# Stress-Induced Transient Cell Cycle Arrest

### Coordinates Metabolic Resource Allocation to

# **Balance Adaptive Tradeoffs**

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## 12 Abstract

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- The ability of a cell to mount a robust response to an environmental perturbation is
- 14 paramount to its survival. While cells deploy a spectrum of specialized counter-measures to
- deal with stress, a near constant feature of these responses is a down regulation or arrest of the
- 16 cell cycle. It has been widely assumed that this modulation of the cell cycle is instrumental in

facilitating a timely response towards cellular adaptation. Here, we investigate the role of cell cycle arrest in the hyperosmotic shock response of the model organism S. cerevisiae by deleting the osmoshock-stabilized cell cycle inhibitor Sic1, thus enabling concurrent stress response activation and cell cycle progression. Contrary to expectation, we found that removal of stress-induced cell cycle arrest accelerated the adaptive response to osmotic shock instead of delaying it. Using a combination of time-lapse microscopy, genetic perturbations and quantitative mass spectrometry, we discovered that unabated cell cycle progression during stress enables the liquidation of internal glycogen stores, which are then shunted into the 24 osmotic shock response to fuel a faster adaptation. Therefore, osmo-adaptation in wild type cells is delayed because cell cycle arrest diminishes the ability of the cell to tap its glycogen 26 stores. However, acceleration of osmo-adaptation in mutant cells that do not arrest comes at the cost of acute sensitivity to a subsequent osmo-stress. This indicates that despite the ostensible advantage faster adaptation poses, there is a trade-off between the short-term benefit of faster adaptation and the vulnerability it poses to subsequent insults. We suggest that cell cycle arrest acts as a carbon flux valve to regulate the amount of material that is devoted to osmotic shock, balancing short term adaptation with long-term robustness.

## **Introduction**

Cells and organisms are constantly challenged in their environment with insults that vary in origin, magnitude and duration. In order to respond to these insults, cells have evolved a large battery of adaptive stress responses that allow them to survive and maintain their homeostasis. Different stress responses show a remarkable diversity in their sensing, regulation, and logic <sup>1–5</sup>. However, an almost constant feature of any stress response is the

involvement of cell cycle slow-down or arrest <sup>6–10</sup>. It is widely assumed that this is because it is advantageous for a cell not to divide during stressful conditions in order to safeguard its own fitness and that of its future progeny <sup>11,12</sup>. Furthermore, by arresting division, resources and energy can be diverted from the replication and division program to the stress response program, allowing it to proceed more efficiently <sup>13</sup>. Despite the near universality of these assumptions, the precise contribution of cell cycle arrest to adaptation remains poorly understood.

46 Upon encountering an environmental perturbation, the model organism Saccharomyces cerevisiae is thought to divert its limited resources such as ribosomes to high-priority transcripts <sup>14,15</sup>. However, this diversion conflicts with other ongoing and resource-intensive processes such as cell cycle progression. Concomitant cell cycle arrest can therefore relieve this resource competition, allowing the cell to adequately mitigate the effects of stress 13. While this model 50 stands to reason intuitively, embedded within that conceptual framework are competing 51 optimization problems that the cell must confront: properly addressing the stress to ensure 52 longevity, but doing so in a time-sensitive manner for cell cycle re-entry to promote reproductive 53 fitness. To understand the tradeoffs involved in a stress response, and interrogate the relative 54 contribution of cell cycle arrest to cellular adaptation and intracellular process optimization, it is necessary to decouple the processes in question.

To explore this paradigm, we used the hyperosmotic glycerol (HOG) response as a convenient framework. The HOG program is a canonical stress response activated by the presence of excess osmolytes in the extracellular environment. The increase in osmotic pressure difference between the inside and outside of the cell drives water out, causing the cellular volume to decrease. At the onset of a step input of hyperosmotic shock, the central HOG mediator, Hog1, rapidly translocates from the cytoplasm to the nucleus where it interacts

with a variety of targets to initiate the production and accumulation of glycerol <sup>16</sup>. The accumulation of glycerol in the cytoplasm re-establishes the osmotic pressure gradient to its basal level, and once volume has been corrected, Hog1 exits the nucleus <sup>17</sup>. Importantly, in addition to the initiation of glycerol synthesis, Hog1 stabilizes Sic1, the stoichiometric inhibitor of b-type cyclins 5 and 6, to transiently arrest the cell in the G1 phase of the cell cycle<sup>11</sup>. Volume restoration and exit of Hog1 from the nucleus also coincides with resumption of cell cycle 68 progression 18. The adaptive translocation pattern of Hog1 has been the subject of many studies 69 for its robust, reproducible and stereotyped pattern, which acts as a real-time reporter of 70 hyperosmotic stress adaptation <sup>19,20</sup>. 71 72 Using HOG as a model system, we investigated the role of transient cell cycle arrest in the adaptive response to hyperosmotic shock. Our approach was to decouple the HOG response program from the canonical cell cycle machinery by removing the stress response-cell 74 cycle link, Sic1, such that both processes proceed simultaneously during osmotic shock. By following Hog1 translocation as a reporter of HOG adaptation, we were able to quantify deviations from Hog1's stereotyped translocation pattern as an indication of an altered stress 77 response. Surprisingly, we found that unabated cell cycle progression during osmoshock accelerated osmo-adaptation as measured by Hog1 translocation. Remarkably, other canonical markers of adaptation such as glycerol production and volume recovery also proceeded faster. These data indicated that cell cycle arrest impedes, rather than facilitates, adaptation to stress. To pinpoint the mechanistic roots of this phenotype, we used mass spectrometry <sup>13</sup>C isotope tracing to probe the differences in metabolic flux between wild type and cell cycle arrest-disabled cells. We discovered that progression in the cell cycle during osmostress initiated catabolism of internal glycogen that is mediated by the enzyme Gph1. Breakdown of 85 glycogen fueled faster glycerol synthesis in the mutant cells, giving them the ability to restore

turgor pressure faster than the wild type. Therefore, cell cycle seems to be the guardian of a
metabolic valve that remains closed when cell cycle is arrested. To investigate what
vulnerabilities arise from opening of this valve, and rationalize why the wild type cells still
implement cell cycle arrest despite the delay it imposes on stress adaptation, we subjected cells
to repeated osmostress pulses. Under repeated pulsing, wild type cells largely return to their
basal morphology while mutant cells lacking cell cycle arrest display physical traits suggestive of
a compromised cell wall. This phenotype accumulates within the mutant population with each
repeated stress pulse. Therefore, adaptation to an osmotic stress proceeds faster when there is
no cell cycle arrest, but this phenotype leaves the cells particularly susceptible to subsequent
osmoshocks. Our findings reveal an example where connection between three important
cellular networks - a stress pathway, cell cycle regulation, and metabolic control - collaborate in
order to strike a balance between mounting a rapid adaptive response to acute threats and

### 100 Results

# Removal of Hog1-mediated cell cycle arrest accelerates adaptation to hyperosmotic shock

To assess the role of cell cycle arrest in adaptation to osmotic shock, we removed the
ability of Hog1 to initiate cell cycle arrest by generating a Sic1 knockout strain (Figure 1A). In
this strain, we tagged Hog1 with mVenus at its endogenous locus to allow visualization of its
nuclear translocation by microscopy. We also incorporated the same Hog1-mVenus construct in
a wild type (WT) strain to allow for comparison of its Hog1 dynamics with those of the *sic1*Δ
mutant. For precise temporal control in applying a step input of osmotic shock, we used a

commercially available microfluidic platform that allowed us to quickly induce osmoshock and monitor Hog1 dynamics in the  $sic1\Delta$  and WT strains by time lapse microscopy. We chose the osmolyte sorbitol as the input to induce hyper-osmotic stress because it is an inert sugar in the presence of glucose  $^{21}$ .

