Matters Arising

Glycerol as the actuator of integral feedback control in yeast osmotic stress signaling

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Abstract

van Oudenaarden and colleagues employ elegant experiments and control theory to model perfect adaptation of the yeast osmotic stress response – precise return of turgor pressure to its optimal, steady-state value despite variation in system parameters and the continued presence of osmotic stress. Their data convincingly show that nuclear signaling and cell volume undergo "robust perfect adaptation" implying integral feedback must restore their steady state values. However they incorrectly map the integrator onto a minimal network that violates assumptions implicit in conventional block diagrams. Using known features of osmotic stress signaling and results presented by the authors, I argue that glycerol concentration – the integral of the rate of glycerol accumulation (synthesis minus leakage) – transforms metabolic energy into increased osmolarity that drives water influx and restoration of turgor pressure. Integral feedback control actuated through glycerol synthesis is logically positioned to provide perfect adaptation and robustness in hyperosmotic stress responses.

Introduction

Robust perfect adaptation is a property of biological feedback control systems that precisely maintains steady-state homeostasis of vital functions in the continued presence of disturbances such as environmental stress or, in the case of desensitization, increases the dynamic range of detection for external signals such as pheromones or environmental nutrients (Yi et al., 2000). In their article "A systems level analysis of perfect adaptation in yeast osmoregulation" Muzzey et al. beautifully demonstrate perfect adaptation of nuclear signaling, cell volume, and thus membrane turgor pressure in osmotic stress. Control theory shows that perfect adaptation predicts and is predicted by integral feedback control (Yi et al., 2000). Muzzey and colleagues argue convincingly and with clever experiments that the data support a single integrating mechanism (or integrators acting in parallel, but not in series). The authors use this point to identify on a minimal network model the location of the integral feedback control mechanism(s).

Analyses of complex control systems depend critically on an accurate working model with clearly defined topology (Riggs, 1970; Romagnoli and Palazoglu, 2006). Here I show that the minimal representation of the osmotic stress response presented in Muzzey et al. violates assumptions implicit in conventional block diagrams (page 115 (Riggs, 1970)). Using known features of osmotic stress signaling and results presented by the authors, I argue that integration of the summed rates of glycerol synthesis and leakage (amount per time) into an accumulating intracellular concentration of glycerol (amount per volume) provides integral feedback control, perfect adaptation, and robustness to the hyperosmotic stress response. In contrast with localization of the integrator in the HOG (high osmolarity glycerol) pathway as proposed by the authors (Muzzey et al., 2009), both HOG dependent and HOG independent processes affect glycerol concentration, which directly drives water influx, osmolarity, and restoration of turgor pressure.

Results

Figure 1 shows a detailed block diagram representation of feedback control based on features of the osmotic stress response using as a general template the feedback control system presented in Figure 10.4 of Romagnoli and Palazoglu (Romagnoli and Palazoglu, 2006). In this representation each block depicts a complete, unidirectional subsystem whose output variables are uniquely and completely determined by their input variables (Riggs, 1970).

It is well documented that the primary survival function of the osmotic stress response is to restore turgor pressure through increased synthesis and accumulation of glycerol (reviewed in (Hohmann et al., 2007)). Like the minimal representation presented by Muzzey et al., I start with measurement of turgor pressure, which is the controlled variable (y). By contrast with the minimal representation, in my detailed model turgor pressure is transduced into biologically meaningful signals through multiple known (and possibly additional unknown) measuring functions (g_m^i) , upstream signal transduction proteins with effects on turgor pressure adaptation (Hohmann, 2002; Hohmann et al., 2007; Rep et al., 1999). For example, the osmosensitive glycerol channel Fps1 closes in response to decreased turgor pressure. Activating conformational changes in the osmosensor proteins Sho1 and Sln1 respond to decreased turgor pressure to initiate the high osmolarity glycerol (HOG) pathway. And inactivating conformational changes in other proteins may respond to directly or indirectly to reduced turgor pressure or downstream changes (e.g. cytoplasmic crowding (Miermont et al., 2013)).

Mapping the minimal model of Muzzey et al. (Figure 2A) onto the detailed representation in Figure 1 shows 1) how their subsystems H and I violate single-input single-output (SISO)assumptions and are therefore invalid transfer functions, 2) where the subsystems overlap, and 3) how their minimal model is not isomorphic with the detailed model of the osmotic stress response including known features discussed in their article (Figure 2B; a minimal model consistent with the detailed block diagram of the response is given in Figure 2C). Moreover, glycerol concentration is by definition an integrator: the cumulative amount (glycerol per cell volume) is proportional to the integral of the rate of glycerol accumulation (amount per time). Glycerol is downstream of the integrator location suggested by Muzzey et al., who demonstrated that the osmotic stress response circuit in yeast must contain exactly one integrator acting in series (or possibly more than one integrator acting in parallel). Therefore, glycerol concentration and not the HOG pathway, must be the locus of integration.

