Expression, stability, and replacement of glucan-remodeling enzymes during developmental transitions in *Saccharomyces cerevisiae*

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**ABSTRACT** Sporulation is a developmental variation of the yeast life cycle whereby four spores are produced within a diploid cell, with proliferation resuming after germination. The GAS family of glycosylphosphatidylinositol-anchored glucan-remodeling enzymes exemplifies functional interplay between paralogous genes during the yeast life cycle. GAS1 and GAS5 are expressed in vegetative cells and repressed during sporulation while GAS2 and GAS4 exhibit a reciprocal pattern. GAS3 is weakly expressed in all the conditions and encodes an inactive protein. Although Gas1p functions in cell wall formation, we show that it persists during sporulation but is relocalized from the plasma membrane to the epiplasm in a process requiring End3p-mediated endocytosis and the Sps1 protein kinase of the p21-activated kinase family. Some Gas1p is also newly synthesized and localized to the spore membrane, but this fraction is dispensable for spore formation. By way of contrast, the Gas2–Gas4 proteins, which are essential for spore wall assembly, are rapidly degraded after spore formation. On germination, Gas1p is actively synthesized and concentrated in the growing part of the spore, which is essential for its elongation. Thus Gas1p is the primary glucan-remodeling enzyme required in vegetative growth and during reentry into the proliferative state. The dynamic interplay among Gas proteins is crucial to couple glucan remodeling with morphogenesis in developmental transitions.

**INTRODUCTION**

The life cycle of the budding yeast *Saccharomyces cerevisiae* is a fascinating example of alternative modes of reproduction and development of a simple eukaryotic cell. Yeast cells can exist in genetically stable haploid or diploid states, and they reproduce asexually by budding. In addition, haploid cells of opposite mating type (a or α) can mate. In the absence of a nitrogen source and in the presence of a nonfermentable carbon source, the resulting diploid a/α cell enters the alternative pathway of meiosis and sporulation. This pathway consists of a round of DNA replication followed by two nuclear divisions (meiosis I and II), which culminate in the formation of four haploid nuclei, each representing a meiotic segregant. Then each nucleus is surrounded by a spore wall to generate mature spores, which remain inside the diploid cell, forming a structure called an ascus. The four haploid spores can be released from the asc by treatment with hydrolytic enzymes that remove the asc wall. Finally, spores can germinate under favorable nutritional conditions, resuming the vegetative growth cycle.
conditions and reenter the mitotic cycle or mate again and restore the diploid state.

Over the years, the process of meiosis and sporulation has been studied in depth (Neiman, 2005). More recently a molecular characterization of the spore wall assembly pathway has been undertaken (Coluccio et al., 2004). The spore wall has a four-layer organization comprising from inside to outside 1) mannoproteins, 2) β(1,3)-glucan, 3) chitosan, and 4) dityrosine. The outer spore wall layers confer the spore much of its resistance to environmental stress. The deposition of spore wall components begins at the end of meiosis I when the prospore membrane (PSM) closure occurs. The PSM is a double membrane sac that originates from an initial priming structure, the meiotic outer plaque of the spindle pole body, to which the vesicles are redirected from the Golgi apparatus through a developmentally modified branch of the constitutive secretory pathway (Neiman, 1998; Morishita et al., 2007). The PSM extends progressively until it engulfs a haploid nucleus. The leading edges of the PSM fuse to form a double membrane compartment inside whose lumen spore wall material is deposited in a tightly regulated manner. At the end of the process, the outer membrane is dissolved. Spore germination is another important morphogenetic process. Under favorable conditions, nondividing haploid spores grow and reenter the mitotic cycle. At the germinating pole, a new wall with the typical three-layer organization of the vegetative cell wall—from inside to outside, chitin, glucan, and mannoprotein—is synthesized (Neiman, 2005).

Gas proteins belong to family 72 of glucanases/transglycosidases (Cantarel et al., 2009). Their β(1,3)-glucanase/transglycosidase dase activity is crucial for the incorporation and remodeling of β(1,3)-glucan into the cell wall and for creating attachment sites for the anchoring of mannoproteins and chitin (Mouyna et al., 2000; Ragni et al., 2007b; Rolli et al., 2009). In the S. cerevisiae genome, the GAS family is constituted by five paralogous genes: 1) GAS1 and GAS5, which are expressed during vegetative growth; 2) GAS2 and GAS4, which are repressed in vegetative growth and specifically induced during sporulation; and 3) GAS3, a weakly expressed gene encoding an inactive member of the family (Rolli et al., 2010). The Gas2 and Gas4 proteins are essential for proper spore assembly and viability (Ragni et al., 2007a). Gas1p plays a major role in cell wall construction during vegetative growth, whereas Gas5p presumably plays an ancillary function. GAS1 mRNA is also four times more abundant than GAS5 mRNA (Rolli et al., 2010). Overexpression of GAS genes has no affect on the phenotype of vegetative or sporulating yeast cells. GAS products are glycosylphosphatidylinositol (GPI)--anchored proteins, and while Gas2p and Gas4p are poorly mannosylated, Gas1, Gas3, and Gas5 are highly mannosylated proteins (Ragni et al., 2007a, 2007b; Rolli et al., 2010). Gas1p is predominantly anchored to the plasma membrane (PM) through GPI, but a fraction is also covalently linked to the cell wall (Yin et al., 2005). Accordingly, Gas1p localizes to microdomains of the PM, but it is also cross-linked to the chitin ring and bud scars, where it remains for many generations (Rolli et al., 2009). Moreover, at cytokinesis Gas1p localizes to the primary septum. At the bud neck and septum region, Gas1p is involved in the maintenance of the neck size and in cell separation (Rolli et al., 2009).

Here we investigated the dynamic changes occurring in the localization of Gas1p during the yeast life cycle with a view to analyze the functional interplay among Gas paralogous proteins. Our results provide novel information about the dynamics of Gas proteins and the roles of glucan-remodeling enzymes during spore morphogenesis and germination.