113 Using this experimental setup, we subjected WT and  $sic1\Delta$  mutant cells to a step input of 1.2 M sorbitol osmoshock and monitored Hog1 nuclear translocation dynamics. Following the 114 sorbitol input, Hog1 rapidly translocated to the nucleus in both strains with similar nuclear influx 116 dynamics (Figure 1B). The degree of maximum nuclear enrichment was virtually indistinguishable in both strains, suggesting that the mutant maintains the ability to sense and 117 respond to acute osmotic shock. In the WT strain, Hog1 exited the nucleus and its cytoplasmic 118 levels adapted to the pre-stimulus values within 45 minutes on average, consistent with 119 120 previous reports <sup>20</sup>. Surprisingly, however, the return of Hog1 to the cytoplasm was much faster in the  $sic1\Delta$  cells, occurring on average within 33 minutes (Figure 1C). This constitutes a 121 significant 30% speed-up compared to WT (Figure 1C, right). This acceleration of Hog1 adaptation in non-arresting cells was not a sorbitol-specific effect, since experiments carried out with 0.6 M NaCl also showed the same phenotype (Supplemental Figure 1A, B). The Sic1 protein is known to be a G1-specific regulator<sup>22</sup>. However, since we observe this phenotype averaged over the entire asynchronous population, we are likely underestimating the effect of its 127 deletion.

The faster Hog1 response in  $sic1\Delta$  cells can be the result of a breakdown of coordination between the regulatory osmotic response and turgor pressure of the cell. If this were the case, then Hog1 would recover its cytoplasmic localization without full recovery in other physiological parameters such as cellular volume and glycerol accumulation necessary for this recovery. On the other hand, if the fast Hog1 adaptation were the result of an acceleration within an intact recovery program, then the profile of volume recovery and glycerol accumulation should also be accelerated. We therefore investigated these critical phenotypes to see if the integrity of the adaptive program is maintained despite accelerated Hog1 dynamics in the  $sic1\Delta$  strain.

136 First, we measured internal glycerol content using a colorimetric assay following a 1.2M sorbitol input over 60 minutes in both the WT and  $sic1\Delta$  strains (Figure 1D). Using this assay, 137 we determined that both WT and mutant cells upregulated their glycerol production by 138 139 approximately 5-fold at the end of the 60 minutes. However, while the WT cells did not begin to dramatically upregulate glycerol production until 30 minutes after the onset of stress, sic1\Delta 140 began increasing glycerol synthesis only 15 minutes after stress (Figure 1D, inset). In 141 agreement with this finding, quantification of cellular surface area as a surrogate for volume also 142 showed a faster recovery for the mutant relative to the WT (Figure 1E). While this volume recovery phenotype was reproducible, its extent was slightly lower than the Hog1 and glycerol 144 145 phenotype, showing ~15% difference between WT and mutant (Figure 1E, right). This could be due to the difficulty in cell tracking and quantification of surface area, or to a strict upper limit on the expansion properties of the cell wall <sup>23</sup>. Taken together, the three phenotypes strongly support the hypothesis that the mutant has the same coordinated osmotic stress response as the WT type, but with faster dynamics that ensue from the inability of these cells to arrest their cell cycle during osmotic shock.

# 151 Glycerol production is accelerated by using internal sources in mutant that lacks cell 152 cycle arrest

Because Hog1 translocation, glycerol production and volume all correlate with faster recovery dynamics in the mutant strain, we next sought to investigate if another cellular process was fueling the accelerated production of glycerol, the effector molecule for osmotic shock recovery. We hypothesized that in the *sic1*Δ strain, a surplus of carbon material from central glycolysis could be shunted into glycerol production, thus resulting in heightened glycerol synthesis.

159 One possible scenario for this to happen is one in which the deletion of Sic1 augments the ability of cells to import extracellular glucose, resulting in a greater amount of carbon material entering glycolysis to be directed towards glycerol production. To test this scenario we 161 performed a mass spectrometry <sup>13</sup>C isotope tracing experiment to compare the external glucose incorporation rate between the mutant and WT strains. In this experiment, cells were grown in 163 <sup>12</sup>C glucose media. At time zero, an aliquot of cells was transferred onto filter paper over a 164 vacuum manifold and continuously perfused with fully-labeled <sup>13</sup>C glucose media with and 165 without 1.2 M sorbitol for various durations before quenching the sample (Figure 2A, 167 Supplemental Figure 2A). A pilot experiment (not shown) suggested that turnover rates of glycolytic intermediates occur on the order of seconds, with intermediates reaching steady-state 168 after 1 minute. Therefore, we chose to quench samples at 10, 20, 30, 45 and 60 seconds in 169 order to capture the rate at which the internally <sup>12</sup>C-enriched glycolysis intermediates are degraded and newly synthetized metabolites incorporate the perfused <sup>13</sup>C. Consistent with a fast turn-over of these metabolites, we observed a rapid decay of <sup>12</sup>C enrichment among the glycolysis intermediates within seconds. In the examples of glucose-6-phosphate (G6P) and fructose bisphosphate (FBP), the  $sic1\Delta$  strain incorporated <sup>13</sup>C with a slower rate than the WT in the presence and absence of osmotic shock (Figure 2B, Supplemental Figure 2B). Using the decay rate of <sup>12</sup>C enrichment as a surrogate for extracellular glucose incorporation, we derived  $^{13}$ C incorporation rates of 0.117 s<sup>-1</sup> and 0.04 s<sup>-1</sup> for WT and sic1 $\Delta$ , respectively, during osmotic 177 shock for G6P. Similarly, FBP incorporated <sup>13</sup>C at rates of 0.12 s<sup>-1</sup> and 0.064 s<sup>-1</sup> for WT and 179  $sic1\Delta$ , respectively. The approximately 2-to-3-fold slower <sup>13</sup>C incorporation rate in the  $sic1\Delta$ 

mutant was consistent for all glycolysis intermediates we targeted, from upper glycolysis in G6P to phosphoenolpyruvate (PEP) in lower glycolysis (Supplemental Figure 2B,C). Therefore, it is clear that the faster recovery of the sic1\Delta mutant is unlikely to be ascribed to import of external glucose, since the mutant is slower at <sup>13</sup>C incorporation.

184 Slower glucose incorporation often has the implication of a reduced doubling rate, which has been suggested to confer an advantage during stress adaptation <sup>24,25</sup>. We measured the 185 growth rate of the sic1∆ mutant and observed that it has a doubling time of 3.4 hours compared 186 to 2.1 hours of the WT in defined media (Supplemental Figure 3A). To assess whether slower 187 growth could be correlated to the accelerated osmoshock adaptation, we compared the growth rate of all strains used in this study (see subsequent sections for different strains, including 189 those with different deletions in metabolic genes) against their respective adaptation time to an 191 osmotic shock (Supplemental Figure 3B). We found a weak correlation (R =0.34) between the growth rate of the strains tested and their adaptation time to osmotic stress, suggesting that while growth may contribute to the accelerated phenotype, it is unlikely to be the only factor.

The difference in external glucose incorporation rates between WT and  $sic1\Delta$  suggest that the mutant is not importing more extracellular carbon material into glycolysis. To further investigate the source of its accelerated glycerol production, we reasoned that if the excess carbon material was not from extracellular sources, then it was conceivable that the glycerol synthesis in the  $sic1\Delta$  strain could be assisted by diverted intracellular carbon stores. To investigate this possibility, we again turned to <sup>13</sup>C mass spectrometry isotope tracing to test whether, and to what extent, intracellular carbon was used for glycerol production in both strains. In order to selectively enrich internal stores with a unique carbon isotope, we grew the cells on <sup>12</sup>C glucose, and 5 minutes before time zero, resuspended the cells in fully-labeled <sup>13</sup>C 202 203 glucose. We then continued this treatment with and without osmoshock (1.2M sorbitol) for the

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remainder of the experiment, collecting samples until 30 minutes after osmoshock (Figure 2C).

