A heritable switch in carbon source utilization driven by an unusual yeast prion

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Abstract

Several well-characterized fungal proteins act as prions, proteins capable of multiple conformations, each with different activities, at least one of which is self-propagating. Through such self-propagating changes in function, yeast prions act as protein-based elements of phenotypic inheritance. We report a prion that makes cells resistant to the glucose-associated repression of alternative carbon sources, [GAR⁺]. [GAR⁺] appears spontaneously at a high rate and is transmissible by non-Mendelian, cytoplasmic inheritance. Several lines of evidence suggest that the prion state involves a complex between a small fraction of the cellular complement of Pma1, the major plasma membrane proton pump, and Std1, a much lower abundance protein that participates in glucose signaling. The Pma1 proteins from closely related Saccharomyces species are also associated with the appearance of [GAR⁺]. This allowed us to confirm the relationship between Pma1, Std1, and [GAR⁺] by establishing that these proteins can create a transmission barrier for prion propagation and induction in S. cerevisiae. The fact that yeast cells employ a prion-based mechanism for heritably switching between distinct carbon source utilization strategies, and employ the plasma membrane proton pump to do so, expands the biological framework in which self-propagating protein-based elements of inheritance operate.
Introduction

The stable inheritance of biological information and phenotype across generations is a fundamental property of living systems. Prions, self-perpetuating and heritable protein conformations that cause multiple phenotypes, represent an unusual mechanism of information transfer that occurs via protein instead of nucleic acid (Wickner 1994). Prion proteins can assume at least two conformations and each conformation alters protein function, resulting in different phenotypes (Wickner et al. 2004; Shorter and Lindquist 2005). When in the self-templating, or prion conformation, prion proteins acquire characteristics normally restricted to nucleic acids. The first prion protein identified, the mammalian protein PrP, can behave as a transmissible pathogen and causes a neurodegenerative disease in its prion form (PrP\textsuperscript{Sc}) (Prusiner 1998). Prion proteins in fungi, which are functionally unrelated to PrP and to each other, act as non-Mendelian elements of inheritance by switching to the self-perpetuating, cytoplasmically transmissible prion conformation (Wickner 1994).

Four prions have been extensively characterized in fungi: [\textit{PSI}⁺], [\textit{URE3}], [\textit{Het-s}], and [\textit{RNQ}⁺]. [\textit{PSI}⁺] (Cox 1965) is the prion form of the translation termination factor Sup35, which causes nonsense suppression (Stansfield et al. 1995; Patino et al. 1996; Paushkin et al. 1996). [\textit{URE3}] (Lacroute 1971) is an altered form (Wickner 1994) of the nitrogen catabolite repressor Ure2 (Courchesne and Magasanik 1988). [\textit{RNQ}⁺] controls the ability of a cell to acquire other prions (Derkatch et al. 2000; Sondheimer and Lindquist 2000; Derkatch et al. 2001). [\textit{Het-s}], found in \textit{Podospora anserina}, causes heterokaryon incompatibility with certain mating partners (Rizet 1952; Coustou et al. 1997).
These four fungal prions, as well as several recently identified prions ([SWT], [MCA], [OCT⁺], and [MOI⁺]) (Du et al. 2008; Alberti et al. 2009; Nemecek et al. 2009; Patel et al. 2009), share key genetic and physical characteristics despite their disparate functions (Chien et al. 2004; Shorter and Lindquist 2005). Their phenotypes appear spontaneously at higher frequencies than those caused by genetic mutations. They are dominant, show non-Mendelian segregation following meiosis, and are also transmissible by cytoduction (cytoplasmic transfer). Physically, they form a self-templating amyloid conformation in the [PRION⁺] state. Further, their inheritance is linked to the activities of chaperones, proteins that mediate conformational changes in other proteins. Transient changes in chaperone levels, particularly Hsp104, are sufficient to eliminate the prions permanently from cells. This occurs because chaperones alter the prion conformations and transmission to daughter cells. Once the prion template is gone cells are “cured” of the elements (Uptain and Lindquist 2002; Shorter and Lindquist 2005). Another unusual feature is that transient overexpression of the prion protein causes permanent inheritance of the prion phenotype. This is because the protein:protein interactions involved in prion formation are more likely to occur at higher protein concentrations (Chernoff et al. 1993; Ter-Avanesyan et al. 1993; Wickner 1994; Serio et al. 2000; Sondheimer and Lindquist 2000; Derkatch et al. 2001; Uptain and Lindquist 2002; Shorter and Lindquist 2005). The yeast prions also share a distinctive feature with mammalian prions, a strong transmission barrier across species. Even subtle differences in amino acid sequence can reduce the ability of prion proteins from one species to convert the homolog from other species, even though the homologous protein is itself capable of forming a prion on its own (Aguzzi et al. 2007; Chen et al. 2007).
The precise nature of the mammalian prion template is not known, but all of the well characterized fungal prions, as well as the newly discovered prions and prion domains (Du et al. 2008; Alberti et al. 2009; Nemecek et al. 2009; Patel et al. 2009) are self-templating amyloid amyloids. The simple and robust character of self-templating amyloids provides a compelling framework for protein-based inheritance (Glover et al. 1997; Shorter and Lindquist 2005). Indeed, in many cases the amyloid has been shown to be the sole determinant needed for prion formation: recombinant amyloid fibers alone are sufficient to convert [prion] cells to [PRION] cells (Maddelein et al. 2002; King and Diaz-Avalos 2004; Tanaka et al. 2004; Brachmann et al. 2005; Patel and Liebman 2007; Alberti et al. 2009). Amyloid structure is therefore commonly held to be a critical feature of all naturally occurring systems for protein-based inheritance. Indeed, a recent genome-wide screen for new prion domains in yeast began by examining proteins likely to be amyloidogenic (Alberti et al. 2009).

Here we took a different approach. We searched the literature for S. cerevisiae phenotypes with prion-like inheritance patterns. One was described many years ago in a screen for cells with an alteration in carbon source utilization (Ball et al. 1976). The basis of the screen was the extreme preference of S. cerevisiae for glucose as a carbon source. In glucose media, cells repress genes necessary to process other carbon sources such as glycerol (Santangelo 2006). Glucosamine, a non-metabolizable glucose mimetic, induces a similar repression. Therefore, yeast cells cannot use glycerol as a carbon source if even small amounts of glucosamine are present (Hockney and Freeman 1980; Nevado and Heredia 1996). Some cells spontaneously acquire the ability to use glycerol in the presence of glucosamine, presumably due to defects in glucose repression. Some of these
Brown and Lindquist exhibit dominant, non-Mendelian inheritance (Ball et al. 1976). Further, the phenotype is neither carried by the mitochondrial genome nor by a plasmid (Kunz and Ball 1977). Employing a variety of methods, we show here that this factor, \([GAR^+]\), exhibits all of the genetic characteristics of a yeast prion, and we use a broad range of biochemical and genetic methods to identify proteins that play a key role in \([GAR^+]\) inheritance.

