Elements of a single MAP kinase cascade in *Saccharomyces cerevisiae* mediate two developmental programs in the same cell type: mating and invasive growth

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Diploid *Saccharomyces cerevisiae* strains starved for nitrogen undergo a developmental transition from a colonial form of growth to a filamentous pseudohyphal form. This dimorphism requires a polar budding pattern and elements of the MAP kinase signal transduction pathway essential for mating pheromone response in haploids. We report here that haploid strains exhibit an invasive growth behavior with many similarities to pseudohyphal development, including filament formation and agar penetration. Haploid filament formation depends on a switch from an axial to a bipolar mode of bud site selection. Filament formation is distinct from agar penetration in both haploids and diplodips. We find that the same components of the MAP kinase cascade necessary for diploid pseudohyphal development (*STE20, STE11, STE7,* and *STE12*) are also required for both filament formation and agar penetration in haploids. Thus, haploid yeast cells can enter either of two developmental pathways: mating or invasive growth, both of which depend on elements of a single MAP kinase cascade. Our results provide a novel developmental model to study the dynamics of signal transduction, with implications for higher eukaryotes.

**Key Words:** Signal transduction, MAP kinase; fungal dimorphism; budding

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Mitogen-activated protein kinase (MAPK or ERK) signal transduction pathways are critical for many developmental events in eukaryotes, and require the sequential activation and phosphorylation of a series of protein kinases (for review, see Blumer and Johnson 1994). Activation of MAPK requires phosphorylation by activated MAPK/ERK kinase (MEK or MAPKK) (Crews et al. 1992). MEK activation is achieved by phosphorylation by RAF1 or MEK kinase (MEKK or MAPKKK; Kyriakis et al. 1992; Lange-Carter et al. 1993). Homologs of each of these components have been identified in a large number of eukaryotes. In the nematode *Caenorhabditis elegans*, vulval induction requires signaling through the MAPK homolog *Sur1/Mpk1* (Lackner et al. 1994; Wu and Han 1994). In *Drosophila melanogaster*, the MAPK homolog *rolled* is critical for photoreceptor development and dorsoventral patterning during embryogenesis (Brunner et al. 1994). In mammalian cells, MAPKs are important in a wide array of cytokine and growth factor responses (Pazin and Williams 1992; Schlessinger and Ullrich 1992; Foster 1993).

In the yeast *Saccharomyces cerevisiae*, several independent MAP kinase pathways have been characterized. One pathway mediates the response to hyperosmotic conditions through the MAP kinase pathway homologs *PBS2* [MEK] and *HOG1* [MAPK], which stimulate glycerol accumulation (Brewster et al. 1993). A second pathway controls cell wall biosynthesis through *PKC1* [a protein kinase C homolog] and the protein kinases *BCK1* [MEKK], *MKK1* and *MKK2* [MEK], and *MPK1* [MAPK] (Blumer and Johnson 1994). The best defined MAP kinase pathway controls the mating of haploid cells through *STE11* [MEKK], *STE7* [MEK], and *FUS3* and *KSS1* [MAPK).

In *S. cerevisiae* haploid cells, mating is triggered by pheromones and leads to the fusion of two cells of opposite mating type to form a diploid. The primary signaling event in this pathway, the binding of the peptide pheromone to its cognate receptor [Ste2p in MATα cells and Ste3p in MATα cells], results in GDP/GTP exchange on the α subunit [Gpa1p] of a heterotrimeric G protein [Dietzel and Kurjan 1987]. The β [Ste4p] and γ [Ste18p] subunits then stimulate the activation of a series of protein kinases [Whiteway et al. 1989]. Ste20p is proposed to link the G protein to the kinase cascade [Leberer et al. 1992; Ramer and Davis 1993]. The MEKK homolog Ste11p [Rhodes et al. 1990; Stevenson et al. 1992] and the MEK homolog Ste7p are sequentially activated [Cairns et al. 1992], and the latter phosphorylates the MAPK homologs Fus3p and Kss1p [Gartner et al. 1992; Zhou et al. 1994].
phosphorylates the cyclin-dependent kinase inhibitor transcription factor Stel2p (Elion et al. 1993). Fus3p also the assembly of the Stellp, Ste7p, and Fus3p protein kinases into a complex (Choi et al. 1994, Kranz et al. 1994, Marcus et al. 1994), which phosphorylates the transcription factor Ste12p [Elion et al. 1993]. Fus3p also phosphorylates the cyclin-dependent kinase inhibitor Far1p [Elion et al. 1993], which mediates cell cycle arrest (Chang and Herskowitz 1990). Ste12p activates the transcription of mating-specific genes including FUS1 [Do- lan and Fields 1990] and genes linked to the transposon Ty [Van Arsdell et al. 1987; Ciriacy et al. 1991]. Thus, activation of the MAP kinase pathway in two haploid cells of opposite mating type arrests vegetative growth and leads to the fusion of the two cells to form a diploid.