Based on the rapid turn-over of glycolytic intermediates (Supplemental Figure 2B, C), we

reasoned that within 5 minutes all glycolysis intermediates will be <sup>13</sup>C enriched, but

macromolecules such as storage carbohydrates would remain enriched in <sup>12</sup>C due to their

slower turn-over <sup>26,27</sup>. Given that the cell is only provided <sup>13</sup>C carbon at the onset of osmotic

shock, and glycolysis intermediates likely enriched <sup>13</sup>C in the 5 minutes before stress, any

detection of <sup>12</sup>C after HOG activation would implicate the liquidation of an internal

macromolecule.

212 By monitoring the panel of targeted metabolites in glycolysis and glycerol precursors, we 213 detected low <sup>12</sup>C enrichment along central glycolysis, suggesting that both strains liquidated internal stores during osmoshock. The dynamic pattern of <sup>12</sup>C enrichment of metabolites for the 215 WT cells showed a pattern of low and unchanged level (FBP and PEP) or a level that declined as a function of time (G6P and DHAP) (Figure 2D, Supplemental Figure 4B). The temporal enrichment dynamics of the  $sic1\Delta$  mutant, however, were markedly different showing a transient 217 peak of <sup>12</sup>C enrichment (indicating incorporation of internal carbon stores) followed by a decrease for all metabolites (Figure 2D, Supplemental Figure 4B). In the first step of glycolysis, the metabolite G6P started with a low <sup>12</sup>C enrichment, which increased and peaked between 1 220 and 5 minutes and subsequently decreased. The same pattern was present for glycolysis 221 intermediates FBP and PEP. More importantly, this same transient <sup>12</sup>C signature was present in the glycerol production branch represented by dihydroxyacetone phosphate (DHAP). This 223 transient  $^{12}$ C enrichment pattern in the  $sic1\Delta$  mutant suggests that  $^{13}$ C in glycolysis 224 intermediates is temporarily replaced by <sup>12</sup>C from internal stores in a flux that traverses the 225 metabolic route to glycerol production. However, after this wave, external <sup>13</sup>C is incorporated 226 again in metabolites, underlying the subsequent decline in <sup>12</sup>C enrichment.

It is worth noting that in the absence of stress, the WT had a higher basal  $^{12}$ C enrichment across the metabolites targeted, likely due to a higher basal turnover rate of macromolecules, and consistent with its faster growth rate compared to the  $sic1\Delta$  strain  $^{26}$ . Only during the onset of stress did the  $sic1\Delta$  mutant have a brief, higher  $^{12}$ C enrichment with the aforementioned peak throughout (Figure 2C, Supplemental Figure 4B). Collectively, these data strongly suggest that, unlike the WT,  $sic1\Delta$  briefly shunts internal carbon stores as extra flux into glycerol production at the beginning of osmotic shock.

Interestingly, the immediate precursor to glycerol, glycerol-3-phosphate (GL3P), had an order of magnitude greater proportion of  $^{12}$ C compared to the other metabolites tested (Figure 2D). The increase in  $^{12}$ C could be attributed to back-flux from existing  $^{12}$ C-enriched glycerol by way of a futile cycle to degrade excess ATP during severe stress  $^{28}$ . Despite the discrepancy between GL3P and DHAP, the  $sic1\Delta$  strain was still enriched with a greater amount of  $^{12}$ C than the WT, likely reflecting the convergence of the aforementioned futile cycle and internal carbon liquidation.

## 242 Internal glycogen is liquidated using the Gph1 enzyme and shunted into glycerol 243 production in mutant that lacks cell cycle arrest

244 Previous studies have established links between cell cycle progression and central
245 metabolism as integral to cellular physiology <sup>29–31</sup>. These links are mediated mechanistically
246 through biochemical interactions where CDK1 (Cdc28) activates storage catabolism enzymes
247 for trehalose and glycogen (Nth1 and Gph1, respectively) <sup>32,33</sup>. Given these data, we
248 hypothesized that during osmotic shock, the *sic1*Δ mutant could activate storage catabolism
249 enzymes through unabated cell cycle progression, resulting in a burst of glycolytic flux that was
250 then shunted into excess glycerol production (Figure 3A). Further, we predicted that by coupling

sic1Δ with either a Nth1 or Gph1 knockout, we could rescue the mutant to the WT phenotype,
 for example as measured by Hog1 localization dynamics.

253 Cellular trehalose levels have been widely established as mediators of stress recovery <sup>34</sup>. However, surprisingly, in a  $sic1\Delta nth1\Delta$  mutant, Hog1 nuclear levels still adapted significantly faster than in WT following osmotic stress (Supplemental Figure 5A, B), suggesting that trehalose is not the main liquidated internal carbon source. However, when we coupled the 256  $sic1\Delta$  deletion with a knockout of Gph1 ( $sic1\Delta gph1\Delta$  mutant), Hog1 adapted nearly 30% slower, 257 258 closely resembling WT recovery time (Figure 3B, C). When we combined these genetic perturbations in a  $sic1\Delta nth1\Delta gph1\Delta$  strain, Hog1 again adapted in time comparable to WT, 259 supporting the notion that the Gph1-mediated breakdown of glycogen is the main driver of the 260 accelerated  $sic1\Delta$  phenotype (Supplemental Figure 5C). To ensure that the  $gph1\Delta$  rescue is 261 specific to mitigate the effect of sic1\Delta, and not a broad glycolytic flux perturbation irrespective of 262 genetic background, we tested whether the absence of Gph1-mediated glycogen breakdown 263 affected an otherwise WT Hog1 response. Because glycogen catabolism is halted due to 264 normal cell cycle arrest,  $qph1\Delta$  cells should show a WT adaptation. Indeed, Hog1 adaptation 265 time following osmostress in  $gph1\Delta$  cells is nearly indistinguishable from the WT response 266 267 (Supplemental Fig 5A, B). Consistent with the rescue in Hog1 dynamics, we observed that the deletion of Gph1 counter-acted the  $sic1\Delta$  effect and reduced the glycerol accumulation at 15 minutes in a manner commensurate to the WT rate (Figure 3D). We attempted to measure cellular glycogen levels and did not observe a significant difference within the first 15 minutes of 270 response between strains (data not shown). This is consistent with the notion that potentially 271 undetectable changes in internal glycogen can be converted to substantial changes in glycerol 272 content, given the massive polysaccharide nature of glycogen and the 3-carbon composition of 273 274 glycerol.

## 275 Accelerated recovery due to glycogen storage liquidation during osmotic shock prioritizes faster adaptation over robustness to repeated insults

277 Bypassing cell cycle arrest during osmotic shock results in accelerated recovery due to cell cycle-mediated carbon flux shunted into glycerol production. This suggests that the sic1Δ mutant might have an advantage upon one instance of osmotic stress. Conceivably, multiple 279 instances of osmotic shock can amplify the advantage that  $sic1\Delta$  cells have over WT cells. 280 Alternatively, the faster recovery advantage of  $sic1\Delta$  cells might come at the cost of other 281 282 vulnerabilities that are only revealed dynamically 35.

