

Glycolysis serves three major fluxes resulting in the production of ethanol plus acetate, biomass, and glycerol. Although active Hog1 leads to cell cycle arrest [26,27], potentially decreasing carbon flux towards biomass, a role of glycolysis in the osmotic stress response has not been considered in detail. Data presented by Dihazi et al. [17] as well as recent gene expression data [42] indicate that glycolytic flux may play a role in osmoadaptation. Our preliminary theoretical work [22] indicated that *Pfk26/Pfk27* participate in maintaining the flux towards pyruvate upon adaptation to hyper-osmotic conditions. Glucose consumption and ethanol production rates are similar in all strains studied here,

regardless which node of the glycerol accumulation network was removed. The *pfk26/27Δ* strain is the only exception: the slightly lower ethanol production and glucose consumption observed in this mutant indicate that *Pfk26/27* influence glycolysis rather than osmotic adaptation. We conclude that under hyper-osmotic stress part of the carbon flux, which is normally directed towards biomass production, is used for increasing glycerol production. One of the elements in this metabolic prioritization mechanism is *Pfk26/27*, although it does not seem to be the only factor. The ratio between the production of glycerol and pyruvate (resulting in the formation of ethanol and acetate) is crucial for a proper energy and redox balance. In all strains, the flux towards pyruvate remains unaffected by osmoadaptation while flux towards biomass decreases. Apparently, energy production and redox balance are maintained while growth is temporarily stopped for faster adaptation. This pattern is likely to hold for a wide variety of stresses and adaptation mechanisms.

The relative contribution of each individual glycerol accumulation mechanism depends on the environmental conditions and the physiological state of the cell (Fig. 3A). In their natural environment, cells have to cope with combinations of stresses demanding for dynamic modulation of response mechanisms. The different mechanisms contributing to osmoadaptation might make the process robust but also allow choosing the ‘cheapest’ adaptation strategy under different conditions.

Response coefficient analysis is an excellent tool to elucidate this kind of interdependences. We found that introducing an individual deletion into the glycerol accumulation network is compensated. Our experimental data show that the adaptation takes longer for



**Figure 5. Effect of salt stress on growth rate.** **A:** *In vivo* doubling times (–: before, +: after addition of 0.4 M NaCl) strongly differ between strains. **B:** Model simulations of the flux towards biomass production (left) and glycerol production (right) in the different strains at 0, 20, and 90 minutes after osmotic upshift to 0.4 M NaCl indicate a link between insufficient glycerol accumulation and a prolonged decrease in growth rate.  
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single knock-out strains, compared to wild type. The activation of the remaining glycerol accumulation mechanisms is prolonged (active component of the compensation) and the remaining mechanisms become more important (passive compensation component).

**Table 1. Yeast strains used in this study.**

Strain name	Strain genotype	Origin
W303-1a	MATa leu23/112 ura31 trp11 his311/15 ade21 can1100 GAL SUC2	Thomas and Rothstein (1989) [45]
YMR84	W303-1A with <i>gpd1Δ::URA3</i>	Martijn Rep (Amsterdam)*
YSH1583	W303-1A with <i>pfk26Δ::KanMX</i>	This study
YSH1585	W303-1A with <i>pfk27Δ::KanMX</i>	This study
YSH1586	W303-1A with <i>pfk26Δ::KanMX pfk27Δ::KanMX</i>	This study
YMT101	W303-1A with <i>fsp1Δ::LEU2 Ylp-URA3-fps1-Δ1</i>	Tamas et al., (1999) [8]
YSH2293	W303-1A with <i>stl1Δ::KanMX</i>	This study
YSH444	W303-1A with <i>hog1Δ::TRP1</i>	Albertyn et al., (1994) [12]
	W303-1A with <i>hog1-att</i>	This study

\*Strain YMR84 was kindly provided by Martijn Rep (Amsterdam) and contains a replacement of the GPD1 upstream region (–883 to +91) by the URA3 gene. The strains was generated using a PCR approach and does not express the GPD1 gene product.  
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Minimalistic models of biological processes, such as the osmoadaptation model presented by Mettetal et al. [3] are of great value for understanding general principles underlying a given process. With a sufficient degree of generalization, conclusions drawn from minimalistic models can be applicable for characterizing other cellular events. This approach neglects the individual mechanisms contributing to a cellular function. On the other extreme, the approach of hierarchical control analysis [43] measures the effects of changes on a cellular process in an enormous amount of detail. This approach is crucial for understanding the role of individual components and events. Our work attempts to bridge between the two paradigms, which can be viewed as an intermediate stage of different magnifications of the same picture.