Elements of the mating pheromone response MAP kinase pathway are also essential for pseudohyphal development in diploid strains [Liu et al. 1993]. When starved for nitrogen, diploid cells undergo a developmental transition from a single cell yeast form to a filamentous pseudohyphal form. Pseudohyphal filaments are composed of chains of long thin cells that radiate away from the colony and penetrate the agar substrate on which they are grown [Gimeno et al. 1992]. Mutations in STE20, STE11, STE7, or STE12 block the filament formation, cell elongation, and agar penetration of pseudohyphal cells. However, mutations in the pheromone receptors and G protein do not affect pseudohyphal development, suggesting that other molecules generate and transduce the signal that triggers filamentous growth [Liu et al. 1993].

The finding that both mating and filamentous growth use elements of the same MAP kinase cascade raises the question of how the same signal transduction pathway can specify two different developmental sequences. Specifically, the Ste12p transcription factor must activate mating-specific genes in the haploid and pseudohyphal-specific genes in the diploid. In one model, stimulation of the MAP kinase cascade in haploid MATa or MATa cells results in the activation of mating-specific events, whereas stimulation of this pathway in diploid MATa/α cells results in pseudohyphal development. According to this model, factors specific to each cell type determine the ensemble of genes that are regulated by Ste12p. For example, α1 and α2 may repress the activation of mating-specific genes, such as FUS1, in the MATa/α diploid cell, allowing the induction of genes specific to filamentous growth under conditions of nitrogen starvation.

In addition to elements of the mating MAP kinase cascade, filament formation also requires a series of polarized cell divisions determined by the site of bud emergence of each new cell in the growing chain. Diploid cells bud in the bipolar mode, where the first bud of a virgin mother emerges from the free end of the cell and subsequent buds emerge from either the birth or free end of the cell [Freifelder 1960]. The birth end is defined as the region where each cell is attached to its mother; the opposite end is the free end [Freifelder 1960]. Pseudohyphal cells bud in a unipolar mode rather than the bipolar mode typical of diploid yeast form cells [Kron et al. 1994]. In the unipolar mode, the first bud and all subsequent buds emerge from the free end of the cell. Successive rounds of unipolar budding produce a chain of first daughters oriented away from the cell that initiated the chain. In contrast, most haploid cells bud in an axial mode of bud site selection and do not form filaments readily when starved for nitrogen [Gimeno et al. 1992]. In the axial mode, each new bud emerges from the birth end of a cell [Freifelder 1960]. Successive rounds of axial budding produce a cluster of cells rather than the ordered chain of cells required to form a filament [Gimeno and Fink 1992]. The bud site selection mode is determined by a regulatory cascade of BUD genes [Chant and Herskowitz 1991]. A mutation in BUD1 (RSR1) disrupts the bipolar budding pattern of diploids [Bender and Pringle 1989] and drastically reduces their ability to form filaments [Gimeno et al. 1992].

In this paper we describe a haploid invasive growth behavior with several similarities to diploid pseudohyphal development. Specifically, haploids switch their pattern of bud site selection from an axial to a bipolar pattern and consequently form filaments. In addition, they penetrate the agar substrate on which they are grown. Haploid filament formation and agar penetration depend on the same pheromone response MAP kinase cascade components required for diploid pseudohyphal growth. Thus, in a single cell the same signal transduction pathway can mediate two distinct developmental sequences showing that the differential outputs of the pathway do not require cell type-specific factors.