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To test the endurance of the  $sic1\Delta$  mutant upon a series of osmotic shocks, we subjected cells to the same 90 minute step input of 1.2 M sorbitol as before, but followed by three 45 minute shocks separated by 5 minutes (Figure 4A). At the end of each step input, we calculated the relative adaptation time using Hog1 nuclear residence as a metric and also assessed the visible physical characteristics of cells. After the first osmotic shock, both strains adapt as previously shown, with the  $sic1\Delta$  mutant recovering faster than its WT counterpart (Figure 4B). Following the first step input, Hog1 adaptation to subsequent osmostress inputs proceeded faster (around 50% faster in this case) in both WT and mutant strains due to accumulation of glycerol from the previous cycle and consistent with previous reports 35 (Figure 4C). However, in the subsequent pulses, many  $sic1\Delta$  mutant cells started exhibiting morphological differences from their WT counterparts. These cells displayed a deflated phenotype with visible material accumulated in the nearby vicinity, suggesting the cell lysed and released intracellular debris. This morphology is consistent with a breakdown of cell wall integrity (Supplemental Video), which is a common mechanism of death in serial osmotic shock perturbations <sup>23</sup>. Assessment of this phenotype revealed a marked increase in cells that have a breakdown in their cell wall integrity for each subsequent pulse in  $sic1\Delta$  cells. By the end of the 298

fourth step input of osmotic shock, 25% of  $sic1\Delta$  cells displayed a morphology consistent with a 300 compromised cell wall, while only 5% of WT cells displayed a similar phenotype (Figure 4D, E). Interestingly, the  $sic1\Delta gph1\Delta$  genetic background shares the same acute vulnerability as  $sic1\Delta$ 301 302 to repeated osmotic pulses despite its adaptation that resembles that of the WT. However, the triple mutant  $sic1\Delta nth1\Delta gph1\Delta$  is able to recover its morphology after sequential osmoshocks in 303 a manner that is more similar to the WT (Supplemental Figure 6D). This difference in 304 305 responsiveness suggests unique roles for the breakdown of trehalose and glycogen that warrant further investigation. Nonetheless, it is clear that while the accelerated adaptation ostensibly 306 307 provides an advantage in the face of a single step input of osmotic shock, the  $sic1\Delta$  mutant is 308 severely ill-suited for repeated insults.

## Discussion

310 Alterations in cell cycle dynamics have a fundamental presence in many adaptations to stress, but a mechanistic understanding of its role has long been outstanding. Here we 311 attempted to understand some aspects of the role of cell cycle arrest in the context of the 312 well-studied HOG pathway and associated osmotic stress. Our approach was to decouple the 313 HOG stress program from the cell cycle machinery and monitor the nuclear localization dynamics of HOG master effector (Hog1). This experiment revealed that the stereotyped 315 316 behavior of Hog1 accelerated when the cell cycle could still proceed during osmoshock. We confirmed that the HOG program was still competent under these circumstances by measuring 317 other canonical hallmarks of osmoshock recovery - internal glycerol content and cellular volume. Observing that both proceed faster in the mutant, we utilized quantitative mass spectrometry to 319 implicate internal glycogen liquidation by the mutant as a route by which glycerol synthesis

increases and mediates faster adaptation to the stress. In strong agreement with this insight, deletion of the glycogen catabolism enzyme Gph1 in a sic1Δ background rescues the Hog1 translocation and glycerol accumulation phenotypes. Intrigued by the observation that the WT is 324 not optimized with respect to the speed of its recovery, we hypothesized that the mutant, which has faster dynamics, might have vulnerabilities that the WT can circumvent. Following this reasoning, we identified a critical failure mode of the mutant by subjecting it to multiple step 326 inputs of osmotic shock. The mutant adapts faster and recovers its basal morphology after the 327 first osmotic shock, but trades its faster initial adaptation for a compromised cell wall in 328 subsequent osmoshocks. Meanwhile, the WT adapts slower after the initial osmotic shock, but 329 maintains nearly 95% consistent physical traits throughout the experiment, thus highlighting the 330 dichotomy between apparent short-term gain versus long-term resilience against a dynamic 331 332 environment.

We believe that the main contribution of this work is two-fold. First, our investigations provide a higher resolution dissection of the interconnection between three crucial cellular pathways: the cell cycle, the HOG stress response, and carbon metabolism. Contrary to expectation, this connection is not perfectly tuned to maximize the speed of adaptation to stress. In fact, the connection of the HOG pathway to the cell cycle diminishes the ability of the cell to recover rapidly following osmostress, and ablation of this connection allows the cell to recover a substantial 30% faster. The cell cycle seems to be the gatekeeper of a metabolic valve that can augment carbon flow into glycerol from internal resources, but this valve remains shut in WT cells. Opening of this valve in mutant ( $sic1\Delta$ ) cells seems to mediate their faster recovery, as evidenced by a  $gph1\Delta$  mutant in which deletion of the enzyme presiding over the internal flux 342 343 from glycogen to glycerol abolishes the fast osmostress recovery. Therefore, our data provide

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344 additional mechanistic details to an intricate interplay of pathways that together set the cellular 345 recovery tempo.

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The second contribution of this work is to formulate an instance in which cells seem to navigate a delicate functional balance, sacrificing the brief advantage of faster recovery from an insult for robustness to future environmental changes (Figure 4F). This is perhaps a prompt to revise our view of how to interpret the measured dynamics of stress responses and our assumptions about how cells mobilize their resources to combat stress. It is clear that at least in the example of osmostress, S. cerevisiae cells do not maximally mobilize their internal carbon sources to combat the stress, and hence sacrifice substantial speed in their recovery. It is also evident that the cell cycle serves as an arbiter and enforcer of this suboptimal performance. Since the mutant that evades the speed limitation shows tremendous vulnerability to repeated stress, one is compelled to at least hypothesize that this cell cycle control has evolved to alleviate such vulnerability in an environment where repeated or oscillating stress might be more probable.

The work presented here reframes cell cycle arrest in a mechanistic light as being a mediator of a slower adaptation response to hyperosmotic shock. In future investigations, it 360 would be interesting to use a similar approach with conditional or inducible mutants to test for the role of cell cycle arrest in other stresses, potentially discovering similar metabolic flux control mechanisms or roles more tailored to specific stresses. Alternatively, expanding the scope of this question to higher eukaryotes could further illuminate the complex relationship between cell cycle, metabolism and stress response, which has been implicated in several pathologies <sup>36</sup>. 364 More broadly, it is clear that as we begin to explore how multiple pathways collaborate to allow a 365 cell to navigate its complex environment, we need to revisit statements about functional 366 allocations and re-explore plausible but exceedingly simple assignment of roles and 367

assumptions of unifunctional optimality of any one pathway. We hope that the data presented in this work help to form a basis for such investigations, initiated by our quantitative exploration of the ubiquitous role that cell cycle arrest plays in stress adaptation.

## Materials and Methods

#### 372 Strain Construction

The base *S. cerevisiae* strain used in this study is w303. Hog1-mVenus at the
endogenous locus was generated by ordering oligos of 40 bp homology 5' upstream and 40 bp
homology downstream of the stop codon, PCR amplifying the mVenus-HIS3 cassette, and
transforming as previously described<sup>37</sup>. To knockout genes, 80 bp of homology 5' to the start
codon and 3' of the stop codon was used to PCR amplify a selection marker cassette and
transformed as described above. PCR products using oligos in the 5' UTR and internal to the
selection cassette were used to verify knockouts and insertions.

Name	Background	Genotype	Description
yARB001	w303a	HOG1-mVenus-HIS3	"Wild type" strain
yARB002	w303a	HOG1-mVenus-HIS3,SIC1::TRP1	sic1∆ strain
yARB003	w303a	HOG1-mVenus-HIS3,SIC1::TRP1, NTH1::NAT	sic1∆nth1∆ strain
yARB004	w303a	HOG1-mVenus-HIS3,SIC1::TRP1, GPH1::LEU2	sic1∆gph1∆ strain
yARB005	w303a	HOG1-mVenus-HIS3,SIC1::TRP1, GPH1::LEU2, NTH1::NAT	sic1Δnth1Δgph1Δ strain

yARB006	w303a	HOG1-mVenus-HIS3, GPH1::LEU2	gph1∆ strain

#### 380 Growth Conditions

Single colonies were picked and inoculated from auxotrophic SD (6.7 g/L Bacto-yeast nitrogen base without amino acids, 2 g/L complete amino acid mix, and 20 g/L dextrose) agar plates. SDC liquid media used throughout the study consisted of 6.7 g/L Bacto-yeast nitrogen base, 2 g/L complete amino acid mix, and 20 g/L dextrose.