RESULTS

Expression profiles of GAS1 and GAS5 during meiosis and sporulation

GAS1 and GAS5 mRNA levels were monitored in the strain AN120, derived from the sporulation-proficient strain SK-1 (Kane and Roth, 1974). Total RNA was extracted at different time intervals after the induction of sporulation and used for a quantitative reverse transcriptase PCR (qRT-PCR) analysis. Actin mRNA was chosen as a reference transcript because the expression of ACT1 does not fluctuate significantly in sporulating cells (Primig et al., 2000). As illustrated in Figure 1A, the GAS1 and GAS5 transcripts showed very similar profiles. They were detectable at time 0 and remained roughly constant until 6 h after induction of sporulation, when they started to decrease, and at 10 h the levels were reduced by more than fourfold. These results are in accordance with previous Northern hybridization and transcriptome analyses (Popolo et al., 1993a; Chu et al., 1998; Primig et al., 2000).

As an internal control, we compared the expression profiles of GAS1 and GAS5 with that of GAS2 as a representative member of the GAS2-GAS4 gene pair (Figure 1A). Consistent with a previous analysis, we detected a peak of expression at 7 h (Ragni et al., 2007a). In conclusion, upon the induction of sporulation, either the expression of GAS1 and GAS5 is switched off and/or GAS1-GAS5 mRNAs are degraded.

Gas1 protein is stable during meiosis and sporulation

Total protein extracts were prepared at different time intervals after induction of sporulation and analyzed by immunoblot (Figure 1B). Gas1p was detected as a 130-kDa polypeptide. As shown in Figure 1B, the relative Gas1p level was constant throughout the sporulation process except for a slight increase in the first 10 h, which was probably due to completion of maturation of the precursors. The level of Gas1p remained constant during sporulation also in the W303 genetic background (Supplemental Figure S1). Thus the Gas1p present at time 0 appears to persist after the shift to the sporulation medium (SPM). Surprisingly, Gas1p was still detected at 45 h after sporulation induction, indicating that it is a highly stable protein (Figure 1, B and C). The persistence of Gas1p prompted us to examine its localization. As shown in Figure 1D, Gas1p was localized to the spore periphery in mature ascis. In conclusion, although GAS1 transcription is repressed, the product persists over time and its localization changes.

Localization of Gas proteins during meiosis and spore formation

To study the localization of other Gas proteins during meiosis and sporulation, Gas2p was tagged with three hemagglutinin (HA) epitopes, whereas Gas4p was recognized by an anti-Gas4p serum. Sporulating cells were collected during sporulation and processed for indirect immunofluorescence. Gas2 and Gas4 proteins could be visualized only when overexpressed. Previous studies showed that their overexpression had no effect on sporulation (Ragni et al., 2007a). At time 0, no fluorescence was detected (data not shown), in agreement with the absence of expression of GAS2 and GAS4 during vegetative growth (Ragni et al., 2007a). During meiosis I, Gas2p-HA exhibited a diffuse fluorescence in the cytoplasm that was more intense around the two nuclei (Figure 2A, a–c), in agreement with the absence of expression of GAS2-GAS4 mRNAs are degraded.

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ment with their role in the assembly of the glucan layer of the spore wall (Ragni et al., 2007a).

Gas1p localization was analyzed in more detail by taking advantage of fluorescent versions of Gas1p. In Gas1p–green fluorescent protein (GFP), GFP was inserted internally to leave intact the N- and C-terminal signal sequences required for protein maturation. This hybrid has been shown to be partially functional in vegetative cells (Rolli et al., 2009). Whereas gas1Δ diploid cells harboring the wild-type GAS1 on a centromeric plasmid sporulated normally, those carrying the GAS1-GFP fusion on the same vector were unable to sporulate and displayed the defective phenotype of gas1Δ cells (discussed later in this article). In contrast, gas1Δ cells transformed with a multicopy plasmid harboring GAS1-GFP (strain ER335) fully complemented the phenotype, as shown by the high sporulation efficiency (∼95% in a shift from semidefined presporulation medium [SA] to SPM). Accordingly, the experiments were performed with strain ER335. At different time points after the induction of sporulation, cells were collected to analyze Gas1p-GFP localization. In Gas1p-HA and Gas4p localization during meiosis and sporulation. (A) Gas2p-HA localization in cells at different stages of meiosis. Cells of strain ER314 (gas2Δ diploid overexpressing GAS2-3xHA), pregrown in SA, were examined at 8 h of sporulation by indirect immunofluorescence using anti-HA mAb (red fluorescence). (B) Gas4p localization during meiosis II. A culture of strain ER316 (a parental strain overexpressing Gas4p) was sporulated at 30°C, and cells were examined by indirect immunofluorescence with anti-Gas4p serum (green fluorescence). Images of DAPI staining (blue fluorescence), the immunofluorescence signal, and the merging of the two previous images are shown. Bar: 3 μm.

**FIGURE 1:** Time course of GAS1 and GAS5 transcription during meiosis and sporulation. (A) Cells were grown in YPD, precultured in YPA, and induced to sporulate in SPM. GAS1, GAS2, and GAS5 mRNA levels were measured by qRT-PCR. Relative mRNA levels are expressed as log2 of the ratio between the relative level at time 0 and the relative level at each subsequent time point. The C_T values in the same experiment at time 0 for GAS1, GAS5, and GAS2 were respectively 23.8, 22.2, and 31.8. The percentages of sporulated cells were 0% (time 0), 12% (8 h), 58% (10 h), 70% (12 h), 80% (15 h), and 85% (20 and 24 h). (B) Levels of Gas1p during sporulation. Equal amounts of protein extracts (100 μg) were analyzed by immunoblotting using anti-Gas1p serum and anti-actin mAb. (C) Relative Gas1p levels during sporulation. Actin was used for normalization. The relative level at time 0 was set at 1. (D) Localization of the wild-type Gas1p to the spore periphery as determined by indirect immunofluorescence in AN120 cells at 24 h of sporulation. Bar: 3 μm.
A. Gas1p-GFP localization changes during sporulation. A gas1Δ/gas1Δ mutant expressing Gas1p-GFP (strain ER337) was shifted from SA to SPM and allowed to sporulate. (A) At different time points, sporulating cells were stained with DAPI and observed under a fluorescence microscope to detect both the DAPI and Gas1p-GFP fluorescence. Bar: 3 μm. (B) Spores released from the ascis by treatment with Zymolyase at 24 h after the induction of sporulation. Bar: 3 μm. (C) An ascus ghost left after Zymolyase treatment of asci. To determine whether the fluorescence was associated with the spores, ascis were mildly treated with Zymolyase. As shown in Figure 3B, Gas1p-GFP decorated the spore periphery, indicating that the fluorescent labeling was associated with the spores. However, the fluorescence was less intense than in the intact ascus, suggesting that a fraction of Gas1p-GFP was lost during ascus wall digestion. Consistent with the removal of Gas1p-GFP from the mother PM during sporulation, the ascus ghosts were negative except for the presence of very bright rings (Figure 3C, right). These structures contained Gas1p-GFP cross-linked to the bud scars produced during vegetative growth (Rolli et al., 2009). Thus during sporulation the GPI-anchored Gas1p is completely removed from the PM of the mother cell and internalized, whereas the Gas1p cross-linked to the chitin of the bud scars remains static.