**Results**

**Haploid cells are capable of agar penetration and filament formation on rich medium**

**MATa** and **MATa** haploid strains of the Σ1278b background grown on rich medium plates (YPD) for several days begin to penetrate the agar beneath each colony [Fig. 1; Table 1]. A population of cells that have penetrated the agar can be observed by washing the top portion of the colony off of the agar surface with water. The remaining cells are judged to have penetrated the agar surface because they are inaccessible to a microdissection needle unless the needle pierces the agar. After 3 days of growth, when the mass of cells on top of the agar has already slowed its growth, the number of cells that have penetrated the agar increases steadily, suggesting that agar penetration may be triggered by nutrient limitation. However, haploid agar penetration may not be related to ammonia deprivation because the addition of ammonium sulfate to YPD does not diminish agar penetration. Remarkably, isogenic **MATa/α** diploid cells do not penetrate agar efficiently under these conditions, except after extremely long incubations, and the few cells that remain in the agar have the yeast form [Fig. 1].

The haploid cells remaining in the agar after washing the plate form networks of filaments underneath each colony [Fig. 2]. The cells of these haploid filaments are not as elongated as those observed in diploid filaments,
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Figure 1. Invasive growth of haploid strains. Haploid MATα (L5487) and MATa (L5528) strains were patched on YPD with an isogenic diploid MATα/a strain (L5721) and incubated as described in Materials and methods. The plate was photographed before (total growth) and after (invasive growth) washing the cells off of the agar surface.

although some may have a slightly greater length-to-width ratio than they do during exponential growth (1.43±0.02 as compared with 1.07±0.01) (mean±S.E.M.). Even after prolonged incubations, these haploid filaments do not extend beyond the perimeter of the colony but are restricted to the agar immediately beneath it. Thus, we define the combination of filament formation and agar penetration that occurs underneath each colony on rich medium as haploid invasive growth.

Bud site switching during haploid invasive growth

Haploid filament formation is unexpected because haploid cells bud in the axial mode, a pattern inconsistent with filament formation. Even if the cells fail to separate after cytokinesis, reiteration of this budding pattern produces clumps of cells, not the sequential extensions required for filament formation (Fig. 2). To determine the growth pattern of our strains, we compared the budding pattern of cells during exponential and invasive growth by staining their bud scars with calcofluor, a fluorescent compound that binds the chitin ring at the mother-daughter junction.

Cells were divided into three classes based on the distribution of their bud scars: axial, bipolar, and anomalous (see Materials and methods). Haploid cells in exponential phase either on plates or in liquid bud in a very strict axial mode, where all the bud scars are clustered together at one end of the cell (Fig. 3, Table 2). Even after extended growth periods in liquid medium, these cells continue to bud in an axial mode (Table 3). Remarkably,

| Strain | Genotype | Source |
|--------|-----------|--------|
| L5528  | MATα, ura3-52, his3::hisG | this study |
| L5487  | MATα, ura3-52, leu2::hisG | this study |
| L5466  | MATα, ste2::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5549  | MATα, ste4::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5579  | MATα, ste18::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5585  | MATα, ste20::TRP1, ura3-52, trp1::hisG | Liu et al. (1994) |
| L5554  | MATα, ste5::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5565  | MATα, ste11::URA3::TRP1, ura3-52, trp1::hisG | Liu et al. (1994) |
| L5559  | MATα, ste7::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5743  | MATα, fus3::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5491  | MATα, kss1::URA3::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5724  | MATα, fus3::LEU2, kss1::URA3::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5723  | fus3::LEU2, ste7::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5724  | fus3::LEU2, ste12::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5573  | MATα, ste12::LEU2, ura3-52, leu2::hisG | Liu et al. (1994) |
| L5742  | MATα, far1::URA3, ura3-52, his3::hisG | Liu et al. (1994) |
| L5721  | MATα/a, ura3-52, ura3-52, his3::hisG/HIS3, leu2::hisG/LEU2 | this study |
| L5729  | MATα/a, bud1::URA3/bud1::URA3 ura3-52/ura3-52, his3::hisG/HIS3, leu2::hisG/LEU2 | this study |
| L5741  | MATα/a, ste12::LEU2/STE12, leu2::hisG/leu2::hisG | this study |
| L5726  | MATα, bud1::URA3, ura3-52, his3::hisG | this study |
| L5727  | MATα, bud2::LEU2, ura3-52, leu2::hisG | this study |
| L5725  | MATα, bud3::URA3, ura3-52, his3::hisG | this study |
| F1432  | MATα, bud4, ura3-52 | Gimeno and Fink (1994) |
| L5740  | MATα, bud5::URA3, ura3-52, his3::hisG | this study |
| L5728  | ura3-52, his3::hisG, ura3::FUS1-lacZ::URA3 | this study |