The principles of osmoregulation are conserved from yeast to mammalian cells. The mammalian stress-activated protein kinase p38 is a homolog of Hog1. Like Hog1 it plays a critical role in mounting the adaptive response to stress by controlling metabolism, gene expression and cell cycle progression [44]. Hence, the approaches employed in the present work as well as the conclusions drawn may have consequences for studies on mammalian cells as well.

By integration of experimental data for different strains into a medium-size model and reliably estimating its parameters, we were able to achieve a better understanding of the contribution of individual players to a cellular response – osmoadaptation – in a quantitative and time-resolved manner. Moreover, our analysis has revealed the trade-off between growth control and glycerol accumulation in the adaptation process.

## Materials and Methods

### Experimental methods

**Yeast strains and culturing.** Strains used in this work originate from W303-1A (MATa leu2-3/112 ura3-1 trp1-1 his3-11/15 ade2-1 can1-100 GAL SUC2 mal0) [45]. The strains include: wild type, *gpd1Δ*, *pfk26Δpfk27Δ*, *fps1-Δ1* (kindly provided by Markus Tamás, University of Gothenburg), *stl1Δ*, *hog1Δ*, *HOG1-att* (kindly provided by Jeremy Thorner, University of California at Berkeley). A complete list of strains used in this study is presented in Table 1. Yeast cultures were grown until mid-exponential phase ( $OD_{600} = 0.7-1.0$ ) in YPD medium (Yeast Peptone D-glucose; 1% yeast extract (Bacto), 2% peptone (Bacto), 2% glucose), then NaCl was added from a stock solution of 5 M in water to a final concentration of 0.4 M at  $t = 0$  min.

**Western blot analysis.** Samples of 1 ml were collected at the indicated time points, sedimented and frozen in liquid nitrogen after removing the supernatant. Proteins were extracted by boiling for 10 min in extraction buffer (100 mM Tris-HCl pH 6.8, 20% glycerol, 200 mM DTT, 4% SDS, 10 mM NaF, 0.1 mM Na<sub>3</sub>V04 (sodium orthovanadate), protease inhibitor (Complete EDTA-free Protease Inhibitor Cocktail tablets, Roche), and 20 mM mercaptoethanol). The extracts were clarified by centrifugation (13 000 rpm in 4°C for 10 min). For each sample 40 µg of protein was separated by electrophoresis on a 10% polyacrylamide gel (SDS-PAGE) and transferred (semi-dry) to a nitrocellulose membrane (Hybond-ECL, Amersham). Membranes were blocked with Odyssey Blocking Buffer (Li-Cor Biosciences) and incubated sequentially with primary antibodies: first primary antibody - phospho-p38 MAPK (Thr180/Tyr182) monoclonal rabbit antibody (Cell Signalling), 1:1000 in Odyssey Blocking Buffer with TBST (1:1000), overnight at 4°C; second primary antibody -  $\gamma$ C20 total Hog1 polyclonal goat antibody (Santa Cruz Biotechnology Inc.), 1:2000 in Odyssey Blocking Buffer with TBST (1:1000), 1 h at room temperature, third primary antibody - rabbit polyclonal antisera Gpd1-A (Innovagen), 1:2000 in Odyssey Blocking Buffer with TBST (1:1000), 1 h at room temperature, and simultaneously with secondary antibodies: donkey anti-goat IR Dye 680, 1:12 500 and donkey anti-rabbit IR Dye 800CW 1:12500 (Li-Cor Biosciences), in Odyssey Blocking Buffer with TBST (1:1000), for 45 min at room temperature.

The membranes were scanned using Odyssey Infrared Imaging System (Li-Cor Biosciences) and quantified using Multi Gauge 3.0 (FujiFilm) software.