**Results**

\([GAR^+]\) shows non-Mendelian, infectious inheritance

We obtained cells able to utilize glycerol as a carbon source despite the presence of glucosamine, as did Ball and colleagues (Ball et al. 1976; Kunz and Ball 1977), by selecting for cells that could grow in 2% glycerol in the presence of 0.05% glucosamine. Colonies appeared at a frequency of approximately 5 in \(10^4\) cells in the W303 genetic background, well above the predicted mutational frequency (figure S01). Some recessive mutations allow growth on glycerol in the presence of glucosamine ((Ball et al. 1976) see table S1 table) but the novel phenotypes described by Ball and colleagues were dominant. Therefore, we first crossed our cells to wild-type cells. All diploids exhibited an unstable semi-dominant phenotype (figure 1a). Specifically, a mixed population was produced in which some diploids showed “strong” phenotypes (large colonies) and others “weak” phenotypes (small colonies; figure S02a). Cells with weak phenotypes invariably converted to strong over approximately 25 generations (data not shown). Notably, both mammalian and fungal prions exhibit “strong” and “weak” strains (Aguzzi et al. 2007).

In yeast, chromosomally inherited traits show 2 : 2 segregation following meiosis. Both strong and weak \([GAR^+]\) phenotypes, however, exhibited non-Mendelian 4 \([GAR^+]\):
non-Mendelian segregation (figure 1b). That is, all meiotic proeny exhibited a capacity to grow on glucose in the presence of glucosamine. Spores produced from cells with weak phenotypes generally converted to strong phenotypes (figure S02b, bottom). We named the responsible genetic element responsible for this trait [GAR\(^+\)], for "resistant to glucose-associated repression", with capital letters indicating dominance and brackets ([]) its non-Mendelian character.

To determine whether [GAR\(^+\)] is transmissible by cytoduction (that is, "infectious"), we used a mutant defective in nuclear fusion (kar1-1). During mating kar1 cells fuse but nuclei do not (Conde and Fink 1976). Selecting for a particular nucleus and cytoplasm of interest after mating accomplishes cytoplasmic exchange without the transfer of nuclear material. We mated a [GAR\(^+\)] strain carrying the nuclear markers URA3\(^+\) his3\(^-\) and the cytoplasmic marker \(\rho^+\) to a kar1-1 [gar\(^-\)] strain that was ura3\(^-\) HIS3\(^+\) and \(\rho^0\). We then selected for cells containing the nucleus originally associated with [gar\(^-\)] cells and the cytoplasm originally associated with [GAR\(^+\)] cells. All ten strains tested were [GAR\(^+\)] (figure 1c). Thus, [GAR\(^+\)] exhibits an “infectious,” non-nuclear pattern of inheritance.

\[\text{[GAR}^+]\text{ appears at high frequency in a variety of genetic backgrounds}\]

We next asked whether [GAR\(^+\)] was an oddity of specific strains or could appear in diverse genotypes. Cells able to utilize glycerol in the presence of glucosamine appeared at a frequency of ~9 in 10\(^5\) cells in the BY background, ~1 in 10\(^4\) cells in 74D, ~5 in 10\(^4\) cells in W303, and ~7 in 10\(^4\) cells in Sigma. In the SK1 background, [GAR\(^+\)] appeared at the astonishingly high rate of ~4 in 10\(^3\) cells (figure 1d). In comparison, the
frequency of heritable phenotypic change due to genetic mutation is generally ~1 in 10^6 haploid cells (Ohnishi et al. 2004).

We tested dozens of variants from each background for dominance. All exhibited the semi-dominant pattern observed in W303 (figure 1b and data not shown). \([GAR^+]\) cells of the 74D background did not sporulate, preventing us from testing segregation pattern. In W303 and W303/BY hybrids, \([GAR^+]\) only delayed sporulation (data not shown). In every tetrad tested from these backgrounds (more than 25 of each genotype), \([GAR^+]\) showed 4 \([GAR^+]\) to 0 \([gar^-]\) segregation (figure 1b and data not shown). Together, these data establish that yeast strains of diverse genetic backgrounds commonly switch carbon-utilization strategies in a heritable way by acquiring a non-Mendelian element of inheritance.

\([GAR^+]\) is curable by transient changes in chaperone protein levels

The inheritance of prions is based upon self-perpetuating changes in protein conformations. In contrast to other non-Mendelian elements, a hallmark of prion phenotypes is the ability of transient changes in the expression of chaperones to cause a heritable loss of the phenotype. Other yeast prions, as well as 18 of 19 newly identified protein domains with prion-forming capability, require Hsp104 for propagation (Chernoff et al. 1995; Derkatch et al. 1997; Moriyama et al. 2000; Shorter and Lindquist 2004; Jones and Tuite 2005; Du et al. 2008; Alberti et al. 2009; Patel et al. 2009). To test the influence of Hsp104 on \([GAR^+]\), we crossed \([GAR^+]\) cells to cells carrying a knockout of \(hsp104\) and sporulated them. Hsp104 was not required for \([GAR^+]\) inheritance: \(\Delta hsp104\) segregants remained \([GAR^+]\) (figure 1e). \([GAR^+]\) was also not curable by growth on
guanidinium hydrochloride, which inhibits Hsp104’s ATPase activity (Ferreira et al. 2001; Jung and Masison 2001), nor by overexpression of \textit{HSP104} (data not shown).

We next tested the Hsp70 proteins Ssa1 and Ssa2 (Werner-Washburne et al. 1987), mutations in which affect the inheritance of other prions (Sweeny and Shorter 2008). These mutations are also a good measure of general chaperone sensitivity, as they induce production of most chaperone proteins (Oka et al. 1997). Strikingly, all \textit{\Delta ssa1\Delta ssa2} meiotic products lost the ability to grow on glycerol in the presence of glucosamine (figure 1f). Was this due to curing of the \textit{[GAR]} genetic element or did the \textit{\Delta ssa1\Delta ssa2} mutations simply mask the phenotype? To test this we restored \textit{SSA1} and \textit{SSA2} to the glucosamine-sensitive \textit{\Delta ssa1\Delta ssa2} progeny by mating them back to wild-type \textit{[gar]} cells (see figure S03a for diagram of cross). Restoring Hsp70 function did not result in the reappearance of the \textit{[GAR]} phenotype (data not shown). However, when the cells were plated on medium with glucosamine, colonies able to grow on glycerol could be recovered at normal frequencies (figure S03b). Thus, a transient change in chaperone proteins was sufficient to cure cells of \textit{[GAR]} and this curing was reversible, both hallmarks of prion biology (Wickner 1994). \textit{[GAR]} therefore exhibits all of the distinguishing genetic characteristics of yeast prions.