#### 385 Microscopy

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386 Time-lapse microscopy was collected on a Nikon Ti inverted scope 40x air objective, with Sutter XL lamp illumination and a Hamamatsu Flash 4.0 camera. YFP (515 ex/528 em) 387 channel was collected using a Chroma CFP/YFP filter set with an exposure time of 300 ms. Automated image acquisition was controlled by Nikon NIS Elements proprietary software. The 389 CellAsics Onix2 Microfludic platform was used to control the changing of media. Pressure of the 390 media perfusion was held constant at 10.8 kPa in a microfluidic plate designed to trap haploid 391 yeast (Millipore). To ensure that the yeast cells adapted to conditions within the microfluidic 392 chamber, cells were perfused with normal SDC media for 90 minutes prior to the osmotic shock 393 in all experiments. 394

#### 395 Image Processing

The nucleus of each cell was defined as the mean pixel intensity of the brightest 5% of pixels over the segmented cell in the YFP channel. The remainder of the segmented cell outside the brightest 5% was defined as the cytoplasm. Cell tracking and quantification of nuclear/cytoplasmic enrichment was done using automated yeast cell tracking software implemented in Matlab <sup>38</sup>. Nuclear enrichment is plotted as the population average nuclear to cytoplasmic ratio divided by the average three time points before the onset of osmotic shock

minus one, as previously described <sup>18</sup>. Cellular volume was calculated from segmentation of an out of focus brightfield image using the Nucleaizer web interface (<a href="www.nucleaizer.com">www.nucleaizer.com</a>). The surface area of each cell in pixels was then converted into an approximate volume as described previously <sup>18</sup>. This volume was normalized by the three time points before the onset of osmotic shock.

#### Mass Spectrometry

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Samples were grown overnight to saturation and diluted in SDC media. Cultures were grown to mid-log and 1 mL was transferred to a 0.45 µm PVDF membrane filter paper (Millipore), fixed atop a vacuum manifold. Cells were continuously perfused with either normal SDC media or 1.2 M sorbitol in SDC media for the indicated durations (Figure 2A). At the end of the perfusion period, the filter paper containing cells was immediately transferred to a 2 mL quenching solution of 40:40:20 Methanol:Acetonitrile:H<sub>2</sub>O chilled to -20 °C. After 2 hours incubation at -20 °C, quenching solution plus cells was transferred to a conical tube and dried for approximately 7 hours *in vacuo* and resuspended in 45 µL H<sub>2</sub>O.

In the experiment described in Figure 2C, five minutes prior to time zero 50 mL culture were transferred to a conical tube, spun down at 2000 RPM for 2 minutes and resuspended in Land 18 25 mL fully labeled 13 C glucose SDC. After time zero, at the indicated time points, 1 mL of culture was transferred to the same filter paper vacuum manifold described above, and after media washed through the sample was quenched as described above.

Collected compounds were analyzed using an LC-MS/MS mass spectrometer system
consisting of a 1290 Infinity LC (Agilent Technologies) coupled to a 5500 QTRAP triple
quadrupole mass spectrometer (AB Sciex) in negative mode and with multiple reaction
monitoring (MRM) scan type. Five µL of metabolite extracts were injected on an Agilent
PoroShell 120 HILIC-Z column (150 x 2.1 mm, 2.7 µm; Agilent, Santa Clara, CA) using a mobile

phase A (water, 10 mM ammonium acetate, 5 μM medronic acid, pH 9) and mobile phase B (90% acetonitrile, 10% water, 10 mM ammonium acetate, 5 μM medronic acid, pH 9) at a constant flow rate of 250 μl/min; Initial conditions: 10% A, 2 min: 10% A, 12 min: 40% A, 15 min: 40% A, 16 min: 10% A, 24 min: 10% A. The MRM settings were adapted from Yuan et al <sup>39</sup>. The raw data were processed and analyzed using custom software in Matlab (Mathworks), and <sup>13</sup>C fractional labeling was corrected for natural abundance of <sup>13</sup>C isotopes as previously described <sup>40</sup>.

#### Intracellular glycerol assay

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Each strain was inoculated into SDC media, grown overnight at 30 °C, split and diluted 434 into six 600 µL 0.1 OD600 cultures in a 96 well 2 mL plate. Once the cells were in log phase 435 growth, 600 µL of 2.4 M sorbitol was added to one well at time points of 60 minutes, 30 minutes, 436 15 minutes, 10 minutes, 5 minutes and 0 minutes. After time zero, 200 μL of each culture was 437 transferred to a separate Corning 3904 96-well assay plate plate for an OD600 reading, and the 438 remaining 1 mL immediately spun down for 2 minutes at 2000 RPM. Cells were washed in 400 439 μL H<sub>2</sub>O, and spun down again for 2 minutes at 2000 RPM. The culture was then resuspended in 440 150 μL H<sub>2</sub>O, and left to incubate for 15 minutes at 95 °C. Following incubation at 95 °C, cells 441 were vortexed for 2 minutes and promptly spun down for 10 minutes at 4000 RPM. After the pelleting of cell debris, 100 µL of supernatant was carefully removed and transferred to a separate plate and kept at 4 °C. Colorimetric glycerol assays were acquired using a commercial kit (Sigma) where the provided assay powder was resuspended in 40 mL of distilled H<sub>2</sub>O. For 445 each sample, 5 µL of supernatant was added to 400 µL of glycerol free reagent solution, and left 446 to incubate at room temperature for 15 minutes hidden from light. After 15 minutes, 200 µL of 447 the glycerol free reagent solution-sample mixture was transferred to a separate plate and the 448 OD540 was acquired for each sample on a Tecan Spark 10M plate reader. To account for 449

differences in cell density across samples, the 540 nm readings were normalized by their OD 600 nm reading values.

#### 452 Growth Assay

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Each strain was inoculated into SDC media overnight, reached saturation and diluted the following morning to an OD600 of 0.1. After dilution, 200 μL of culture was transferred to a Corning 3904 96-well assay plate and grown at 30 °C while shaking. Optical density readings were collected at 600 nm every 20 minutes until saturation on a Tecan Spark 10M plate reader.

Cell Morphology Quantification

At the end of each 45 minute step input of osmotic shock, each cell was manually assessed for cell cycle re-entry and return to basal morphology. Cells that either did not show continued cell cycle progression, or displayed a visible change in refractive index reflective of a change in morphology were labeled with altered morphology.

#### 462 Main Figure Captions:

Figure 1: Removing cell cycle arrest by deletion of Sic1 accelerates the HOG adaptation 463 program during osmotic shock. A) A simplified schematic of the HOG pathway showing its 464 465 coupling to cell cycle arrest and glycerol production. In a cell cycle mutant strain  $sic1\Delta$ , we asked whether the removal of stress-induced cell cycle arrest affects the adaptive response. B) 467 Representative time lapse images of endogenously-tagged Hog1-mVenus following step input of 1.2 M sorbitol osmotic shock to WT (top) and  $sic1\Delta$  (bottom) cells. Quantification of all cells presented in Panel C. C) Top: Cartoon depicting the translocation dynamics and quantification 469 of adaptation time. Bottom: Quantification of the WT (blue) and  $sic1\Delta$  (orange) Hog1-mVenus 470 471 nuclear enrichment in the experiment described in Panel B. Shaded regions represent the 472 standard error of the mean (SEM) of n=3 biological replicates. Right: Quantification of Hog1

adaptation for WT (blue) and  $sic1\Delta$  (orange). Values are normalized to the average of WT. Error bars represent the SEM of n=3 biological replicates. \*P-value<0.05; two-sided Student's t-test. D) Top: Cartoon schematic depicting the intracellular accumulation of glycerol. Bottom: 475 Quantification of internal glycerol as a function of time for WT (blue) and  $sic1\Delta$  (orange) to a 476 step input of 1.2 M sorbitol osmotic shock. Measurements are taken using a colorimetric assay. 477 478 Error bars represent the standard deviation for n=3 biological replicates. Inset: the change in glycerol, calculated as the difference between two time points for data in Panel D, is plotted as a 479 function of time. \*P-value<0.05; two-sided Student's t-test. E) Top: Cartoon schematic depicting 480 volume recovery and quantification of its adaptation time. Bottom: Change in volume of the WT 481 (blue) and  $sic1\Delta$  (orange) strains in response to a step input of 1.2 M sorbitol osmotic shock. 482 Shaded regions represent the SEM of n=4 biological replicates. Right: Quantification of volume 483 adaptation time of WT (blue) and  $sic1\Delta$  (orange) volume. Values are normalized to the average 484 485 of WT. Error bars represent the SEM of n=4 biological replicates.