At steady-state the concentration of glycerol is by definition constant. If the concentration of glycerol is constant, then net glycerol accumulation must be zero, and the rate of glycerol leakage must be equal to its rate of synthesis. As shown by Muzzey et al., similar logic applies to all upstream components; if the integrator is the most downstream element in the network (relative to the input of a disturbance in external osmolarity), all error deviations, Hog1 nuclear enrichment, and steady-state viability are expected to display perfect adaptation (Muzzey et al., 2009). Indeed, we observed perfect adaptation of viability before and after adaptation to an osmotic challenge (correlation between early mortality and recovery of viability of 50 different yeast strains is over 0.98; http://biorxiv.org/content/early/2016/03/07/039982). Given enough time (depending on the time constants of each response) cells adapt to the hyperosmolar media with increased intracellular glycerol concentrations. Once adaptation has occurred

and turgor pressure is restored, sensors relax and all deviations from steady state activity return to zero.

Discussion

The molecular mechanisms behind error sensing have been a source of mystery to bioengineers (e.g. "the molecular mechanisms behind error sensing are little understood" (Xiao et al., 2009)). However, protein conformational shifts between relaxed and activated states in response to changes in turgor pressure are easily interpretable as error sensing mechanisms in the osmotic stress response (Rutherford and Zuker, 1994). Conformational distortions caused by less than optimal turgor pressure are thought to activate the high osmolarity glycerol (HOG) pathway in proportion to osmotic stress. Similar activation or loss of activity of cytosolic proteins plausibly occur in conditions of molecular crowding (Miermont et al., 2013). In that case, restoration of cell volume and turgor pressure would allow proteins to return to pre-stress steady conformations and levels of activity. Finally, the intracellular concentration of any biomolecule would similarly be the integral of its positive rate of accumulation, providing a general mechanism for integral feedback and homeostatic actuators, converting the energy of cellular metabolism into the energy inherent in concentration gradients and other concentration dependent processes.

In addition to revealing potentially general features of integral feedback control in biology, the block diagrams in Figures 1 and 2c show how the osmotic stress response may work over different time scales and levels of stress. Upon a shift to hyperosmotic media, the initial response is the rapid closure of constitutive leakage channel Fps1, closing the smallest negative feedback loop in Figure 1 and effectively removing the negative input to the controller in the circuit shown in Figure 2c. Figure 1 also shows clearly and intuitively how the successive activation of sensors with longer time scales and additional sources of negative feedback on glycerol accumulation could occur, consistent with the observed longer delays and increasingly stronger adaptive responses proportional to the degree of osmotic stress (Hohmann et al., 2007).

Figure Legends

Figure 1. The yeast osmotic stress response as an error actuated linear control system. At steady-state (time 0⁻), turgor pressure (y) is at its normal, steady-state value and osmostress sensitive proteins (sensors 1-5) are in their relaxed, non-induced conformations, with all error deviations (e) equal to 0. An abrupt step in external

Figure 2. Minimal model violates single-input single-output (SISO) assumptions of conventional block diagrams. A. The minimal circuit model of Muzzey et. al. with 4 blocks identified as 1) the H subsystem that contains the MAP kinase cascade and "links an osmotic disturbance at the membrane with Hog1 nuclear enrichment", 2) the D subsystem that contains "Hog1 dependent mechanisms that promote glycerol accumulation including the transcriptional activation of genes encoding glycerol producing enzymes and interactions initiated by Hog1 in the cytoplasm or nucleus that lead to increased glycerol synthesis", 3) the I subsystem that contains "Hog-independent mechanisms that contribute to osmolyte production" and 4) the G subsystem representing "metabolic reactions involved in glycerol synthesis and any other reactions that contribute to glycerol accumulation" (Muzzey et al., 2009). A mathematical implementation of the model shows how turgor pressure can return to pre-stimulus values even in the continued presence of osmotic stress (Muzzey et al., 2009) but does

transfer functions are assumed (Riggs, 1970; Romagnoli and Palazoglu, 2006).

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Rutherford Figure 1

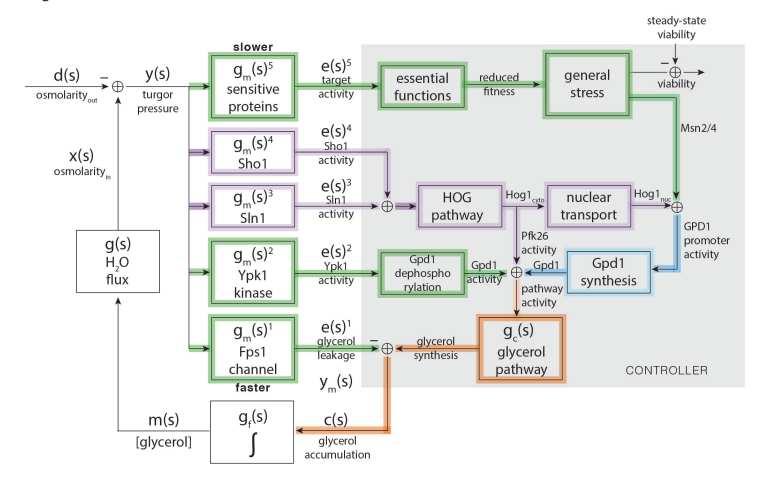


Figure 2