\textit{[GAR]} is regulated by the Rgt2/Snf3 glucose signaling pathway

We performed gene expression profiling to identify transcriptional consequences of \textit{[GAR]}. In glucose-grown cultures tested just prior to the diauxic shift, only one gene showed a detectable difference between \textit{[gar]} cells and \textit{[GAR]} cells on our arrays, but that gene was very strongly affected. \texttt{Hexose Transporter 3} (\texttt{HXT3}) was approximately
36-fold down-regulated in \([GAR^+]\) cells compared to \([gar]\) cells (figure S04). No other transcript exhibited more than a two-fold change. We used an Hxt3-GFP fusion protein under the control of the endogenous \(HXT3\) promoter to examine protein levels. Hxt3-GFP was easily visible at the plasma membrane in late log phase \([gar]\) cells but extremely difficult to detect in \([GAR^+]\) cells (figure 2a). The loss of \(HXT3\) expression (\(\Delta hxt3\)) alone did not allow cells to utilize glycerol in the presence of glucosamine (figure 2b) and thus, does not explain the \([GAR^+]\) phenotype. However, it led us to hypothesize that the causal agent of \([GAR^+]\) is a regulator of \(HXT3\) expression.

To define the protein(s) required for \([GAR^+]\) inheritance, we took advantage of two things. First, transient overexpression of each of the known prion proteins dramatically increases the frequency at which the corresponding prion appears (Uptain and Lindquist 2002). Second, the \([GAR^+]\) determinant exerts a strong effect on \(HXT3\) expression, and \(HXT3\) predominantly controlled by the Snf3/Rgt2 pathway (Kim et al. 2003; Santangelo 2006). When glucose is present, transmembrane glucose sensors Snf3 and Rgt2 transmit a signal to the Yck1 and Yck2 complex, which then phosphorylates Mth1 and Std1, marking them for degradation (figure 2c) (Moriya and Johnston 2004). When glucose is not present, Mth1 and Std1 accumulate and interact with Rgt1. This complex then binds to the \(HXT3\) promoter and represses transcription of \(HXT3\) (Lakshmanan et al. 2003).

We tested each gene in the Snf3/Rgt2 regulatory pathway for induction of \([GAR^+]\) when overexpressed from a plasmid with a strong constitutive promoter, GPD (figure 2d). In every strain test, \(STD1\) caused an extraordinary increase in the appearance of colonies able to grow on glycerol in the presence of glucosamine. In W303, for example,
the increase was ~900 fold over empty vector; more than one in ten cells in these cultures converted to $[\text{GAR}^+]$. This is at the high end of prion inductions obtained by analogous experiments with other proteins (Masison and Wickner 1995; Derkatch et al. 1996). While no other gene in this pathway induced $[\text{GAR}^+]$, overexpression of the STD1 paralog $\text{MTH1}$ blocked its appearance, further confirming the importance of members of this pathway in $[\text{GAR}^+]$ biology.

**Transient STD1 overexpression induces $[\text{GAR}^+]$ but is not required for maintenance**

Next we asked if transient expression of STD1 was sufficient to create a heritable change in phenotype, a defining feature of prion biology. When ~100 cells that had lost the over-expression plasmid were tested, all retained the $[\text{GAR}^+]$ phenotype (confirmed by marker loss; data not shown). Thus $\text{STD1}$ is not simply a dynamic regulator of glucose repression. Rather, its transient overexpression induces a new, heritable state of carbon utilization.

These data suggested that Std1 is the determinant of the $[\text{GAR}^+]$ prion, but further date indicated it could not be the sole determinant. First, most prion phenotypes mimic loss-of-function phenotypes of their prion determinants. However, $\Delta\text{std1}$ strains derived from a $[\text{gar}^-]$ background were not able to grow on glycerol in the presence of glucosamine (figure 2b and data not shown). Further, $\Delta\text{std1}$ cells derived from a $[\text{GAR}^+]$ background were able to do so, indicating that they kept the prion (data not shown). Finally, such cells were able to pass the [GAR] element onto progeny in tester crosses for inheritance of the prion element (figure 2e). Therefore, $[\text{GAR}^+]$ maintenance does not
require STD1. This makes [\(GAR^+\)] highly unusual among yeast prions in that its transient inducing agent is not required for propagation.

We next examined all other members of the Rgt2/Snf3 pathway. None behaved as would be expected for the causal agent of [\(GAR^+\)]. All knockouts were capable of propagating [\(GAR^+\)] (figure S05). Cells with \(rgt1\) knockouts did not exhibit the prion phenotype, but they maintained it in a “cryptic” form. It reappeared when cells were crossed to \([\text{gar}^-]\) \(RGT1\) cells. Therefore, \(RGT1\) is required for the manifestation of [\(GAR^+\)] phenotype but is not necessary for its propagation.

**Identification of genes that modify the frequency of [\(GAR^+\)] appearance**

We conducted genome-wide screens for affecters of [\(GAR^+\)] induction. We screened the \(S.\ cerevisiae\) haploid deletion library (Giaever et al. 2002) for mutants that were incapable of inducing [\(GAR^+\)] (table S2), caused a high frequency of appearance of [\(GAR^+\)] (table S3), or that themselves exhibited an ability to grow on glycerol in the presence of glucosamine (table S1). Four of the eight members of the Snf3/Rgt2 pathway showed a phenotype in this screen (\(p = 8 \times 10^{-6}\); Fisher’s exact test). \(\Delta snf3\) grows on glycerol with glucosamine (table S1) and \(\Delta std1, \Delta mth1,\) and \(\Delta rgt1\) exhibited lower than normal [\(GAR^+\)] induction (figure 2b. table S2). However, none of these genes were required for the maintenance of [\(GAR^+\)] in strains already carrying the element (figure S05).

Finally, we screened a library of \(~5000\) ORFs (\(~85\%\) of yeast ORFs) on a galactose-inducible single copy plasmid (Leonardo et al. 2002) to find genes that induce [\(GAR^+\)] following overexpression. \(STD1\) was the only clone that caused strong [\(GAR^+\)]
induction, ~1000 fold when retested under the regulation of the \textit{GPD} promoter. A second gene, \textit{DOG2}, caused a 10-fold induction (figure S06).

\textit{Pma1 associates with Std1 and is a component of [GAR\textsuperscript{+}]}

Since neither the deletion nor the overexpression screen identified a protein that by itself could embody the [GAR\textsuperscript{+}] prion, we turned to biochemical methods. \textit{STD1} had been implicated in [GAR\textsuperscript{+}] in three ways: 1) the highly specific down-regulation of \textit{HXT3} pointed to members of the Rgt2/Snf3 glucose signaling pathway; 2) transient \textit{STD1} overexpression caused huge increases in [GAR\textsuperscript{+}] appearance; and 3) deletion of \textit{std1} reduced the spontaneous appearance of [GAR\textsuperscript{+}] to the frequency of genetic mutations. We hypothesized, therefore, that Std1 might physically interact with an unknown propagating agent.