Figure 2: An internal carbon store is shunted towards excess glycerol production during 486 osmotic shock in the sic1∆ mutant. A) Cartoon schematic of the experiment to measure 487 488 extracellular glucose incorporation rates. Cells were inoculated overnight, diluted and outgrown in <sup>12</sup>C glucose. At time zero a 1 mL sample of cells was transferred to filter paper above a vacuum manifold and continuously perfused with fully-labeled <sup>13</sup>C glucose media. A 1.2 M 490 sorbitol input was also administered at time 0. Samples were taken at 10 s, 20 s, 30 s, 45 s, and 491 60 s and transferred to quenching solution. B) <sup>12</sup>C enrichment over time of central glycolysis 492 metabolite glucose-6-phosphate (G6P) (Left) and fructose bisphosphate (FBP) (Right). Traces 493 shown are WT (blue) and  $sic1\Delta$  (orange) strains for experiment described in Panel A. Error bars 494 495 represent the standard deviation of n=2 technical replicates. C) Cartoon schematic of

496 experiment to test internal carbon enrichment of targeted metabolites. Cells were inoculated overnight, diluted and outgrown in <sup>12</sup>C glucose. Five minutes prior to time zero, cells were 497 resuspended in fully-labeled <sup>13</sup>C glucose. At time zero the culture was diluted 1:1 with 2.4 M sorbitol in fully-labeled in <sup>13</sup>C glucose. At the indicated time points, 1 mL of culture was placed 499 on filter paper above a vacuum manifold for the media to wash through, transferred to 500 quenching solution and then measured. D) <sup>12</sup>C enrichment over time for a panel of select 501 metabolites in glycolysis and glycerol production. Traces shown are the WT (blue) and sic1Δ 502 (orange) strains for experiment described in Panel C. Error bars represent the standard 503 deviation of n=2 technical replicates. 504

Figure 3: Glycogen catabolism enzyme Gph1 mediates expedited glycerol synthesis to 505 506 fuel acceleration phenotype in sic1∆ mutant. A) Schematic depicting the hypothesis that liquidation of glycogen by activation of Gph1 accelerates the Hog1-mediated glycerol 507 production. B) Left: Traces of Hog1 nuclear enrichment over time following 1.2 M sorbitol 508 osmotic shock in the WT (blue),  $sic1\Delta$  (orange),  $sic1\Delta gph1\Delta$  (purple) cells. Shaded regions 509 represent the SEM of n=3 biological replicates. C) Quantification of adaptation time of Hog1 510 nuclear enrichment computed as in Figure 1B. Values are normalized to the average WT. Error bars represent the SEM of n=3 biological replicates. \*P<0.05; two-sided Student's t-test. D) Measurement of internal glycerol over time for the strains shown in Panel B in response to a step input of 1.2 M sorbitol osmotic shock. Measurements are taken using a colorimetric assay. Inset: the change in glycerol, calculated as the difference between two time points for data in Panel D, is plotted as a function of time. Error bars represent the standard deviation for n=3 516 biological replicates. 517

Figure 4: Cell cycle arrest mediates tradeoffs between fast recovery and resilience to multiple instances of osmotic shock. A) Top: experiment schematic representing a series of 519 1.2 M sorbitol osmotic shock step inputs. The first input lasts for 90 minutes and subsequent inputs last 45 minutes, and are separated by 5 minutes. B) Time traces of Hog1 nuclear 521 enrichment of WT (blue),  $sic1\Delta$  (orange). Shaded regions represent the SEM of n=3 biological 522 replicates. C) Quantification of adaptation time of Hog1 nuclear enrichment for WT and sic1Δ 523 strains for data presented in Panel B. Values are normalized to the average first response for 524 the WT strain. Error bars represent the SEM of n=3 biological replicates. D) The percent of cells 525 with altered morphological phenotypes at the end of each step input of 1.2 M sorbitol. Error bars 526 represent the SEM of n=3 biological replicates. \*\*P<0.005; two-sided Student's t-test. 527 E) Representative brightfield images corresponding to the timepoints quantified in Panel D. 528 529 Images depict the compromised cell wall morphology indicated by red arrows in the WT (top) and  $sic1\Delta$  (bottom) strains. F) Conceptual model of the role of cell cycle arrest in hyperosmotic 530 531 shock response.

## 532 Author Contributions

533 A.R.B. and H.E-.S. conceived of the study. A.R.B., K.K, and M.D. collected and processed data.

34 A.R.B., K.K., and H.E-.S. interpreted results. A.R.B. and H.E-.S. wrote and edited the

535 manuscript with input from all authors.

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## **References**

- 545 1. Berry, D. B. & Gasch, A. P. Stress-activated Genomic Expression Changes Serve a
- Preparative Role for Impending Stress in Yeast. *Molecular Biology of the Cell* vol. 19
- 547 **4580–4587** (2008).
- 548 2. Gasch, A. P. et al. Genomic expression programs in the response of yeast cells to
- environmental changes. *Mol. Biol. Cell* **11**, 4241–4257 (2000).
- 550 3. Hohmann, S. & Mager, W. H. Yeast Stress Responses. (Springer Science & Business
- 551 Media, 2007).
- 552 4. Hao, N. & O'Shea, E. K. Signal-dependent dynamics of transcription factor translocation
- controls gene expression. *Nat. Struct. Mol. Biol.* **19**, 31–39 (2011).
- 554 5. Lin, Y., Sohn, C. H., Dalal, C. K., Cai, L. & Elowitz, M. B. Combinatorial gene regulation by
- modulation of relative pulse timing. *Nature* **527**, 54–58 (2015).
- 556 6. Lew, D. J. & Reed, S. I. A cell cycle checkpoint monitors cell morphogenesis in budding

- yeast. J. Cell Biol. **129**, 739–749 (1995).
- 558 7. Moreno-Torres, M., Jaquenoud, M. & De Virgilio, C. TORC1 controls G1–S cell cycle
- transition in yeast via Mpk1 and the greatwall kinase pathway. *Nature Communications* vol.
- 560 6 (2015).
- 8. Rowley, A., Johnston, G. C., Butler, B., Werner-Washburne, M. & Singer, R. A. Heat
- shock-mediated cell cycle blockage and G1 cyclin expression in the yeast Saccharomyces
- cerevisiae. *Mol. Cell. Biol.* **13**, 1034–1041 (1993).
- 564 9. Bellí, G., Garí, E., Aldea, M. & Herrero, E. Osmotic stress causes a G1 cell cycle delay and
- downregulation of Cln3/Cdc28 activity in Saccharomyces cerevisiae. Mol. Microbiol. 39,
- 566 1022–1035 (2001).
- 10. Yano, K. et al. Mih1/Cdc25 is negatively regulated by Pkc1 in Saccharomyces cerevisiae.
- 568 Genes Cells **18**, 425–441 (2013).
- 11. Escoté, X., Zapater, M., Clotet, J. & Posas, F. Hog1 mediates cell-cycle arrest in G1 phase
- by the dual targeting of Sic1. *Nat. Cell Biol.* **6**, 997–1002 (2004).
- 571 12. Clotet, J. et al. Phosphorylation of Hsl1 by Hog1 leads to a G2 arrest essential for cell
- survival at high osmolarity. *EMBO J.* **25**, 2338–2346 (2006).
- 573 13. Ho, Y.-H., Shishkova, E., Hose, J., Coon, J. J. & Gasch, A. P. Decoupling Yeast Cell
- 574 Division and Stress Defense Implicates mRNA Repression in Translational Reallocation
- during Stress. *Curr. Biol.* **28**, 2673–2680.e4 (2018).
- 576 14. Chasman, D. et al. Pathway connectivity and signaling coordination in the yeast
- stress-activated signaling network. *Mol. Syst. Biol.* **10**, 759 (2014).
- 578 15. Zid, B. M. & O'Shea, E. K. Promoter sequences direct cytoplasmic localization and
- translation of mRNAs during starvation in yeast. *Nature* **514**, 117–121 (2014).
- 580 16. Saito, H. & Posas, F. Response to hyperosmotic stress. *Genetics* **192**, 289–318 (2012).