To exclude any artificial effect due to overexpression or type of fusion, we used a hybrid of Gas1p with the monomeric red fluorescent protein (mRFP-Gas1p). In this chimera, mRFP was inserted immediately after the signal peptide and the hybrid protein was fully functional (Rolli et al., 2009). One copy of the fusion gene was integrated at the LEU2 locus in the W303 genetic background (strain WER375). During vegetative growth, mRFP-Gas1p showed the typical localization sites of Gas1p at the PM, bud neck (Figure 4A), and scars (not shown in the figure). During sporulation, mRFP-Gas1p was found inside the sporulating cells and around the prospores in addition to the PM staining that was detected in cells with not yet visible spores (Figure 4B). In mature ascis, an intense staining was observed in the epiplasm (Figure 4C). The protein uniformly labeled the periphery of the released spores (Figure 4D). Thus both the relocation of Gas1p from the PM to the epiplasm and the association to the spore periphery are independent of the fusion protein used and the genetic background. In addition, because the mRFP-GAS1 fusion was carried by only one parental strain, from which the diploid was generated and an equivalent partitioning of the hybrid protein to the spores was observed (fluorescent spores > 99%), we can conclude that transport of either the preexisting or the new synthesized protein could occur.

To answer the question of the origin of the spore-associated Gas1p, the GAS1 promoter of the GAS1-GFP construct was replaced by the CLB2 promoter, which is repressed upon induction of sporulation. In this strain, the fate of the vegetative protein present at the PM can be analyzed without the interference of the protein that can be synthesized in the first hours of sporulation. In vegetative cells Gas1p-GFP was detected at the typical sites (Figure 5A). At 24 h of sporulation, a bright fluorescence was present around and between the spores (Figure 5B). However, no labeling was observed at the periphery of the released spores (Figure 5C). Taken together, these data support the idea that Gas1p localization at the spore depends on newly made protein, whereas the protein present at the PM during vegetative growth is transported to the cytoplasm but excluded from the developing spores.

To confirm these results, we also examined the level of untagged Gas1p present in the epiplasm and spore extracts. Mature ascis from a wild-type diploid strain (AN120) were treated with Zymolyase, and no fluorescence is detectable except for the two bright rings corresponding to the Gas1p cross-linked to the bud scars. Bar: 5 μm.
Thus Gas1p molecules localized to the spores represent a minor fraction of the total Gas1p present in the ascus. The newly synthesized mRFP-Gas1p localizes to the PSMs. A colocalization analysis with GFP-Spo20p, a sporulation-specific member of the (SNAP25)-t-SNARE family that specifically marks the PSMs in developing spores, was performed (Neiman, 1998; Nakanishi et al., 2004, 2006). Cells coexpressing mRFP-GAS1 and SPO2051–91-GFP were induced to sporulate. The Spo20p fragment contains the membrane localization signal but lacks the functional domain of Spo20p. During spore development, GFP-Spo20p resolved into ringlike structures representing the PSMs (Supplemental Figure S2). mRFP-Gas1p gave origin to a diffuse staining pattern in the cytoplasm in addition to a membrane labeling pattern overlapping that of Spo20p (Supplemental Figure S2, merge). These results support the existence of a pool of newly synthesized Gas1p molecules that are localized to the PSMs. A sufficient amount of protein is required to detect the fluorescence around the spores and to distinguish it from the cytoplasmic mRFP-Gas1p fluorescence.

The newly synthesized mRFP-Gas1p localizes to the PSMs

We further examined whether the newly synthesized Gas1p was incorporated in the PSM. A colocalization analysis with GFP-Spo20p, a sporulation-specific member of the (SNAP25)-t-SNARE family that specifically marks the PSMs in developing spores, was performed (Neiman, 1998; Nakanishi et al., 2004, 2006). Cells coexpressing mRFP-GAS1 and SPO2051–91-GFP were induced to sporulate. The Spo20p fragment contains the membrane localization signal but lacks the functional domain of Spo20p. During spore development, GFP-Spo20p resolved into ringlike structures representing the PSMs (Supplemental Figure S2). mRFP-Gas1p gave origin to a diffuse staining pattern in the cytoplasm in addition to a membrane labeling pattern overlapping that of Spo20p (Supplemental Figure S2, merge). These results support the existence of a pool of newly synthesized Gas1p molecules that are localized to the PSMs. A sufficient amount of protein is required to detect the fluorescence around the spores and to distinguish it from the cytoplasmic mRFP-Gas1p fluorescence.