All strains are congenic to the Σ1278b genetic background (Grenson et al. 1966; Liu et al. 1993).
invasive haploid cells from washed plates exhibit a bipolar pattern of bud scar distribution, with scars clustered at both ends of the cell. The frequency and appearance of these bipolar haploids is comparable to that of diploids in the yeast form [Fig. 3; Table 2]. Cells in the population that do not manifest the bipolar pattern could be either cells in the axial mode that have not switched or cells that have switched but are in the unipolar pattern rather than the bipolar pattern (Kron et al. 1994). Because the distribution of bud scars is the same for cells dividing in the axial and unipolar patterns, these two modes could not be distinguished by this method.

The finding that haploid cells can switch from axial to bipolar budding on plates suggested that this switch is critical for filament formation. To determine directly whether the cells in haploid filaments have the diploid cell division motif, we recorded the growth of filaments in the plate by time-lapse photography. In a high proportion of the cells in haploid filaments (52/70), the first bud of a virgin mother emerges from the free end of the cell. This pattern, typical of bipolar budding in diploid strains, permits the formation of filaments through a series of polarized cell divisions. The bipolar pattern contrasts with that of haploids in exponential growth, where all new buds emerge from the birth end of the cell (78/78). The bipolar cells are not diploids because, when excised from the agar, they give rise to cells capable of mating with cells of the appropriate mating type. Thus, haploids grown on plates can switch their budding pattern from an axial to a bipolar mode and form filaments. This switch demonstrates that bud site selection is not a fixed parameter determined by cell type but, rather, a dynamic process sensitive to environmental influences.

Table 2. Haploid bipolar budding

| Sample               | axial | bipolar | anomalous | n   |
|----------------------|-------|---------|-----------|-----|
| MATα exponential     | 98    | 1       | 1         | 500 |
| MATα below agar      | 30    | 62      | 8         | 800 |
| MATα/a exponential   | 28    | 66      | 6         | 200 |

The distribution of bud scars was scored for the haploid strain L5528 during exponential and invasive growth on plates, and for the diploid strain L5721 during exponential growth in liquid. Numbers represent the percentage of cells in each class (see Materials and methods) and the sample size [n].
Three gene products, BUD1, BUD2, and BUD5, are required for both the axial and bipolar budding patterns. Haploid bud1, bud2, and bud5 mutant strains exhibit a random distribution of bud emergence sites on the cell surface. Unlike isogenic Bud+ strains, these mutants do not switch to the bipolar budding pattern (Table 4) or form filaments in our plate assay [data not shown]. Thus, a bipolar budding pattern is necessary for filament formation.

### Agar penetration does not require BUD gene functions

Haploid bud1, bud2, and bud5 mutants fail to form filaments on rich medium, but they penetrate agar as well as Bud+ strains [Fig. 4]. Therefore, a polar mode of bud site selection is not necessary for agar penetration, although it is necessary for filament formation. To test whether BUD genes are also required for diploid filamentous growth, we constructed diploid strains homozygous for a deletion of BUD1. These strains bud randomly and fail to form filaments on plates (SLAD) that induce pseudohyphal growth in diploid Bud+ strains, although they form normal proportions of long cells that penetrate the agar below each colony [data not shown]. On rich medium used to observe haploid invasive growth (YPD), diploid Bud+ strains do not penetrate agar efficiently [Figs. 1 and 4], although they exhibit the bipolar budding pattern necessary for filamentous growth. Taken together these data suggest that filament formation and agar penetration are distinct functions in both haploids and diploids. Thus, a polar mode of bud site selection is neither necessary nor sufficient for agar penetration; other factors must regulate this behavior.