We sought proteins that interacted with Std1 by co-immunoprecipitation with an HA-tagged derivative. A high molecular weight band was recovered from [GAR\textsuperscript{+}] protein lysates but not from [\textit{gar}] lysates (figure S07). Mass spectrometry analysis identified the protein as Pma1, a large, highly abundant P-type ATPase with 10 transmembrane domains that is the major controller of membrane potential and cytoplasmic pH (Morsomme et al. 2000). When the same assay was performed with isogenic \textit{\Delta std1} cells, Pma1 was not detected. Notably, if Pma1 is indeed a constituent of the prion, we would not have identified it in our genetic screens. It is essential (Serrano et al. 1986) and therefore absent from the deletion library. Moreover, it is already the most abundant membrane protein in yeast and notoriously difficult to overexpress (Eraso et al. 1987).
Transient overexpression of *STD1* induced [\(GAR^+\)] and transient overexpression of its paralog, *MTH1*, inhibited [\(GAR^+\)] conversion. We therefore asked whether Pma1 exhibited heritable differences in association with Std1 and Mth1 in [\(gar^-\)] and [\(GAR^+\)] cells. As a multipass transmembrane protein, Pma1 is intractable to most methods for analyzing protein complexes, but it migrates as an oligomeric species when digitonin lysates are separated on Blue Native gels (Gaigg et al. 2005). Most Pma1 in [\(GAR^+\)] and [\(gar^-\)] cells migrated as heterogenous high molecular weight (HMW) complexes but a smaller fraction migrated as two distinct complexes of (very roughly) 600 and 700kDa (figure 3a, top). The lower bands (especially the 600kDa species) were associated with Std1 in [\(GAR^+\)] cells but with Mth1 in [\(gar^-\)] cells (figure 3a, bottom). Std1 is much less abundant than Pma1. Consistent with the fact that only a small fraction of Pma1 is associated with Std1 in [\(GAR^+\)] cells, Pma1 showed a minor but statistically significant but minor change in protease sensitivity between [\(gar^-\)] and [\(GAR^+\)] cells (figure S08).

Next we asked whether mutations that affect Pma1 oligomerization and trafficking to the plasma membrane alter [\(GAR^+\)] frequency. Mutants that affect phospholipid synthesis and protein trafficking but not Pma1 oligomerization – *LCB3, LCB4, DPL1* and *ATG19* (Lee et al. 2002; Mazon et al. 2007) – did not change the appearance of [\(GAR^+\)] (figure 3b and S09a). Mutants that do affect Pma1 oligomerization and trafficking — *SUR4* and *LST1* (Roberg et al. 1999; Lee et al. 2002) decreased the appearance of [\(GAR^+\)] (figure 3b and S09a). These genes were not, however, required for [\(GAR^+\)] maintenance (figure S09b).

We explored the relationship between Pma1, [\(GAR^+\)], and the Rgt2/Snf3 glucose signaling pathway. Carbon sources regulate Pma1's phosphorylation state (Lecchi et al. 2005).
2005), its ATPase activity (Serrano 1983), and its conformation (Miranda et al. 2002) through residues S899, S911, and T912 in the C-terminal tail, which faces the cytosol (Eraso et al. 2006; Lecchi et al. 2007). We mutated S899, S911, and T912 to alanine, which cannot be phosphorylated, or to aspartic acid, which mimics constitutive phosphorylation. (Phosphorylated S911 and T912 are commonly observed in glucose media and the non-phosphorylated forms when cells are starved of glucose (Lecchi et al. 2007)) S899 mutations and S911D and/or T912D mutations had no effect on \([GAR^+]\) frequency. However, S911A and S911A/T912A increased the frequency of \([GAR^+]\) appearance by several fold (figure 3c). Notably, these same mutants also reduced levels of an Hxt3-GFP reporter, both a readout for the Rgt2/Snf3 pathway and the only change in gene expression detected in \([GAR^+]\) cells (figure 3d). These results indicate that Pma1 affects glucose signaling to regulate \(HXT3\). In any case, the fact that such subtle mutations in the Pma1 protein affect \([GAR^+]\) induction confirms that Pma1 plays a key role in \([GAR^+]\) biology.

The unstructured N-terminus of Pma1 is involved in \([GAR^+]\) propagation

A characteristic of prions is that transient overexpression is sufficient for induction. However, Pma1 is the most abundant plasma membrane protein in yeast (Morsomme et al. 2000) and overexpression is not well tolerated (Eraso et al. 1987). We found that we could obtain a three-fold increase in Pma1 protein levels with a \(CEN\) plasmid and a \(GPD\) promoter. This caused a corresponding increase in \([GAR^+]\) frequency (figure 4a). Introducing stop codons at amino acid positions 23 or 59 eliminated this effect (figure S10). Thus, it is not the nucleic acid sequence but the Pma1 protein that
contributes to \([GAR^+]\) induction. Finally, when the inducing \(GPD\ PMA1\) plasmid was lost, the cells remained \([GAR^+]\). Thus a transient increase in \(PMA1\) was sufficient to induce a heritable change in phenotype.

Pma1's N- and C-termini face the cytosol. The C-terminus is predicted to be \(\alpha\)-helical and the N-terminus unstructured (Morsomme et al. 2000), the latter a characteristic of prions. An N-terminally truncated (\(\Delta40\)) mutant of \(PMA1\) did not increase \([GAR^+]\) appearance, although the protein was expressed at wild-type levels, (figure 4a). A C-terminally truncated \(PMA1\) did increase \([GAR^+]\) induction, even though its levels were reduced.

\([GAR^+]\) could be propagated through cells whose only source of Pma1 was a \(GAL1\)-regulated N-terminal deletion, \(PMA1\Delta40N\) (figure S11). Strikingly, however, it did not propagate through a double mutant of \(PMA1\Delta40N\) and \(\Delta std1\), and it did not reappear when wild-type \(PMA1\) and \(STD1\) function were restored with crosses (figure 4b). (The few glucosamine-resistant colonies that remained were not \([GAR^+]\) but contained conventional recessive; data not shown). Thus, once \([GAR^+]\) has been established, it is maintained in the absence of either Std1 or the N-terminus of Pma1, but not in the absence of both.

\([GAR^+]\) is sensitive to a Pma1-dependent “species barrier”

Previously described yeast prion proteins exhibit changes in localization and solubility in the prion state (Uptain and Lindquist 2002) and affect the induction of other prions by cross templating (Derkatch et al. 2000; Derkatch et al. 2001). There was no difference in localization of Pma1 or Std1 between \([gar^+]\) and \([GAR^+]\) (figure S12).
Neither formed a detectable SDS-resistant species in \([GAR^+]\) (figure S13). Furthermore, the frequency of \([GAR^+]\) appearance did not change in backgrounds carrying \([PSI^+]\), \([RNQ^+]\), or \([URE3]\), prions that broadly affect the appearance of amyloid-based prions (figure S14). Analysis of protein extracts by 2D gel electrophoresis did not reveal any proteins that changed solubility between \(\text{gar}^-\) and \([GAR^+]\) (figure S15). \([GAR^+]\) was not affected by Hsp104 expression (figure 1e). Whatever the manner by which Pma1 and Std1 contribute to the prion state, it is not likely by forming amyloid.