Gas1p internalization is impaired in end3Δ mutants

Internalization of GPI-anchored Gas1p could be mediated by endocytosis. Thus Gas1p-GFP localization was analyzed in an end3Δ mutant, which is defective in endocytosis. In this mutant, owing to a
failure in the localization of the glucan and chitin synthase to the PSMs the deposition of glucan and chitosan layers does not occur (Morishita and Engebrecht, 2005). At 30°C, the sporulation efficiency of the end3Δ mutant was only ∼45% (n = 200), with an enrichment of asci with one, two, or three spores, in agreement with previously reported data (Morishita and Engebrecht, 2005). At 24 h after the induction of sporulation, Gas1p-GFP localized to the periphery of the spores and between the spores in the wild-type strain (Figure 7, a and a'), whereas it was mislocalized in the majority of end3Δ cells (Figure 7, b–f). Gas1p-GFP was detected mainly in the cytoplasm of abnormally shaped cells and excluded from the spores (Figure 7, b and b'). In some cells, Gas1p-GFP remained at the PM of the mother cells even when refractive spores were already visible (Figure 7, c–f). In ∼35% of the sporulated cells, Gas1p-GFP showed only a faint fluorescence on the spores, indicating that some new synthesis of the spore-associated protein occurred (Figure 7, e' and d'). In conclusion, End3p-mediated endocytosis contributes to the efficient removal of Gas1p from the PM although other pathways are likely to be involved.

Gas1p is not internalized in an sps1Δ mutant

Sps1p encodes a sporation-specific serine/threonine protein kinase with homology to members of the p21-activated kinases (PAKs) (Friesen et al., 1994; Park and Bi, 2007). It has previously been shown that sps1Δ mutants are not affected in the progression of meiosis or PSMs formation but form only aberrant spore-like compartments (Friesen et al., 1994; Iwamoto et al., 2005). Sps1p does not affect the developmentally regulated branch of the secretory pathway but regulates the trafficking of the enzymes involved in spore wall biogenesis to the PSMs (Iwamoto et al., 2005). To determine whether Gas1p internalization is regulated by Sps1p, we analyzed Gas1p localization during sporulation in an sps1Δ null diploid mutant. Gas1p-GFP fluorescence was detected in the PM in most cells that had completed meiosis II, and it was also slightly diffused throughout the cytoplasm, although it was never observed at the PSMs or in spore-like compartments (Figure 7, g–g”). Thus the sps1Δ mutant appears to be defective in triggering the internalization step of PM-associated Gas1p-GFP.

Functional characterization of GAS1 during sporulation

The presence of a fraction of Gas1p in the spore suggests a potential involvement of the protein in spore wall assembly or in subsequent germination. We first examined the effect of GAS1 inactivation on sporulation. The phenotypic analysis of gas1Δ/gas1Δ cells during sporulation was complicated by the fact that the mutant was unable to grow in presporulation media containing acetate as a carbon source. Cells appeared swollen and underwent lysis. A spot assay on solid medium indicated that the viability of gas1Δ/gas1Δ cells was very low on acetate, glycerol, and ethanol with respect to
glucose (Supplemental Figure S3). Thus gas1Δ phenotype is worsened in the presence of nonfermentable carbon sources. This defect was common to gas1/gas1 null mutants derived from SK-1 or W303 genetic backgrounds (Supplemental Figure S3). In an attempt to prevent growth defects of gas1Δ/gas1Δ cells, 1 M sorbitol was included in 5A as an osmotic stabilizer. Under this condition, cells grew slowly (doubling time of ~8 h), and the lysis phenotype of the mutant was partially mitigated (Figure 8A). To induce sporulation, cells were shifted to SPM supplemented with 1 M sorbitol. Both the mutant and the isogenic strain completed the process in 48 h. The sporulated population of mutant cells was composed of asci of different sizes, many lysed cells forming large aggregates, and also multisporic asci (Figure 8B). The sporulation efficiency of the gas1Δ mutant was ~36%, whereas in the wild type it was ~78% (Table 1). Thus the defects carried on from vegetative growth on acetate affect the morphology and sporulation efficiency of the gas1Δ mutant.

To check whether mutant spores were affected in spore wall maturation, the dityrosine layer was analyzed. The gas1Δ mutant showed normal dityrosine accumulation and deposition, suggesting cells undergoing sporulation have a normal deposition of the cell wall, including the outermost layer (Figure 8C). Accordingly, we monitored the organization of septins, a cytological marker of spore formation. Septins are organized at the leading edge of the PSM and facilitate spore wall deposition (Fares et al., 1996). Their localization is altered in mutants in which PSM closure is not sensed or has failed (Tachikawa et al., 2001). The localization of Spr28p, a sporulation-specific septin and a marker of PSM closure, was examined (De Virgilio et al., 1996). The septins formed parallel bar structures and then eventually surrounded the spores in the wild type (Figure 8D, left). An analogous pattern was observed in gas1Δ cells, suggesting that septins are properly organized in the absence of Gas1p (Figure 8D, right). Thus PSMs undergo closure in the gas1Δ mutant. In conclusion, Gas1p does not play any appreciable role during spore morphogenesis and spore wall assembly.

A striking phenotypic trait of the gas1Δ/gas1Δ mutant was the presence of ascii with more than four nucleate spores (Table 1, Figure 8B). This trait could be explained by the presence of ~3% of bi- and trinucleate cells during vegetative growth, probably derived from endomitosis caused by the cell wall defects (unpublished data). Hence, multisporic asci were the meiotic products of these cells.

**Gas1p-GFP is highly polarized during spore germination**

Because Gas1p appears to play a dispensable role in spore wall assembly, we analyzed its possible involvement in germination. Spore germination consists of the exit from quiescence and adaptation to the new nutritional condition, followed by the entry into the proliferative state (Joseph-Strauss et al., 2007). Asci were collected after 24 h of sporulation and treated with Zymolyase, and the released spores were induced to germinate. As shown in Figure 9, at time 0 Gas1p-GFP uniformly decorated the spore periphery. One hour later, Gas1p-GFP was still localized to the spore periphery. At 2 h, the protein concentrated in one part of the spore, with a crescent-like aspect. Notably, the fluorescent protein marked the site of growth polarization. As germination proceeded, the Gas1p-GFP signal was almost undetectable in the nongrowing half, and it concentrated in the growing half as a bright crescent (3 h). At 4 h, bud emergence was observed, and Gas1p-GFP was still highly polarized and decorated the periphery of small new buds. From this stage, it was possible to recognize the Gas1p-GFP localization pattern typical of vegetative growth (Rolli et al., 2009) (5 h). These results