### Elements of the pheromone response pathway are required for haploid invasive growth

Because mutations in the mating signal transduction pathway block both the filament formation and the agar penetration of diploid strains, we reasoned that this pathway might control these behaviors in haploids as well. Haploid strains carrying deletions in various pheromone response pathway genes were tested for their ability to penetrate agar. Strikingly, the same components of the MAPK cascade necessary for diploid pseudohyphal growth are required for haploid invasive growth. As in diploids, the pheromone receptor (STE2) and the G protein (STE4, STE18) are not required for haploid agar penetration. Similarly, the scaffold protein (STE5) that nucleates assembly of the signaling kinases is not required nor is the cyclin-dependent kinase inhibitor (FAR1) [Fig. 5; data not shown]. In contrast, other components of the MAP kinase cascade, including STE20, STE11 (MEKK), STE7 (MEK), and the transcription factor STE12, are essential for agar penetration (Fig. 5). As in diploids (Liu et al. 1993), the ste20 mutant shows the strongest phenotype of all the mutants in this pathway. In each case, the defect in the invasive growth of these strains is completely linked to the ste deletion (Fig. 6), and agar penetration can be restored by introduction of the appropriate STE gene. It is difficult to assess whether the noninvasive ste mutants are completely defective in filament formation because our assay requires examination of the cells remaining in the agar. Nevertheless, the few cells that do remain after washing the plate do not appear to form filaments [data not shown]. Therefore, we conclude that these mutant strains are defective in agar penetration and probably also filament formation.

### A MAPK is required for haploid invasive growth

Haploid strains containing deletions of the Fus3p and Kss1p MAPK homologs were tested for their ability to

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**Table 3. Haploids do not switch budding patterns in liquid**

| Sample         | Bud scar distribution | n  |
|----------------|-----------------------|----|
|                | axial | bipolar | anomalous |    |
| MATa exponential| 98    | 1.0     | 1.0       | 400 |
| MATa 2 days    | 80    | 15      | 4.3       | 391 |
| MATa 20 days   | 85    | 8.0     | 7.0       | 100 |

The distribution of bud scars was scored for the haploid strain L5528 after growth in YPD liquid for various amounts of time. Numbers represent the percentage of cells in each class (see Materials and methods) and the sample size (n).

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**Table 4. The budding pattern switch requires BUD genes**

| Sample | exponential growth | invasive growth |
|--------|-------------------|----------------|
|        | axial | bipolar | anomalous | axial | bipolar | anomalous |
| Bud+   | 98    | 1.5     | 0.0       | 26    | 66      | 8.0       |
| bud1   | 14    | 14      | 72        | 9.0   | 20      | 72        |
| bud2   | 12    | 13      | 75        | 2.0   | 21      | 77        |
| bud3   | 21    | 64      | 15        | 26    | 59      | 15        |
| bud4   | 23    | 65      | 12        | 20    | 65      | 15        |

The distribution of bud scars was scored for haploid strains [Bud+ L5528], [bud1 L5726], [bud2 L5727], [bud3 L5725], and [bud4 F1432] during exponential growth in liquid and invasive growth on plates. Numbers represent the percentage of cells in each class (see Materials and methods) for a sample size of n = 200.
penetrate agar. These genes are partially redundant for an essential mating function (Elion et al. 1991a,b) but are not required for filament formation in diploids (Liu et al. 1993). Haploid kss1 strains show greatly reduced agar penetration [Fig. 7], demonstrating that Kss1p promotes haploid invasive growth. In contrast, fus3 strains penetrate agar more vigorously than FUS3 strains, suggesting that Fus3p inhibits haploid invasive growth [Fig. 7]. Surprisingly, the double mutant fus3 kss1 penetrates agar as well as the wild-type FUS3 KSS1 strain. Because the fus3 kss1 double mutant penetrates agar better than the kss1 single mutant, we hypothesized that there might be a novel MAPK, partially redundant with Kss1p, that is inhibited by the presence of a functional FUS3 gene. To test whether the enhanced agar penetration of the fus3 mutant strain is dependent on a functional signal transduction pathway, we analyzed fus3 ste7 and fus3 ste12 mutants. These double mutant strains, like the ste7 and ste12 single mutant strains, fail to penetrate agar [Fig. 7]. We tested deletions of the MAPK homologs HOG1 and MPK1 and found that these kinases do not play a role in agar penetration (data not shown). Taken together these data support a positive role for Kss1p in haploid invasive growth and raise the possibility of a heretofore unidentified MAPK.