The extremely stable nature of amyloids allows them to be confirmed as prion determinants by "protein only" transformation (Maddelein et al. 2002; Tanaka et al. 2004). The lack of an identifiable amyloid in \([GAR^+]\) cells precluded the use of this procedure for \([GAR^+]\). Instead, to rigorously test the relation between Pma1, Std1, and \([GAR^+]\), we performed a classic “transmission barrier” experiment. Small differences in amino acid sequence cause prions that originate in one species to fail in transmission to another (Santoso et al. 2000; Bagriantsev and Liebman 2004; Chen et al. 2007). If Pma1 and Std1 contribute to a transmission barrier for \([GAR^+]\), it would establish that they are integral to the propagating element.

We chose to study a possible \([GAR^+]\) transmission barrier using \(S.\) bayanus and \(S.\) paradoxus, two closely related \textit{sensu stricto} species that also exhibit glucose-mediated repression of the utilization of other carbon sources. First, we asked whether diploids of these species could also acquire the ability to utilize glycerol in the presence of glucosamine (figure 5a). They could, and they did so at a higher frequency than expected for mutation. Indeed, \([GAR^+]\) appeared in \(S.\) bayanus at an astonishingly high rate (greater than one in 1,000 cells). Moreover, the \([GAR^+]\) phenotype was very stable in
these cells. Thus, the ability to heritably switch carbon utilization strategies through this prion is broadly utilized.

We asked whether the Pma1 proteins from *S. bayanus* and *S. paradoxus* can propagate \([GAR^+]\) in *S. cerevisiae*. Sequence differences between the species are slight (figure S16): *S. bayanus* Pma1 and *S. paradoxus* Pma1 are 96% and 99% identical to *S. cerevisiae* Pma1, respectively. Most of these changes are in the N-terminal region, which is required for prion induction.

First, we transformed *S. bayanus* or *S. paradoxus* *PMA1* plasmids into an *S. cerevisiae* strain in which a deletion of the essential *PMA1* gene was covered by a plasmid encoding *S. cerevisiae* Pma1. The *S. cerevisiae* *PMA1* plasmid was then selected against. All cells grew at the same rate on glucose, indicating that the Pma1 protein from these species was fully functional in *S. cerevisiae*. However, when \([GAR^+]\) cells were selected by plating these cells to glycerol-glucosamine medium, the resultant phenotypes were weak, unstable, and appeared at a low frequency. When putative \([GAR^+]\) cells were passaged on non-selective medium and then plated back onto glucosamine-containing medium, many fewer cells with *S. bayanus* or *S. paradoxus* *PMA1* maintained the resistant phenotype than cells with *S. cerevisiae* *PMA1* (data not shown). Thus, in a background where the entire genome otherwise remains the same, changing the species of origin for Pma1 had a critical effect on \([GAR^+]\) induction and propagation.

Next we asked whether the *S. bayanus* or *S. paradoxus* Pma1 proteins could propagate a \([GAR^+]\) state received from the *S. cerevisiae* protein. We performed another plasmid shuffle, this time starting with cells already carrying a strong *S. cerevisiae* \([GAR^+]\) element. We selected against the plasmid carrying the *S. cerevisiae* *PMA1* after
approximately 25 generations. After another 25 generations, cells were tested for the ability to grow on glycerol in the presence of glucosamine. Most retained a strong \([GAR^+]\) phenotype. Thus, strains with \textit{S. bayanus} and \textit{S. paradoxus} \textit{PMA1} were capable of accepting and propagating \([GAR^+]\) from strains with \textit{S. cerevisiae} \textit{PMA1} (figure 5b), at least after co-expression of both proteins for 25 generations.

Finally we tested how efficiently \([GAR^+]\) elements from cells expressing \textit{S. bayanus} or \textit{S. paradoxus} \textit{PMA1} could be transmitted back to cells expressing only \textit{S. cerevisiae} \textit{PMA1}. Multiple \([GAR^+]\) strains carrying the three \textit{PMA1} genes were mated to wild-type \([\text{gar}^-]\) cells. Cells expressing \textit{PMA1} from \textit{S. paradoxus} could not transmit \([GAR^+]\) at all, and cells expressing \textit{PMA1} from \textit{S. bayanus} transmitted it very inefficiently. Controls expressing \textit{S. cerevisiae} \textit{PMA1} transmitted \([GAR^+]\) efficiently (figure 5b). Thus, the \textit{PMA1} species of origin creates a strong transmission barrier for \([GAR^+]\) propagation.

Might Std1, the \([GAR^+]\) induction factor that is complexed with Pma1 in \([GAR^+]\) cells, create an induction barrier? Std1 is 81% identical between \textit{S. cerevisiae} and \textit{S. bayanus} but much more divergent in \textit{S. paradoxus} (figure S17). We transiently overexpressed \textit{STD1} from each organism in \([\text{gar}^-]\) \textit{S. cerevisiae} cells carrying each of the three Pma1 genes. \textit{STD1} alleles of \textit{S. cerevisiae} and \textit{S. bayanus} acted as general inducers. They increased the appearance of \([GAR^+]\) ~1000 fold in strains producing the Pma1 protein of any of the three species (figure 5c). In contrast, \textit{S. paradoxus} \textit{STD1} did not induce \([GAR^+]\) in any. Presumably, some other factor contributes to \([GAR^+]\) induction in \textit{S. paradoxus}. Most importantly, however, this experiment demonstrates that Std1 creates
a strong species barrier for $[GAR^+]$ induction, confirming its intimate involvement in the prion.

**Discussion**

The ability of cells to sense and adapt to nutrients is crucial to survival in highly competitive and rapidly fluctuating environments. Here we describe a cytoplasmically inherited element, $[GAR^+]$, that is involved in the fundamental processes of glucose sensing and signaling and carbon source utilization. $[GAR^+]$ arises spontaneously in every *S. cerevisiae* strain tested as well as sibling species separated by ~5 million years of evolution (Kellis et al. 2004), *S. paradoxus* and *S. bayanus*, at frequencies much higher than genetic mutations.

$[GAR^+]$ fulfills all of the genetic criteria established for prions: it is dominant (or at least semi-dominant). It exhibits non-Mendelian inheritance. It can be transferred via cytoplasmic exchange. Transient changes in the levels of chaperone proteins are sufficient to heritably cure cells of the $[GAR^+]$ state. Transient changes in the expression of proteinaceous determinants heritably induce $[GAR^+]$. The non-Mendelian mechanism of inheritance that best describes $[GAR^+]$ is that of a prion.

In other ways, however, $[GAR^+]$ seems very different from previously described yeast prions. It has at least two components, the plasma membrane proton pump Pma1 and the glucose signaling factor Std1. Transient overexpression of either *PMA1* or *STD1* is sufficient to establish a heritable conversion to $[GAR^+]$, yet once $[GAR^+]$ is established, either is sufficient for propagation. Cells lacking *std1* and also carrying a small deletion in the N-terminus of Pma1 cannot propagate $[GAR^+]$ at all. Pma1 and Std1 associate in an
oligomeric complex in \([GAR^+]\) cells but this complex is barely detectable in \([gar]\) cells. The integral relationship between these proteins and the \([GAR^+]\) state was tested and confirmed by transmission barrier experiments. Substituting the \(PMA1\) gene from \(S. bayanus\) or \(S. paradoxus\) for that of \(S. cerevisiae\) blocked propagation of \([GAR^+]\) to \(S. cerevisiae\) Pma1. Substituting Std1 from \(S. paradoxus\) eliminated its potency in \([GAR^+]\) induction.