![FIGURE 8: Phenotype of gas1Δ/gas1Δ sporulating cells. (A) The gas1Δ/gas1Δ cells (strain ER320) growing exponentially in SA with 1 M sorbitol (time 0). In the inset, the morphology of a wild-type cell (strain AN120). (B) The gas1Δ/gas1Δ cells at 24 h after the shift to SPM, supplemented with 1 M sorbitol. In the inset, a wild-type ascus is shown. A multisporic ascus (m) is indicated (arrow). (C) Microscopic analysis of intact asci for the study of the coherence of the dityrosine layer in the presence of 1 M sorbitol. (D) Septin organization is normal in gas1Δ/gas1Δ sporulating cells. Bar: 3 μm.](image)

| Strains          | Medium          | Unsporulated | >4 spores | Distribution of ascus type (%)a |
|------------------|-----------------|--------------|-----------|--------------------------------|
|                  |                 |              |           | Monad | Dyad | Triad | Tetrad |
| SK-1             | SMP-1 + sorbitol| 22.5 ± 9.7   | 0         | 1.3 ± 1 | 17.6 ± 5.4 | 24.9 ± 3.5 | 34.2 ± 13 |
| gas1Δ/gas1Δ      | SMP-1 + sorbitol| 64.4 ± 10.3  | 6 ± 6.3   | 0.3 ± 0.6 | 7.4 ± 1.1 | 9.4 ± 4.4 | 12.9 ± 3.9 |

Values are expressed as means ± SD of data obtained from three independent experiments at 48 h after the induction of sporulation.

**TABLE 1: Effects of GAS1 deletion on sporulation.**
Growth polarization is defective in gas1Δ germinating spores

We next analyzed the effects of the loss of Gas1p during spore germination. Wild-type and gas1Δ spores were released from the ascus and induced to germinate. As shown in Figure 10A, at time 0 no significant morphological difference could be detected between the parental and gas1Δ spores. At 1 h, the spores were still round. From 2 to 7 h, polarized growth became progressively more evident in the wild type, with the presence of elongated cells, whereas gas1Δ spores maintained their round shape. Also, the buds that emerged from gas1Δ cells at later times (5–7 h) were rounder than the wild-type buds. These morphological defects are in agreement with the previously described phenotypic traits of the mutant during vegetative growth (Popolo et al., 1993b) and support a primary role for Gas1p in determining cell shape.

A quantitative analysis of gas1Δ defects in germination was performed by measuring the length of the long and short axes of the germinating spores (Figure 10B). The longest segment along the direction of growth was defined as the long axis, while the short axis was taken as the longest segment in a direction perpendicular to the long axis. As shown in Figure 10B, in wild-type cells the long and short axes started to increase at 2 h, but the long axis increased more rapidly than the short one, in good agreement with previously published data (Kono et al., 2005). In the mutant, both the long and short axes remained more or less constant over the first 2 h, after which they increased slightly until reaching approximately the same length. For the wild type, the long:short axis ratio was 1.19 ± 0.15 at time 0 and increased to a value of 1.6 ± 0.28 at 7 h, whereas for the mutant it was 1.12 ± 0.24 at time 0 and 0.99 ± 0.17 at 7 h. These results indicate that Gas1p plays a pivotal role in cell elongation in the early stages of spore germination (2–3 h). At the time of bud appearance and during bud maturation (5–7 h), Gas1p is still remarkably implicated in growth polarization.

Dynamic interplay of Gas1p, Gas2p, and Gas4p in sporulation and germination

To study the dynamic relationships among Gas1, Gas2, and Gas4 proteins, the ER314 strain was induced to sporulate. We took advantage of this strain to simultaneously detect wild-type Gas1 and Gas4 proteins and Gas2p-HA, the latter being expressed from a high-copy plasmid. As shown in Figure 11A, the Gas2-3xHA and Gas4 proteins were absent in vegetative growing cells and were present at 8 h after induction of sporulation, in agreement with the time of the maximal expression (7 h) of GAS2 and GAS4 genes (Ragni et al., 2007a and Figure 1A). At 24 h, Gas4p was undetectable, whereas Gas2p-3xHA was still present but its level was reduced. Next Gas2p-3xHA and Gas4p levels were monitored during germination. Gas4p was absent, whereas Gas2p-3xHA decreased steadily over time. In contrast, the level of Gas1p persisted during sporulation and increased during germination (Figure 11A). At 24 h, Gas4p was undetectable, whereas Gas2p-3xHA was still present but its level was reduced. Next Gas2p-3xHA and Gas4p levels were monitored during germination. Gas4p was absent, whereas Gas2p-3xHA decreased steadily over time. In contrast, the level of Gas1p persisted during sporulation and increased during germination (Figure 11A). These results point to a lack of new synthesis and the concomitant degradation of Gas2 and Gas4 proteins during sporulation and early germination (see Discussion). Moreover, the results show that wild-type Gas1p persists during sporulation and early germination and its level increases rapidly upon entry in germination, consistent with the induction of GAS1 transcription by transfer from SPM to yeast peptone dextrose (YPD) (see Discussion).

Gas1p levels in old asci

To explore the physiological significance of the persistence of Gas1p in the mature ascus, we examined the levels of Gas1p during aging of the spores. Wild-type cells were pregrown either in yeast peptone acetate (YPA) or in SA and then induced to sporulate in old asci.
S. cerevisiae

Multigene families offer a good model to study the interplay among paralogous proteins and their interactions for the fine-tuning of different biological processes. One such family is the Gas family of paralogous proteins and their interactions for the fine-tuning of different biological processes. One such family is the Gas family of paralogous proteins and their interactions for the fine-tuning of different biological processes.

DISCUSSION

Multigene families offer a good model to study the interplay among paralogous proteins and their interactions for the fine-tuning of different biological processes. One such family is the Gas family of paralogous proteins and their interactions for the fine-tuning of different biological processes.