**Metabolite measurements.** Samples of 1 ml were collected at the indicated time points. Three types of samples were collected for each time point: intracellular - centrifuged cell pellet without supernatant, extracellular - pure supernatant, removed from intracellular samples, total - 1 ml cells in medium, and frozen in liquid nitrogen. Extracellular samples did not require further processing; total extracts were boiled for 10 min and cleared by centrifugation, cell pellets were extracted with sterile water by boiling for 10 min and cleared by centrifugation. The concentrations of glucose, trehalose, glycerol, acetate, succinate, pyruvate and ethanol were measured by high performance liquid chromatography (DIONEX) with an Aminex HPX-87H ion exchange column (Bio-Rad, Hercules, USA). An isocratic condition was performed with 5 mM H<sub>2</sub>SO<sub>4</sub> as mobile phase at flow rate of 0.6 ml/min and oven temperature of 65°C. Glucose, trehalose, glycerol and ethanol were quantified by a refraction index detector (Waters 410 Differential Refractometer Millipore, CA, USA) and acetate, succinate and pyruvate by ultraviolet-1 visible light absorbance detector (Waters 486 Tunable Absorbance Detector set at 210 nm, Millipore, CA, USA).

**Data processing and additional data.** Experimental data was processed to account for cell density increase and the reliability of HPLC measurements was assessed by comparison with enzyme assay quantification. For a full description of data processing and additional data, see Text S1.

## Mathematical modeling

System dynamics were described by ordinary differential equations (ODEs). A complete list of model equations and parameter estimation procedures is provided in SI. Time-dependent response coefficients have been calculated as described in [34]. Temporal simulations were performed with Mathematica7 (Wolfram Research. Mathematica edition: Version 7.0, 2008). Parameter estimation was done with PottersWheel [46] and SBML-PET [47].

## Supporting Information

**Dataset S1** Raw and processed metabolite measurements for different strains, stress 0.4 M NaCl added at  $t=0$ . (XLS)

**Dataset S2** Overview of experimental data as used in fitting. (XLS)

**Dataset S3** Comparison of intracellular trehalose measurements by HPLC and enzyme assay, stress 0.4 M NaCl added at  $t=0$ . (XLS)

**Dataset S4** Western Blot data for different strains and stress strengths. (XLS)

**Dataset S5** Intracellular glycerol quantification by enzyme assay in different strains for different stresses added at  $t=0$ . (XLS)

**Dataset S6** Intracellular trehalose quantification by enzyme assay in different strains, stress 0.4 M NaCl added at  $t=0$ . (XLS)

**Dataset S7** Northern Blot results for CTT1, GRE2, STL1, 18S in different strains. (XLS)

**Figure S1** Cell density and optical density in stressed (0.4 M NaCl added at  $t=0$ ) and control experiments (A) and normalized to the values at  $t=0$  (B). Experimental setup as explained in main text. (PDF)

**Figure S2** Plot of OD versus cell density values as obtained from control and stressed time course experiments and the function for computing cell density from OD. (PDF)

**Figure S3** Measured (blue) and inferred (pink) intracellular glycerol for different experiments. Errors in the inferred values due to inconsistencies in measurements are visible in *FPS1-Δ1*, *hog1Δ* and WT1. (PDF)

**Figure S4** Comparison of glycerol time courses for WT1 and WT4. (A) intracellular glycerol, (B) extracellular glycerol. (PDF)

**Figure S5** Comparison of enzyme assay (blue) and HPLC (pink) intracellular glycerol quantifications. Underlying data is given in Supplemental Dataset S2. (PDF)

**Figure S6** Comparison of intracellular trehalose levels obtained with different methods. Enzyme assay data is in g D-glucose/l per µg protein/ml, HPLC data in mol/l is normalized to the enzyme assay value at  $t=30$ . Underlying data is given in Supplemental Dataset S3. (PDF)

**Figure S7** Experimental data as used for model fitting. Measured entities are indicated on y-axes. Shown are representative experiments for each strain. Stress of 0.4 M NaCl is added at  $t=0$ . A: intracellular glucose, B: extracellular glucose, C: Optical density, D: intracellular ethanol, E: extracellular ethanol, F: extracellular acetate, G: intracellular trehalose, H: extracellular trehalose, I: extracellular glycerol. (PDF)