What, then, is the nature of \([GAR^+]\)? It does not involve a detectable amyloid form, at least of the Pma1 or Std1 proteins. It is also not sensitive to overexpression or deletion of the general amyloid-remodeling protein Hsp104. Hsp104 severs amyloid filaments to ensure orderly inheritance of prion templates to daughter cells. It is required for the propagation of all known prions as well as for 18 of 19 recently discovered prion candidates (Chernoff et al. 1995; Patino et al. 1996; Derkatch et al. 1997; Ness et al. 2002; Cox et al. 2003; Kryndushkin et al. 2003; Shorter and Lindquist 2004; Jones and Tuite 2005; Shorter and Lindquist 2006; Tipton et al. 2008; Alberti et al. 2009). Thus, the absence of dependence on Hsp104 makes it rather unlikely that \([GAR^+]\) involves any amyloid-based element.

One possibility is that \([GAR^+]\) inheritance and propagation results from heritable alterations in Rgt2/Snf3 signaling involving a self-sustaining feedback loop. Indeed, Std1 and its paralog, Mth1, are subject to many feedback mechanisms involving their own transcription and degradation (Lakshmanan et al. 2003; Moriya and Johnston 2004; Polish et al. 2005; Kim et al. 2006), and Std1 is found both in the nucleus and on the plasma membrane (Schmidt et al. 1999). Furthermore, Pma1 is very abundant and Std1 is extremely scare (Morsomme et al. 2000). Our data suggest that only a small fraction of
Pma1 contributes to \([GAR^+]\) and that Std1 is the limiting factor. This would be consistent with altered signaling, as only small amounts of the Std1 protein would be necessary to shift the activity of a fraction of Pma1. However, if \([GAR^+]\) is simply due to altered signaling, the mechanism that maintains it must be remarkably robust, as it has been maintained in a highly stable state in some of our strains for six years now, with repeated dilutions into log phase, storage in the freezer and refrigeration, transitions back to room temperature, growth in liquid and on plates, in a wide variety of different media, through repeated rounds of growth into stationary phase (wherein most aspects of carbon metabolism undergo profound changes), and through starvation-induced meiosis.

Another possibility is that \([GAR^+]\) starts with a change in the association of Std1 and Pma1 that induces a conformational change in oligomeric species of each. These can then be maintained in the absence of either Std1 or the Pma1 N-terminus, but not in the absence of both (figure 6). We do not exclude the possibility that another protein contributes to the \([GAR^+]\) state. Indeed, our observations that *S. paradoxus* acquires \([GAR^+]\) at a high frequency, but that the Pma1 and Std1 proteins of *S. paradoxus* do not reconstitute \([GAR^+]\) in *S. cerevisiae*, suggesting the involvement of another protein. (This protein might even form amyloid, but if so it does not require Hsp104 and has escaped detection in our genetic screens.)

Of course, models involving self-perpetuating signaling loops and conformational changes are not mutually exclusive. Associations between Pma1 and Std1 might result in a conformational change that alters signaling and sets up a robust feedback loop that helps maintain the association, either between those same molecules of Pma1 and Std1 or
between other molecules and these proteins (figure 6). It will be of great interest to determine what might render such states stable enough to be so robustly heritable.

Another remaining question is the precise reason why cells carrying \([\text{GAR}^{+}]\) are able to grow on glycerol in the presence of glucosamine. We hypothesize that the \([\text{GAR}^{+}]\) phenotype involves altered signaling through a glucose sensing pathway, likely through Std1’s previously reported ability to interact with the DNA binding protein Rgt1 (Lakshmanan et al. 2003) (figure 6). Experiments investigating gene expression patterns over a much broader range of carbon sources and time points than examined here, as well as chromatin immunoprecipitation experiments with Std1 and Rgt1, may prove illuminating.

Whatever the mechanism may prove to be, Pma1, the major plasma membrane ATPase, and Std1, a much rarer and poorly understood signaling protein, contribute to a prion-like phenotypic state that heritably alters fundamental decisions about carbon source utilization. This heritable element, \([\text{GAR}^{+}]\), has all of the definitive characteristics of a prion. It has been stated that prion-mediated epigenetic states are simply diseases of yeast (Nakayashiki et al. 2005). Our findings that such an element controls something as fundamental to yeast biology as glucose repression, and that this element spontaneously arises at high frequency in diverse strains and sibling species, suggests that such epigenetic switches are actually integral to yeast biology. Clearly, self-propagating protein-based elements (prions) that can stably perpetuate biological states across generations operate over a much broader mechanistic landscape than previously supposed.
Materials and Methods

Yeast strains and genetic manipulations

Strain construction and manipulation followed standard yeast techniques. A list of strains and plasmids used in this study is available in tables S1 and S2. Unless otherwise stated, data shown is from genetic background W303. Five-fold dilutions were used for all spotting assays. Media used were yeast peptone-based medium containing the designated carbon source (YPD, YPglycerol, YPgalactose), synthetic medium lacking a particular amino acid (SD), or glycerol glucosamine medium (GGM; 1% yeast extract, 2% peptone, 2% glycerol, 0.05% D-(+)-glucosamine [Sigma G4875]).

[GAR⁺] frequency assays and isolation of [GAR⁺]

Cultures for [GAR⁺] frequency assays were grown overnight in 2% glucose, either YPD or SD, subcultured in the same, then grown to early exponential phase (OD₆₀₀ = 0.2-0.4). Cultures plated straight to GGM and diluted 10⁴ for plating to YPD. To isolate [GAR⁺] for further study, colonies from GGM were restreaked once to GGM then used in downstream applications. Unless otherwise stated, error bars in [GAR⁺] frequency assays represent the standard deviation and p-values are the binomial distribution of the mean. In all assays for [GAR⁺] propagation, cells were passaged for >100 generations before testing for growth on glycerol in the presence of glucosamine. Sporulation was performed by growing to diauxic shift in YPD or SD, plating to sporulation plates (1% potassium acetate, 0.05% dextrose, 0.1% yeast extract, and 0.01% complete amino acid mix [Bio101]), and incubating at 23°C until sporulated.
Genetic, biochemical, and cell biological analysis

Gene expression profiling, Western blotting, immunoprecipitation, fluorescent microscopy, Blue Native gel analysis, protease sensitivity analysis, and genetic screens were all performed using standard procedures. Detailed descriptions are available in the Supplementary Materials and Methods.