Dynamic interplay of proteins involved in spore wall formation

Overall, the formation of the spore wall requires a specific set of enzymes, some of which are reused from vegetative growth while others are replaced by sporulation-specific isofoms. Moreover, spatial and temporal regulation of protein trafficking is crucial for proper spore formation. Remarkably, Fks1p, the predominant vegetative form of the β(1,3)-glucan synthase during growth on glucose, is replaced by Gsc2p/}

Fks2p (Ishihara et al., 2007). GSC2/FKS2 expression is induced during sporulation (Mazur et al., 1995). Gsc2/Fks2p reaches the PM before being recycled to the PSMs (Morishita and Engebrecth, 2005). Similarly to Fks proteins, Gas2p and Gas4p replace Gas1p function in spore wall assembly and are degraded after execution of their function (Figure 11A). With regard to chitosan synthesis, CHS3, encoding the catalytic subunit of chitin synthase III and responsible for the synthesis of chitosan (Pammer et al., 1992), is not transcriptionally up-regulated during sporulation, but the protein is recycled (Iwamoto et al., 2005), and this is consistent with the notion of it being an enzyme that is mainly regulated at the posttranslational level (Choi et al., 1994; Valdivia and Schekman, 2003). On the contrary, the activator, Chs4p, is quickly degraded upon induction of sporulation and is replaced by the sporulation-specific Schc1p, which is also directed to the PSMs (Sanz et al., 2002; Iwamoto et al., 2005).

Interestingly, in recent years END3-mediated endocytosis has been shown to play an important role in membrane trafficking events required for spore wall formation (Morishita and Engebrecth, 2005). Chs3p and Gsc2p/Fks2p require End3p for their relocalization to the PSMs. In the present work, we demonstrate that in the absence of End3p, Gas1p is also inefficiently internalized. Thus a partially common mechanism could mediate the transport of Gas1p, Chs3p, and Gsc2p inside the sporulating cell,
although the destiny of Gas1p is different from that of Chs3p and Gsc2p, with the latter being diverted to the PSMs while Gas1p is sequestered in the epiplasm.

Internalization of Gas1p-GFP also requires the activity of the Sps1 PAK kinase, as also reported for Chs3p, Shc1p, and Gsc2p/Fks2p (Iwamoto et al., 2005). The defect has been attributed to failure in the intracellular movement because endocytosis appears normal in the sps1Δ mutant (Iwamoto et al., 2005). PAK kinases are activated by the small G-protein Cdc42p (Zhao and Manser, 2005; Perez and Rincon, 2010). In yeast, the best known members are Ste20p, Cla4p, and Skm1p. Once activated, PAK kinases regulate downstream effectors that play a crucial role in cytoskeleton dynamics, pheromone response, cell cycle, and gene transcription. In addition, another protein kinase, the Smk1p mitogen-activated protein kinase, controls the transition between the deposition of the inner and outer layers of the spore wall by negatively regulating Fks2p. There is evidence that Sps1p and Smk1p have distinct functions (Ufano et al., 1999; Neiman, 2005). Because no effector of Sps1p is known, it may be surmised that Sps1p would regulate a crucial step in actin cytoskeleton remodeling required for the endocytic pathway of Gas1p. Alternatively, because the sps1Δ mutant lacks proper coordination between the end of meiosis II and nuclear engulfment, and this causes defects in cell wall formation and transcription, it may be proposed that the same signal would trigger the mobilization of Gas1p.

Interestingly, dynamic changes have also been described for septins (Cdc3p, Cdc10p, Cdc11p, Cdc12p, and Shs1p). During sporulation CDC12 is repressed, but the preformed protein persists, although in the creation of the new structures Cdc12p is replaced by the sporulation-specific septin Spr3p. On germination, only newly synthesized Cdc12 molecules are used for the creation of new structures. By contrast, Cdc10p synthesized during vegetative growth is reutilized to build sporulation-specific structures and then reused again during spor germination (McMurray and Thorner, 2008, 2009). Thus tracking the fate of molecules along the yeast life cycle is becoming a fascinating area of yeast cell biology.

**Physiological significance of Gas1p persistence during sporulation**

The presence of Gas1p at the spore periphery suggests that in addition to its role during vegetative growth, Gas1p might be involved in spore wall assembly. For instance, the cross-linking of the (1,3)-glucan and chitosan layers could require an increase in the branching level of the glucan or in the length of the branches, and hence different proteins of the GH72 family might become necessary. However, the phenotype of gas1Δ/gas1Δ mutants in sporulation indicates that Gas1p plays a dispensable role in spore morphogenesis in S. cerevisiae. Thus the removal of Gas1p from the PM could be necessary to soften the cell wall of the diploid cell in preparation for its collapse on the spores. Indeed, the ascus wall is less resistant than the vegetative cell wall (Coluccio et al., 2008).

Our finding that the rich presporulation medium YPA supports greater stability of wild-type Gas1p than semidefined SA medium also suggests that in the asc Gas1p is slowly degraded at a rate that is dependent on the nutritional state of the diploid cells that generated them.

**Gas1p is required for the elongation of the germinating spore**

Germination can be divided into distinct stages: the uncoating of the spore wall (1 h), polarized growth (2 h), isotropic growth (3 h), and bud emergence (4 h) (Joseph-Strauss et al., 2007). Here we show that Gas1p is essential for growth polarization during germination. In addition, Gas1p-GFP was found to be highly polarized to the growing half of the germinating spores. To check when Gas1p-GFP was present in the database (M. Barkai, personal communication). Interestingly, GAS5 transcription increased very rapidly and reached a maximum within the first 2 h of germination (Joseph-Strauss et al., 2007). Because the present and previous studies indicate that GAS1 and GAS5 share a common pattern of transcription, it may be assumed that GAS1 is also similarly induced during early germination. The lack of elongation in germinating gas1Δ spores points to a primary role of Gas1p in coupling the events required for growth polarization with cell wall biosynthesis.

Although the induction of GAS1 and GAS5 transcription seems to be rapid, protein synthesis and maturation may delay the appearance of newly synthesized enzymes, in particular because Gas1p protein is abundantly modified by N- and O-mannosylation and by GPI-attachment. Thus the spore-associated Gas1p could supply some...
activity at the onset of spore germination before the new molecules become available.