**STE pathway components are not required for bud site switching**

To determine whether the pheromone response MAP kinase pathway controls the bud site switching essential for haploid filament formation, we examined the pattern of bud site selection in ste strains. Because certain ste mutants penetrate agar poorly, we analyzed the budding pattern of cells collected from colonies on the agar surface. All the ste mutants in the pheromone response pathway still switch their budding pattern from an axial to a bipolar mode when grown for several days on YPD plates [Table 5]. Although the proportion of cells that switch from axial to bipolar budding is not as great for the cells on the agar surface as for those below the surface [cf. Tables 2 and 5], it is clear that the ste mutants retain the capacity to switch to the bipolar pattern.

**Mating-specific transcription is not induced during haploid invasive growth**

Because haploid cells express all of the genes required for induction of pheromone-responsive genes, it is possible that mating-specific genes, such as FUS1, are activated in the cells under the colony during the induction of haploid invasive growth. To test this possibility, we compared the expression levels of a FUS1-lacZ construct in haploid cells that had penetrated the agar or were cultured in the presence of mating pheromone. This reporter provides a sensitive assessment of the transcriptional induction of mating-specific genes. Transcription of FUS1-lacZ is induced 100-fold during the pheromone response, and this induction is mediated by the transcription factor Ste12p (Dolan and Fields 1990; Hagen et al. 1991). Our data show that there is no induction of FUS1 in cells above or below the agar in the absence of mating pheromone [Table 6]. Thus, the signaling cascade and the transcription factor Ste12p pre-
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YPD Invasion -Leu

Figure 6. Invasive growth is linked to STE12. The diploid strain (ste12::LEU2/STE12, leu2/leu2 L5741) was sporulated and subjected to tetrad analysis. Spores were assayed for growth on YPD (total growth), agar penetration (invasive growth), and the ability to grow on medium lacking leucine (growth on -Leu).

sumably induce a set of genes specific to haploid invasive growth that is distinct from pheromone-inducible genes.

Discussion

A comparison of haploid invasive growth and diploid pseudohyphal growth

Haploid invasive growth is a developmental program with many similarities to diploid pseudohyphal growth. First, in both haploids and diploids, filament formation and agar penetration appear to be induced by nutrient limitation. Second, both cell types form filaments through a series of polarized cell divisions that depend on a polar pattern of bud site selection. Furthermore, filament formation is distinct from agar penetration in both cell types, as both haploid and diploid bud strains are unable to form filaments but still penetrate agar. Perhaps the most striking similarity is that elements of the pheromone response MAP kinase signal transduction pathway, including STE20, STE11, STE7, and STE12, are required for triggering these developmental events in both haploids and diploids.

However, there are differences between haploid invasive growth and diploid pseudohyphal growth. Diploids form filaments readily on low ammonia medium, but poorly on rich medium. In contrast, haploids do not appear to form filaments on low ammonia medium, but do form them on rich medium. Moreover, diploid filaments extend beyond the perimeter of the colony, whereas haploid filaments are only observed beneath the colony. In addition, cells in diploid pseudohyphal filaments are more elongated than those in haploid filaments. Despite these differences, haploid invasive growth and diploid pseudohyphal growth share several common motifs and require many of the same genes, suggesting that they are different manifestations of a common developmental pathway.

Haploid filamentous growth requires bud site switching

Bud site selection is not a fixed parameter determined by cell type but, rather, a dynamic process sensitive to environmental influences. Haploid cells switch from an axial to a bipolar mode of bud site selection when grown on rich medium for an extended period. In the bipolar mode, the first bud emerges from the free end of the cell, but subsequent buds can emerge from either end. Diploid cells switch from a bipolar to a unipolar budding pattern when starved for nitrogen (Kron et al. 1994). In the unipolar budding pattern, the first bud and all subsequent buds emerge from the free end of the cell. Thus, upon nutrient deprivation, both diploid and haploid cells switch their pattern of bud site selection and form filaments.