Acknowledgements

We would like to thank and acknowledge Neal Sondheimer, whose initial investigation first suggested that \( [GAR^+] \) might be a prion-based phenotype (Sondheimer 2000). We thank Amy Chang for gifts of an antibody and strains, and general advice on working with Pma1; Tom Rapaport for a Sec61 antibody; Eric Spooner for mass spectrometry; and Whitehead Institute Center for Microarray Technology performed cDNA synthesis, labeling, and hybridization reactions for the microarrays. We thank members of the Lindquist lab for providing comments and advice. This work was supported by the Mathers Foundation (to S.L.), Du-Pont MIT Alliance (to S.L.) and N.I.H. grant GM25874. J.C.S.B is supported by an NSF Graduate Research Fellowship and the Arthur Siegel Fellowship from W.I.B.R.
References


Figure Legends

Figure 1: \([\textit{GAR}^+]\) shares the genetic characteristics of yeast prions

a) Mating of \([\textit{gar}^-]\) \textit{MATa} to \([\textit{GAR}^+]\) \textit{MAT\alpha} in the W303 background. Resultant diploids show semi-dominant \([\textit{GAR}^+]\) with a mixed population of large colonies (“strong”) and small colonies (“weak”). All spot tests shown are five-fold dilutions. Diploids are selected for prior to plating to ensure that they are a pure population. b) Tetrad spores from the “strong” \([\textit{GAR}^+]\). Diploids in part A show non-Mendelian segregation of \([\textit{GAR}^+]\). c) Cytoduction shows cytoplasmic inheritance of \([\textit{GAR}^+]\). The \([\textit{GAR}^+]\) donor is 10B \(\textit{URA3}^+\ \textit{his3}^+ \textit{kar1}-1\) and the acceptor is W303 \(\textit{ura3}^-\ \textit{HIS3}^+ \textit{p}^0\) \textit{KAR1}. The \([\textit{GAR}^+]\) donor is therefore capable of growing on glycerol but the \([\textit{gar}^-]\) acceptor is not; “mixed” cells were selected for growth on glycerol (\([\textit{GAR}^+]\) cytoplasm) and SD-his 5-FOA (\([\textit{gar}^-]\) nucleus and counter-selection against the \([\textit{GAR}^+]\) nucleus). d) \([\textit{GAR}^+]\) frequency in various lab strains. Data are shown as mean +/- standard deviation (n=6). e) Tetrad spores from a \([\textit{GAR}^+]\) diploid with the genotype \(\textit{hsp104::LEU2/HSP104}\). \(\Delta\textit{hsp104}\) spores are still \([\textit{GAR}^+]\). f) Tetrad spores from a \([\textit{GAR}^+]\) diploid with the genotype \(\textit{ssa1::HIS3/SSA1 ssa2::LEU2/SSA2}\). \(\Delta\textit{ssa1}\Delta\textit{ssa2}\) spores are no longer \([\textit{GAR}^+]\).

Figure 2: The Snf3/Rgt2 glucose signaling pathway affects \([\textit{GAR}^+]\)

a) Hxt3-GFP signal in \([\textit{gar}^-]\) and \([\textit{GAR}^+]\) cells (S288c background) by fluorescence microscopy. b) Frequency of \([\textit{GAR}^+]\) in knockouts of members the Snf3/Rgt2 glucose signaling pathway. \(\Delta\textit{snf3}\) is completely resistant to glucosamine and therefore \([\textit{GAR}^+]\) frequency could not be measured. Furthermore, the frequency of spontaneous glucosamine-resistant colonies in the \(\Delta\textit{rgt1}, \Delta\textit{std1}, \) and \(\Delta\textit{mths1}\) strains was close to the
rate of genetic mutation and therefore these colonies might not carry the actual \([\text{GAR}^+]\) element. Overall, this pathway is enriched for genes that alter \([\text{GAR}^+]\) frequency when knocked out relative to the library of nonessential genes \( (p = 8 \times 10^{-6}, \text{Fisher’s exact test}) \).

c) The Snf3/Rgt2 glucose signaling pathway (adapted from (Moriya and Johnston 2004)).

d) Measurement of \([\text{GAR}^+]\) frequency following overexpression of Snf3/Rgt2 pathway members. Data are shown as mean +/- standard deviation \((n=6)\). \(STD1\) strongly induces conversion to \([\text{GAR}^+]\) and \(MTH1\) blocks it.

e) Top: tetrad spores from a \([\text{GAR}^+]\) diploid with the genotype \(std1::\text{kanMX}/STD1\). Bottom: spores from top crossed to a \([\text{gar}^-]\) strain with a wild-type \(STD1\) allele.

Figure 3: Pma1 is involved in \([\text{GAR}^+]\)

a) Native gel of Pma1, Std1, and Mth1 in \([\text{gar}^-]\) and \([\text{GAR}^+]\). Either Std1 (left) or Mth1 (right) was tagged with six tandem HA tags and samples were processed as described below from \([\text{gar}^-]\) and \([\text{GAR}^+]\) strains of each background. Total, supernatant (sup.), digitonin soluble (det. sol.), and digitonin insoluble (insol.) fractions were run on SDS gels and probed for Pma1 and Std1 or Mth1 (lower right) as a fractionation control. No differences in Pma1, Std1, or Mth1 levels or localization were detected between \([\text{gar}^-]\) and \([\text{GAR}^+]\). Blots of the total fraction were stained with Ponceau Red to confirm equal amounts of starting material (top right).

b) Measurement of \([\text{GAR}^+]\) frequency in knockout mutants of genes previously shown to affect \(\Delta\text{sur4}, \Delta\text{lst1}\) (Eisenkolb et al., 2002) \((\text{Roberg et al., 1999})\) or not affect \(\Delta\text{lcb3}, \Delta\text{lcb4}, \Delta\text{dpl1}, \Delta\text{atg19}\) (Gaigg et al., 2005) \((\text{Mazon et al., 2007})\) attributes of wild-type Pma1. Graph represents the mean +/- standard deviation \((n=6)\).

c) Mutants in phosphorylation sites at the C-terminus of Pma1
affect $[\text{GAR}^+]$ frequency. Starting strain is haploid, $[\text{gar}^-]$, genotype $\text{pma1}::\text{kanMX}$ with p316-PMA1. p314-PMA1 carrying wild-type $\text{PMA1}$ or mutants of interest were transformed in and then p316-PMA1 plasmid selected against by growth on 5-FOA. Graph represents the mean +/- standard deviation (n=6). P-values are the binomial distribution of the mean. d) Pma1 mutants that increase $[\text{GAR}^+]$ frequency show decreased levels of Hxt3-GFP. Graph represents the mean +/- standard deviation (n=6) and p-values were determined using the chi-squared test. Strain background is a hybrid of W303 and S288C.