- 581 17. Hohmann, S. Osmotic stress signaling and osmoadaptation in yeasts. *Microbiol. Mol. Biol.*
- 582 Rev. **66**, 300–372 (2002).
- 18. Muzzey, D., Gómez-Uribe, C. A., Mettetal, J. T. & van Oudenaarden, A. A systems-level
- analysis of perfect adaptation in yeast osmoregulation. *Cell* **138**, 160–171 (2009).
- 585 19. Mettetal, J. T., Muzzey, D., Gómez-Uribe, C. & van Oudenaarden, A. The frequency
- dependence of osmo-adaptation in Saccharomyces cerevisiae. Science **319**, 482–484
- 587 (2008).
- 588 20. Granados, A. A. et al. Distributing tasks via multiple input pathways increases cellular
- survival in stress. *Elife* **6**, (2017).
- 590 21. Quain, D. E. & Boulton, C. A. Growth and metabolism of mannitol by strains of
- 591 Saccharomyces cerevisiae. *J. Gen. Microbiol.* **133**, 1675–1684 (1987).
- 592 22. Verma, R. et al. Phosphorylation of Sic1p by G1 Cdk required for its degradation and entry
- into S phase. *Science* **278**, 455–460 (1997).
- 594 23. Banavar, S. P. et al. Mechanical feedback coordinates cell wall expansion and assembly in
- 595 yeast mating morphogenesis. *PLoS Comput. Biol.* **14**, e1005940 (2018).
- 596 24. Blank, L. M. & Sauer, U. TCA cycle activity in Saccharomyces cerevisiae is a function of the
- 597 environmentally determined specific growth and glucose uptake rates. *Microbiology* **150**,
- 598 1085–1093 (2004).
- 599 25. Lu, C., Brauer, M. J. & Botstein, D. Slow growth induces heat-shock resistance in normal
- and respiratory-deficient yeast. *Mol. Biol. Cell* **20**, 891–903 (2009).
- 601 26. Suarez-Mendez, C. A. et al. Interaction of storage carbohydrates and other cyclic fluxes
- with central metabolism: A quantitative approach by non-stationary C metabolic flux
- analysis. *Metab Eng Commun* **3**, 52–63 (2016).
- 604 27. Yuan, J., Bennett, B. D. & Rabinowitz, J. D. Kinetic flux profiling for quantitation of cellular

- 605 metabolic fluxes. *Nat. Protoc.* **3**, 1328–1340 (2008).
- 606 28. Blomberg, A. Metabolic surprises in Saccharomyces cerevisiae during adaptation to saline
- conditions: questions, some answers and a model. *FEMS Microbiology Letters* vol. 182 1–8
- 608 (2000).
- 609 29. Özsezen, S. et al. Inference of the High-Level Interaction Topology between the Metabolic
- and Cell-Cycle Oscillators from Single-Cell Dynamics. *Cell Syst* **9**, 354–365.e6 (2019).
- 611 30. Papagiannakis, A., Niebel, B., Wit, E. C. & Heinemann, M. Autonomous Metabolic
- Oscillations Robustly Gate the Early and Late Cell Cycle. *Mol. Cell* **65**, 285–295 (2017).
- 613 31. Tu, B. P., Kudlicki, A., Rowicka, M. & McKnight, S. L. Logic of the yeast metabolic cycle:
- temporal compartmentalization of cellular processes. *Science* **310**, 1152–1158 (2005).
- 615 32. Ewald, J. C., Kuehne, A., Zamboni, N. & Skotheim, J. M. The Yeast Cyclin-Dependent
- Kinase Routes Carbon Fluxes to Fuel Cell Cycle Progression. *Mol. Cell* **62**, 532–545
- 617 (2016).
- 618 33. Zhao, G., Chen, Y., Carey, L. & Futcher, B. Cyclin-Dependent Kinase Co-Ordinates
- 619 Carbohydrate Metabolism and Cell Cycle in S. cerevisiae. *Molecular Cell* vol. 62 546–557
- 620 (2016).
- 621 34. Nwaka, S. & Holzer, H. Molecular biology of trehalose and the trehalases in the yeast
- Saccharomyces cerevisiae. *Prog. Nucleic Acid Res. Mol. Biol.* **58**, 197–237 (1998).
- 623 35. Mitchell, A., Wei, P. & Lim, W. A. Oscillatory stress stimulation uncovers an Achilles heel of
- the yeast MAPK signaling network. Science vol. 350 1379–1383 (2015).
- 625 36. Jones, R. G. & Thompson, C. B. Tumor suppressors and cell metabolism: a recipe for
- 626 cancer growth. *Genes Dev.* **23**, 537–548 (2009).
- 627 37. Lee, S., Lim, W. A. & Thorn, K. S. Improved blue, green, and red fluorescent protein
- tagging vectors for S. cerevisiae. *PLoS One* **8**, e67902 (2013).

- 629 38. Doncic, A., Eser, U., Atay, O. & Skotheim, J. M. An algorithm to automate yeast
- segmentation and tracking. *PLoS One* **8**, e57970 (2013).
- 631 39. Yuan, M., Breitkopf, S. B., Yang, X. & Asara, J. M. A positive/negative ion-switching,
- targeted mass spectrometry-based metabolomics platform for bodily fluids, cells, and fresh
- and fixed tissue. *Nat. Protoc.* **7**, 872–881 (2012).
- 40. Midani, F. S., Wynn, M. L. & Schnell, S. The importance of accurately correcting for the
- natural abundance of stable isotopes. *Anal. Biochem.* **520**, 27–43 (2017).