**Figure S8** Experimental data for extracellular glycerol (A,B) and Gpd1 (C) following addition of 0.4 M NaCl at  $t=0$ . Close inspection reveals a decrease of extracellular glycerol in *gpd1Δ* strain. The decrease intracellular glycerol in *gpd1Δ* is on a similar timescale as the transcriptionally regulated increase of Gpd1 in other strains, indicating that a transcriptionally regulated mechanism is responsible for this decrease as well. (PDF)

**Figure S9** Effects of *HOG1-att* (Hog1 attached to the plasma membrane) compared to wild-type. Under unstressed conditions, inactive Hog1 (dark red stars) is localized throughout the cell, residual active Hog1 (light red stars) is localized in the nucleus. Possible Hog1 interaction partners are depicted: Fps1 (brown triangles), Pfk26/27 (blue spirals) and genes (black waves). In wild-type, osmoadaptation leads to active Hog1 translocating to the nucleus to stimulate transcription. In *HOG1-att*, transcriptional regulation is abolished and cytosolic Hog1-concentration is reduced while possible interactions with membrane-bound proteins are increased. (PDF)

**Figure S10** Model topology in SBGN syntax. The different modules are color coded (red: adaptation, yellow: biophysical, brown: transport, blue: glycolytic, green: growth). Measured entities are indicated by a green background. Perturbations to the model (as stress [NaCl] or different mutations [I: *hog1Δ*, II: *pfk26/27Δ*, III: *HOG1-att*, IV: *gpd1Δ*, V: *FPS1-Δ1*]). (PDF)

**Figure S11** Changes in cell density in a batch culture experiment. Cells covered by an ODE model highlighted in yellow. (PDF)

**Figure S12** Agreement between main model variables and experimental data. Concentrations of intracellular glycerol (dashed) and phosphorylated Hog1 in different strains (A: WT, B: *pfk26/27Δ*, C: *HOG1-att*, D: *FPS1-Δ1*, E: *gpd1Δ*, F: *hog1Δ*) following hyperosmotic stress of 0.4 M NaCl at  $t=0$ . (PDF)

**Figure S13** Experimental data and simulated model variables, stress 0.4 M NaCl added at  $t=0$ . A: phosphorylated Hog1, B: GPD1mRNA, C: Gpd1, D: cell volume, E: abundance of open Fps1, F: intracellular trehalose. Concentrations in A and B are scaled as described in text. (PDF)

**Figure S14** Simulation of model variables, stress 0.4 M NaCl added at  $t=0$ . A: extracellular glycerol, B: intracellular glycerol,

C: extracellular trehalose, D: extracellular glycerol, E: extracellular ethanol, F: extracellular acetate. (PDF)

**Figure S15** Model Simulations and scaled response coefficients for models of different strains. A,D,G: Model variables (solid line: phosphorylated Hog1, dashed line: intracellular glycerol, dotted: abundance of open Fps1) for wild-type, *gpd1Δ* and *hog1Δ*, respectively. B,E,H: scaled response coefficients of osmoshock dependent parameters on intracellular glycerol for wild-type, *gpd1Δ* and *hog1Δ*, respectively. C,F,I: scaled response coefficients of glycolytic parameters on intracellular pyruvate for wild-type, *gpd1Δ* and *hog1Δ*, respectively. (PDF)

**SBML Model S1** Annotated model of osmoadaptation in wild type. (XML)

**SBML Model S2** Annotated model of osmoadaptation in *pfk26/27Δ*. (XML)

**SBML Model S3** Annotated model of osmoadaptation in *HOG1-att*. (XML)

**SBML Model S4** Annotated model of osmoadaptation in *FPS1-Δ1*. (XML)

**SBML Model S5** Annotated model of osmoadaptation in *gpd1Δ*. (XML)

**SBML Model S6** Annotated model of osmoadaptation in *hog1Δ*. (XML)

**Table S1** Optical density (OD) and cell density (CD) for control and stressed WT cultures. OD in arbitrary units, cell density in  $10^6$  cells/ml. (PDF)

**Text S1** Materials and Methods, data processing, and details concerning modeling, parameter estimation, and response coefficients. (PDF)

## Author Contributions

Conceived and designed the experiments: EK SH EPK CK BN. Performed the experiments: EPK CK BN DK KKH TJ PD. Analyzed the data: EPK CK BN JS EK SH JN. Wrote the paper: EPK CK EK SH JN.

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