Figure 4: Alterations to Pma1 affect $[\text{GAR}^+]$

a) $[\text{GAR}^+]$ induction by transient overexpression of $\text{PMA1}$ in a wild-type background. Data is shown as the mean of $[\text{GAR}^+]$ frequency +/- standard deviation (n=6). Western is total protein probed with $\alpha$Pma1 antibody and quantified using Scion Image. The blot was stained with Ponceau Red to confirm equal loading (right). b) Propagation of $[\text{GAR}^+]$ is impaired in $\text{PMA1A}40\text{N Astdl}$ double mutants. Tetrad spores from a $[\text{GAR}^+]$ diploid with the genotype $\text{GAL-PMA1A}40\text{N/PMA1 stdl}:\text{kanMX/STD1}$ were crossed to a $[\text{gar}^-]$ strain with wild-type $\text{PMA1}$ and $\text{STD1}$ alleles. $\text{PMA1A}40\text{N Astdl}$ spores cannot propagate $[\text{GAR}^+]$ to wild-type $[\text{gar}^-]$ yeast. The few glucosamine-resistant colonies found in the $\text{PMA1A}40\text{N Astdl}$ background exhibit standard, Mendelian inheritance of the glucosamine resistance phenotype and thus do not carry the $[\text{GAR}^+]$ element.
Figure 5: \([\text{GAR}^+]\) exhibits a Pma1-dependent species barrier

a) \([\text{GAR}^+]\) frequency of \(S. \text{bayanus}\) and \(S. \text{paradoxus}\) cells grown at 30°C (left), the optimal growth temperature of \(S. \text{paradoxus}\), or 23°C, the optimal growth temperature of \(S. \text{bayanus}\). Data is shown as the mean of \([\text{GAR}^+]\) frequency +/- standard deviation (n=6).

b) Substitution of \(PMA1\) from \(S. \text{cerevisiae}\) with \(PMA1\) from \(S. \text{bayanus}\) or \(S. \text{paradoxus}\) prevents \([\text{GAR}^+]\) propagation. Starting strain is haploid, \([\text{GAR}^+]\), genotype \(pma1::\text{kanMX}\) with p316-PMA1 \(S. \text{cerevisiae}\) as a covering plasmid. p314-PMA1 carrying \(PMA1\) from \(S. \text{cerevisiae}\) (\(S.\text{c.}\), top), \(S. \text{paradoxus}\) (\(S.\text{par.}\), middle), or \(S. \text{bayanus}\) (\(S.\text{bay.}\), bottom) was transformed in and p316-PMA1 \(S.\text{c.}\) selected against by replica plating to 5-FOA (\(S.\text{c.}\) 1N, \(S.\text{p.}\) 1N, or \(S.\text{b.}\) 1N). These haploids were mated to a wild-type \(S. \text{cerevisiae}\) [\(\text{gar}^-\)] background, restreaked two times, and tested for \([\text{GAR}^+]\). Representative data from three independent experiments is shown.

Figure 6: Pma1 and the Rgt2/Snf3 glucose signaling pathway

We propose that Pma1 acts as a part of the Rgt2/Snf3 signaling pathway. a) In \([\text{gar}^-]\) glucose-grown cells, Pma1 associates with Mth1. The glucose signal is propagated through Snf3 and Rgt2 to Yck1 and Yck2, which phosphorylate Mth1 and Std1. This phosphorylation marks Mth1 and Std1 for degradation, leaving their interacting partner, Rgt1, free in the cytosol, where it does not repress transcription at the \(HXT3\) locus. b) Under \([\text{GAR}^+]\) conditions, \(HXT3\) transcription is repressed, which resembles that of cells grown in a carbon source other than glucose. Pma1 associates with Std1, which somehow facilitates the repression of \(HXT3\), possibly by altering the affinity of Std1 for Rgt1. Association with Std1 has previously been shown to facilitate the binding of Rgt1 to
DNA (Lakshmanan et al. 2003). The association between Pma1 can either be transient or stable, but either way it aids in the establishment of an altered signaling pathway. This altered pathway is then maintained either by the contained association between Std1 and Pma1 or by a feedback loop within the signaling cascade itself.
A. [gar] [GAR] diploid 1 diploid 2

B. [gar] [GAR] strong [GAR] tetrad

C. [gar] [GAR] donor [gar] acceptor cytoductant

D. 

E. Δhsp104 Δhsp104 wildtype wildtype

F. Δssa1 Δssa2 wildtype Δssa1Δssa2
A

[gar]

GFP

DIC

[gar]

[Brown127548_Figure 2]

B

\[
\text{frequency relative to vector}
\]

C

D

E

tetrad spores

\Delta std1

wildtype

\Delta std1

wildtype

\Delta std1

\Delta std1

wildtype

\Delta std1

\Delta std1

wildtype

\Delta std1

\Delta std1

wildtype

\Delta std1

\Delta std1

wildtype

2% glucose

2% glycerol

G418
glycerol

0.05% GlcN

[gar]?
Brown127548_Figure 3

A

<table>
<thead>
<tr>
<th></th>
<th>Std1-6HA</th>
<th>Mth1-6HA</th>
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<tbody>
<tr>
<td>[gar]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>[GAR+]</td>
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αPma1

αHA

kDa

B

(IGAR⁺ cfu)/(total cfu)

p ≤ 10⁻⁵

[gar]  
Δsur4  
Δst1   
Δcb3   
Δcb4   
Δdp1   
Δag19

C

(IGAR⁺ cfu)/(total cfu)

p ≤ 2.8x10⁻⁵

p ≤ 4.5x10⁻⁵

wildtype  
S899A  
S899D  
S911A  
S911D  
T912A  
T912D  

D

% cells with Hxt3-GFP

p ≤ 10⁻¹³  
p ≤ 10⁻¹⁰  
p ≤ 10⁻¹⁷

[gar]  
[GAR+]  
S911A  
S911D  
T912A  
T912D  

B

C

D
Figure 4

Panel A: Bar graph showing the protein expression levels of PMA1 constructs. The graph compares the expression of PMA1, PMA1Δ40N, PMA1Δ104N, and PMA1Δ40C constructs. The y-axis represents the protein expression (kDa) normalized to the total cell number (cfu). The x-axis lists the constructs. Error bars indicate the standard deviation, and the p-value is marked as $p \leq 10^{-4}$.

Panel B: Table summarizing the spore growth experiment results. The table compares the growth of PMA1Δ40N std1 and wildtype spores in different conditions: 2% glucose, 2% galactose, SD-his, 2% glycerol, and 2% glycerol with 0.05% GlcN. The [GAR$^+$]? column indicates the growth response, with '-' for no growth and '+' for growth with GAR$^+$.
**A**

![Graph showing (GAR⁺) cfu/(total cfu)]

<table>
<thead>
<tr>
<th></th>
<th>S. bayanus</th>
<th>S. paradoxus</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 °C</td>
<td></td>
<td></td>
</tr>
<tr>
<td>23 °C</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**B**

![Image of yeast strains with different conditions]

- [gar⁺]
- [GAR⁺]
- S. cerevisiae 1N
- S. cerevisiae 1N x S. cerevisiae [gar⁻]
- [gar⁻]
- [GAR⁺]
- S. paradoxus 1N
- S. paradoxus 1N x S. cerevisiae [gar⁻]

**C**

![Bar graph showing (GAR⁺) cfu/(total cfu)]

- **STD1** overexpressed:
  - vector
  - S. cerevisiae
  - S. bayanus
  - S. paradoxus

- **origin of PMA1**: S. cerevisiae
  - S. bayanus
  - S. paradoxus

- **(GAR⁺) cfu/(total cfu)**
  - 0.0001
  - 0.001
  - 0.01
  - 0.1

- **Brown127548_Figure 5**

- S. bayanus