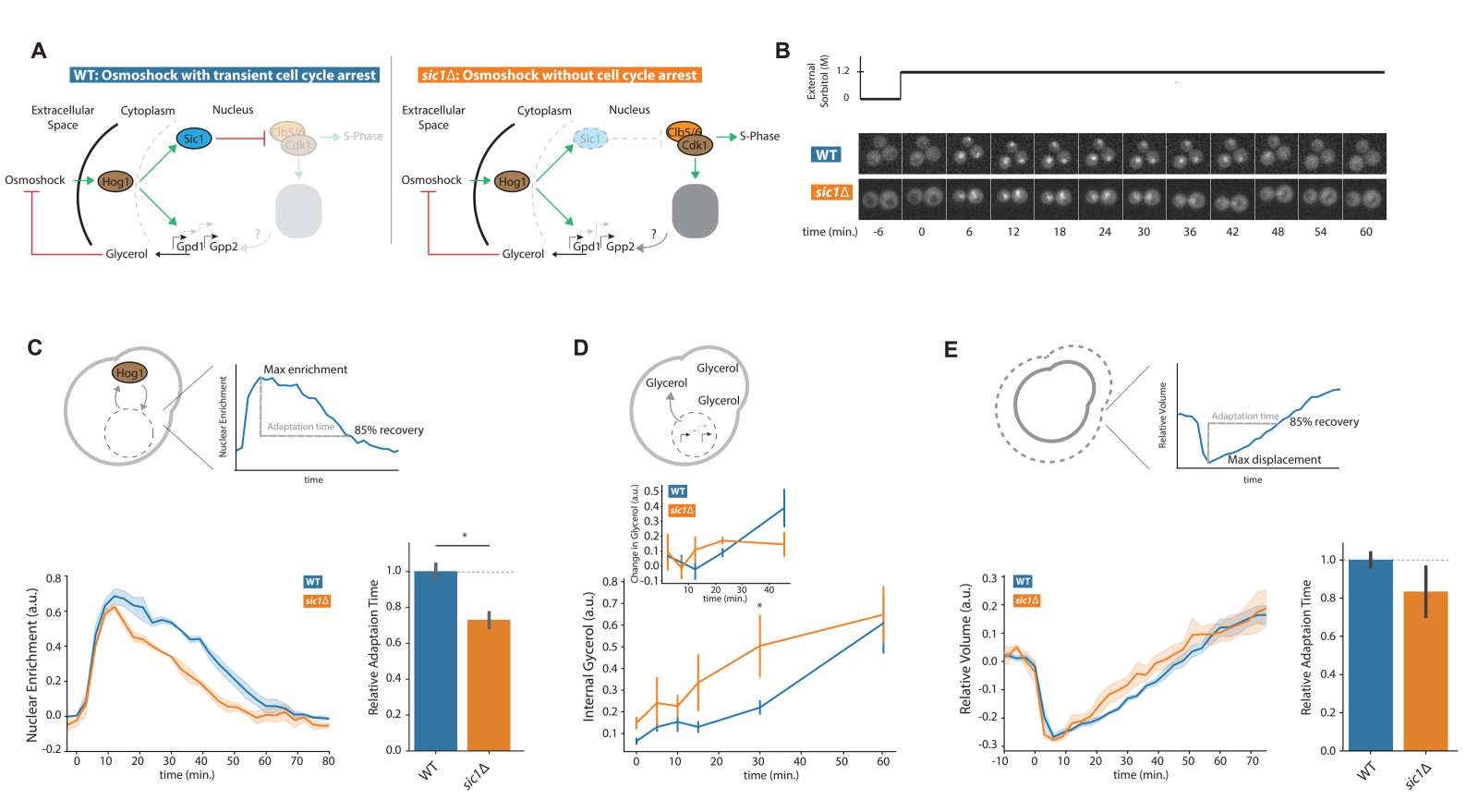
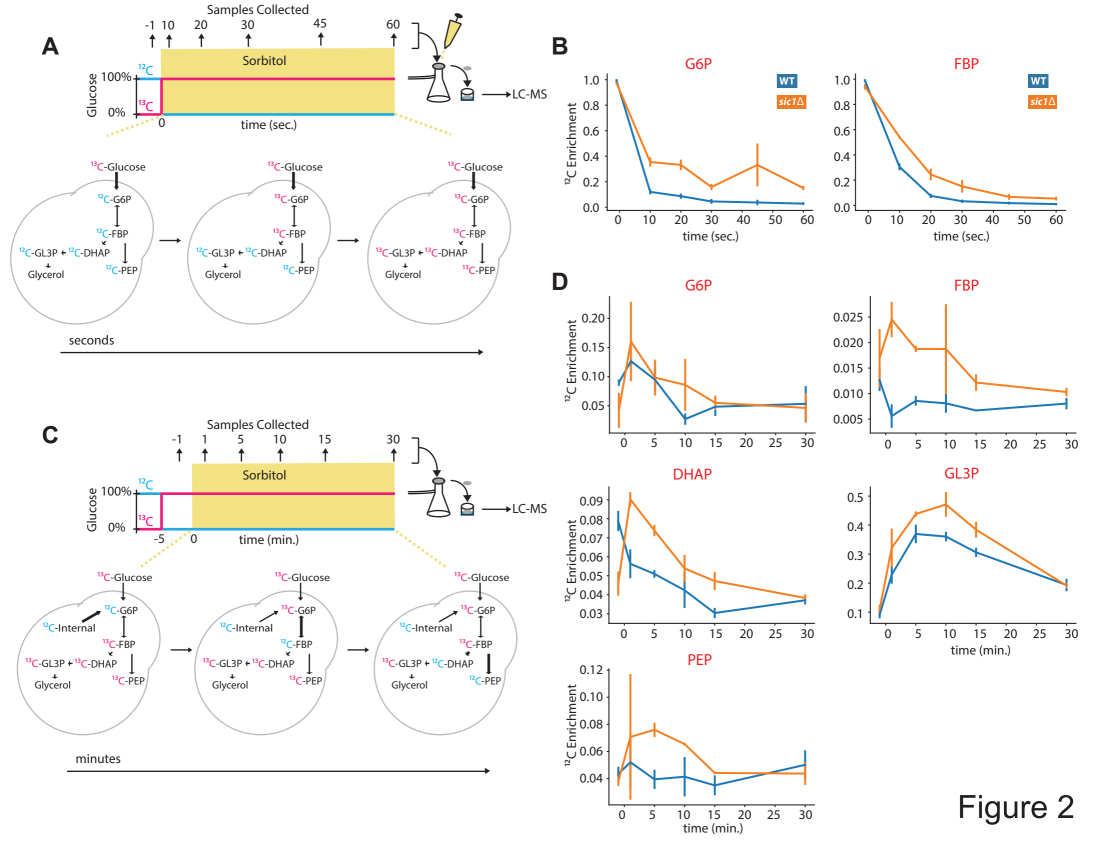


Figure 1



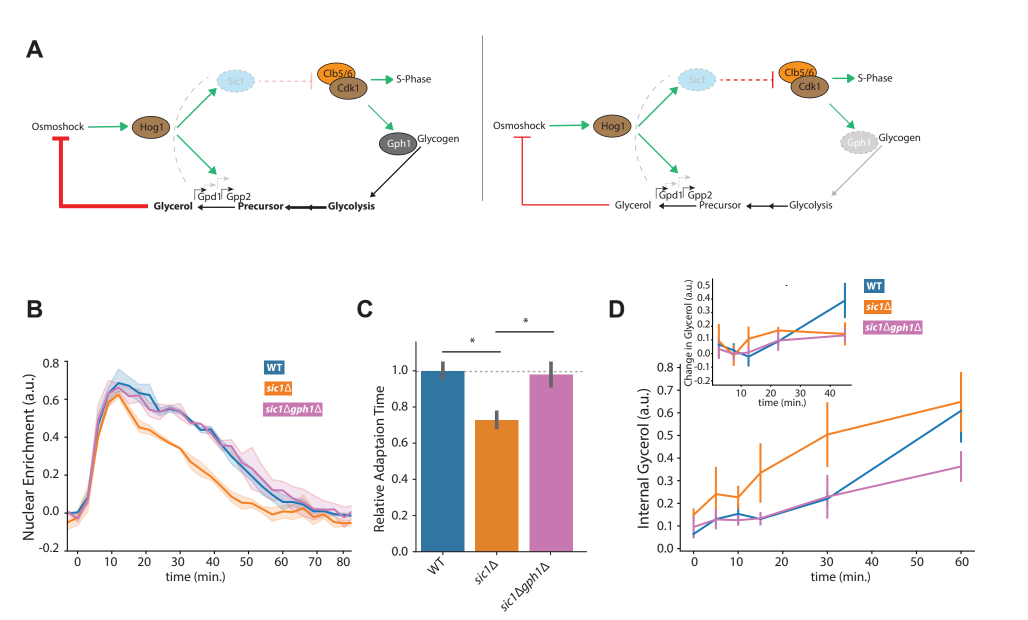


Figure 3