MATERIALS AND METHODS

Yeast strains, growth, sporulation, and germination conditions

The strains are listed in Table 2. Cells were routinely grown at 30°C in YPD (1% yeast extract, 2% Bacto peptone, 2% glucose) or in synthetic dextrose minimal medium (Difco yeast nitrogen base without amino acids at 6.7 g/l, 2% glucose) to which the required supplements were added at concentrations of 50 mg/l for the amino acids and uracil and 100 mg/l for adenine. To induce sporulation, logarithmic cultures were shifted from YPD to YPA (1% yeast extract, 2% Bacto peptone, 2% acetate) or to semidefined presporulation medium (SA) (10 g potassium acetate, 6.7 g yeast nitrogen base without amino acids, 1 g yeast extract in 1 l of 0.05 M phthalate buffer, pH 5) at an initial optical density of 0.2 OD before being shifted to SPM (1% potassium acetate) at a cell density of ~10^7 cells/ml (OD450) at an initial optical density of 0.2 OD450. Cells were grown at 30°C, collected the following morning, and washed with sterile distilled water without amino acids, 1 g yeast extract in 1 l of 0.05 M phthalate buffer, pH 5) at an initial optical density of 0.2 OD before being shifted to SPM (1% potassium acetate) at a cell density of ~10^7 cells/ml (OD450) at a 1:10 ratio of culture volume to flask volume. Cultures were allowed to sporulate under vigorous shaking at 30°C. The percentage of sporulation was determined by microscopic examination (magnification 400x) of the cultures. Sporulation was defined as the presence of ascospores (sporangia) in the culture. The percentage of sporulation was calculated as the ratio of the number of ascospores to the total number of cells observed under the microscope.

| Strain | Genotype | Source |
|--------|----------|--------|
| SK-1   |          |        |
| AN120  | MATα/α ura3/ura3 his3/3 leu2/leu2 trpl/trpl1 ARG4 rme1::LEU2/RME1 | Neiman et al., 2000 |
| ER306  | MATα/α arg4/ARG4 gas2::HIR3/gas2::HIR3 his3/his3 ho::LYS2/ho::LYS2 leu2/leu2 lys2/lys2 rme1::LEU2/RME1 trpl/trpl1 ura3/ura3 | Ragni et al., 2007b |
| ER307  | MATα/α arg4/ARG4 gas2::KanMX2/gas4::KanMX2 his3/his3 ho::LYS2/ho::LYS2 leu2/leu2 lys2/lys2 rme1::LEU2/RME1 trpl/trpl1 ura3/ura3 | Ragni et al., 2007b |
| ER310  | AN120, plus YE24 (2-μM URA3) | Ragni et al., 2007b |
| ER314  | ER306, plus YE24-GAS2-3xHA (pYER-2-HA) | Ragni et al., 2007b |
| ER316  | AN120, plus YE24-GAS4 (YER-4) | Ragni et al., 2007b |
| ER320  | AN120, but gas1::KanMX/gas1::KanMX | Rolli et al., 2009 |
| ER335  | ER320, plus YE24-GAS1 | Rolli et al., 2009 |
| ER337  | ER320, plus YE24-GAS1-GFP | Rolli et al., 2009 |
| ER339  | ER320, plus pRS416-GAS1 | Rolli et al., 2009 |
| ER346  | ER320, plus pRS426-mRFP-GAS1 | This study |
| ER348  | ER320, plus YE24-GAS1-GFP and pRS424-PpSPO25-SEC61-RFP | This study |
| ER350  | ER320, plus pRS426-mRFP-GAS1 and pRS424-G20 | This study |
| Y4733  | AN120, but end3::KanMX/end3::KanMX | Morishita and Engelbrecht, 2005 |
| ER351  | Y4733, plus YE24-GAS1-GFP | This study |
| ER354  | AN120, plus pRS416-SPR28-GFP | This study |
| ER355  | ER320, plus pRS416-SPR28-GFP | This study |
| ER374  | AN120 plus pYE24-PC422-GAS1-GFP | This study |

TABLE 2: S. cerevisiae strains used in this work.
−75–85% when cells were pregrown in YPA and −95% when pregrown in SA. Spore purification was performed as previously described (Esposito et al., 1991). To induce germination, a 10-ml aliquot of a sporulated culture was harvested, washed once with phosphate-buffered saline (PBS; pH 7.2), and then resuspended in PBS containing 20 μg/ml Zymolyase 20T (ICN Biomedicals, Irvine, CA). Cells were incubated for 15 min at 37°C to release the spores. Then the samples were washed twice with PBS, resuspended in YPD at a final concentration of 1–2 × 10⁷ cells/ml, and allowed to germinate at 30°C. For the release of Gas1p spores, ascus digestion was performed in PBS containing 4 μg/ml Zymolyase 20T. To obtain old asci, cultures at 48 h of sporulation were kept static at 20°C for several weeks. To check spore germination, spores were resuspended in YPD at 30°C and increase in OD₄₅₀ was monitored.

Quantification of mRNA using real-time qRT-PCR
Total RNA extraction, cDNA preparation, and real-time PCR were performed as previously described (Ragni et al., 2007a). The primer pairs for GAS1 and GAS5 were GAS1 forward 5′-AGGTAAGTGT-TGATTTGGGTTCand gas1 reverse 5′-AGAAGACCACCCAAGGCTTA; GAS5 forward 5′-CTCCGTGACTTACCAAGGCTC and GAS5 reverse 5′-ATGCGGCGAAGTTGACG. In the case of ACT1 and GAS2, the primers have been described previously (Ragni et al., 2007a). Each cDNA was assayed in at least duplicate PCRs for two independent experiments. Basic analysis was performed using SDS 1.9.1 software (Applied Biosystems, Carlsbad, CA). For further elaboration of the data, the Livak method (Livak and Schmittgen, 2001) was used. Briefly, from each duplicate reaction, a ΔΔCt was calculated by subtracting the average cycle threshold (Ct) value of ACT1 from the average Ct value of the gene of interest for the same time. Then the difference between the ΔCt at any time and the ΔCt at time 0 was calculated (ΔΔCt). The plotted values are 2−ΔΔCt.

Plasmid construction
The plasmids are listed in Table 3. pRS426-mRFP-GAS1 was constructed by cloning the SacI-BamHI fragment from pMF608, kindly provided by Y. Jigami (National Institute of Advanced Industrial Science and Technology, Tsukuba, Japan) and described previously (Rolli et al., 2009), into similarly cut pRS426. To place the GAS1-GFP fusion under the control of the CLB2 promoter (PCLB2), the BamHI/Smal fragment of YEp24-GAS1-GFP containing the GAS1-GFP cassette (including GAS1 promoter and terminator) was cloned in pRS416, generating pRS416-GAS1-GFP. Then the GAS1 promoter was swapped with the PCLB2 by cloning the 1-kb DNA fragment of the −1000/~1 CLB2 region upstream from the GAS1 coding sequence. The pRS416-PCLB2-GAS1-GFP was obtained. The DNA fragment was generated by PCR using the SK-1 genomic DNA as template and the primers CLB2prom-Smal (GCTACTCCGGC-GACCCGTGTTGTTGACCGTC) and Smal restriction site underlined, CLB2 sequence −1000/~983 in italic) and CLB2prom-BstAPI (AAAAGCAGCAGCGGTTGCTAATCCTGGAAAAAGGGATTAAACAACAATCTATAAGATCAATGAAGA, BstAPI restriction site underlined, CLB2 sequence −18/~1 in italics, GAS1 coding sequence +1/+46 in bold). Finally, the Smal-BamHI fragment of pRS416-PCLB2-GAS1-GFP containing PCLB2-GAS1-GFP was transferred into a similarly cut YEp24 to create YEp24-PCLB2-GAS1-GFP.

Microscopy
Cells were observed by phase-contrast microscopy, and sporulation was scored by counting at least 200 cells after a mild sonication. For Gas1p-GFP visualization, 1 ml sporulating culture was centrifuged at 8000 rpm for 2 min at 4°C, washed twice with cold PBS, and incubated for 15 min on ice. If required, 8.3 μg/ml 4,6-diamidino-phenylindole (DAPI) was added to the cells. Samples were then incubated for 15 min at room temperature in the dark before microscopy observation. Cells were observed as wet mounts using an Eclipse 90i (Nikon, Tokyo, Japan) or a DMRXA (Leica, Wetzlar, Germany) microscope equipped with epifluorescence, Nomarski optics, and a Hamamatsu ORCA-ER camera (Nihonbashi, McHenry, IL). The setup, including the microscope and camera, was controlled by MetaMorph software (Molecular Devices, Sunnyvale, CA). Alternatively, cells were examined with an Olympus BX60 microscope (Olympus Optical, Tokyo, Japan) connected to a DC290 Kodak digital camera. The images were analyzed using ImageJ-BMF software (McMaster Biophotonics Facility, Hamilton, ON, Canada) and Adobe Photoshop (San Jose, CA). The observation of the natural fluorescence of ditirosine was performed as described previously (Briza et al., 1986).

For indirect immunofluorescence, sporulating cells (20 OD₄₅₀) were fixed in 3.7% formaldehyde−0.1 M KPO₄ buffer for 30 min. After a 3-min centrifugation at 1500 rpm, the cells were resuspended in the same volume of fixing solution (0.1 M KPO₄, 3.7% formaldehyde) for 2–4 h. Fixed cells were washed and resuspended in SHA buffer (1 M sorbitol, 0.1 M HEPES-KOH, pH 7.5, and 5 mM NaN₃) at a concentration of 10⁸ cells/ml. A small aliquot of fixed cells (150 μl) was centrifuged and resuspended in SHA supplemented...
with 25 μg/ml Zymolyase 20T (ICN Biomedicals) and 0.2% β-mercaptoethanol. After a 30-min incubation at 37°C, removal of the ascus sac was checked before proceeding. Spheroplasts were then permeabilized by incubating in SHA and 0.1% Triton X-100 for 5 min. After attaching to glass slides, cells were plunged into −20°C methanol for 6 min, followed by −20°C acetone for 30 s. After a blocking step in PBS + block (1% milk, 0.5% bovine serum albumin [BSA] in PBS) for 10 min, slides were incubated in primary antibody diluted in PBS + block for 2 h. Primary antibodies were anti-Gas1p rabbit serum (diluted 1:500), anti-β-HA monoclonal antibody (mAb) (diluted 1:1000; Covance, Berkeley, CA), and anti-Gas4p serum. Rabbit immunoglobulin (Ig) G was purified from anti-Gas4p serum using a protein A microspin column and diluted 1:50. Slides were washed 12 times with PBS + 0.5 mg/ml BSA and then incubated in Alexa Fluor 594 goat anti–mouse IgG (1:1000 dilution) or Alexa Fluor 488 goat anti–rabbit IgG (1:1500 dilution) for 1 h. After 12 washes with PBS, or with PBS–0.5% BSA–0.1% Triton X-100 for Gas4p, a mounting medium (Gel Mount; Biomedora Corporation, Foster City, CA) was added to the slides, which were kept at 4°C for 1 h and observed under an epifluorescence microscope.

Extract preparation, electrophoresis, and immunoblotting
Sporulating cells (2 × 10^8) were collected by filtration, washed, and resuspended in ice-cold dH_2O. After a 2-min centrifugation at 4°C, the pellet was rapidly frozen and stored at −80°C. Extract preparation, determination of protein concentrations, and immunoblotting were performed as described previously (Gatti et al., 1994; Ragni et al., 2007a). Anti-Gas1p serum, anti-actin, or anti-β-HA mAbs and an anti-Gas4p serum, diluted 1:1000, were used (Rolli et al., 2009). Peroxidase-conjugated, affinity-purified F(ab')2 fragment donkey anti–rabbit or anti–mouse IgG were used (1:10,000 dilution; Jackson Laboratory, Bar Harbor, ME). Bound antibodies were developed using enhanced chemiluminescence Western blotting detection reagents (Amersham Pharmacia Biotech, Piscataway, NJ). Densitometry measurements were performed using the Scion Image program (Scion, Frederick, MD).

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