The polar budding pattern provides the architectural motif for the growth and extension of a filament. In polar budding the first bud of each virgin mother emerges from the free end of the cell, away from her mother. The switch in diploids from bipolar to unipolar budding ensures that all new growth projects in filaments away from the colony. The switch in haploids from axial to bipolar budding is also consistent with filament formation. Both the bipolar chain and the unipolar chain are composed of a lineage of buds born from the free end of each new cell in the chain. However, in the bipolar pattern the emergence of subsequent buds on the birth end of cells can lead to secondary fronds projecting back toward the cell that originated the chain. This difference may explain why the haploid filaments we observe are
Filament formation is distinct from agar penetration

The distribution of bud scars was scored for haploid MATa strains [Ste+ L5528], [ste2 L5466], [ste4 L5549], [ste18 L5579], [ste20 L5585], [ste5 L5554], [ste11 L5565], [ste7 L5559], [fus3 L5479], [kssl L5491], [fus3 kssl L5722], and [ste12 L5573] after 5 days of growth on a YPD plate. Numbers represent the percentage of cells in each class (see Materials and methods) for a sample size of n = 200.

| Sample      | axial | bipolar | anomalous |
|-------------|-------|---------|-----------|
| Ste+        | 53    | 42      | 5         |
| ste2        | 45    | 46      | 9         |
| ste4        | 43    | 50      | 7         |
| ste18       | 45    | 47      | 8         |
| ste20       | 46    | 48      | 6         |
| ste5        | 43    | 53      | 4         |
| ste11       | 59    | 37      | 4         |
| ste7        | 51    | 43      | 6         |
| fus3        | 50    | 40      | 4         |
| kssl        | 46    | 48      | 6         |
| fus3, kssl1 | 50    | 42      | 8         |
| ste12       | 50    | 42      | 8         |

The role of MAPKs in haploid invasive growth

The FUS3 and KSS1 MAPKs are partially redundant for an essential function in the haploid pheromone response (Ellen et al. 1991a,b). The single mutants are fertile, but the fusi kssl double mutant is completely sterile. The FUS3 kssl single mutant is defective in agar penetration, whereas the fusi KSS1 mutant is enhanced. The role of the FUS3 and KSS1 genes in haploid invasive growth is unexpected because these genes are not required for diploid pseudohyphal growth. Formally, Kssl1p plays a positive role in haploid invasive growth and Fus3p plays a negative role. Epistasis analysis shows that the enhanced agar penetration of the fusi mutant depends on the function of two other components of this pathway, STE7 and STE12.

These data raise the possibility that Kssl1p may be partially redundant with a novel MAPK in the invasive growth signaling pathway; either can activate Ste12p to promote invasive growth. The presence of Fus3p might interfere with the function of the postulated MAPK.

Table 5. Bud site switching does not require STE genes

| Sample | axial | bipolar | anomalous |
|--------|-------|---------|-----------|
| Ste+   | 53    | 42      | 5         |
| ste2   | 45    | 46      | 9         |
| ste4   | 43    | 50      | 7         |
| ste18  | 45    | 47      | 8         |
| ste20  | 46    | 48      | 6         |
| ste5   | 43    | 53      | 4         |
| ste11  | 59    | 37      | 4         |
| ste7   | 51    | 43      | 6         |
| fus3   | 50    | 40      | 4         |
| kssl   | 46    | 48      | 6         |
| fus3, kssl1 | 50 | 42 | 8 |
| ste12  | 50    | 42      | 8         |

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Table 6. FUS1-lacZ is not induced during haploid invasive growth

| Mating transcription induction | β-galactosidase activity |
|-------------------------------|-------------------------|
| sample                        |                         |
| MATa exponential growth + αF  | 250                     |
| MATa exponential growth        | 2.2                     |
| MATa on agar surface           | 0.40                    |
| MATa beneath agar surface      | 0.39                    |

β-Galactosidase activity was assayed in cells of the haploid MATa strain L5728 during exponential growth in YPD liquid in the presence and absence of mating pheromone, and after 5 days of growth on a YPD plate. Numbers represent β-galactosidase activity normalized to total protein (see Materials and methods).
thereby reducing the invasive growth signal. This model explains the strength of the haploid invasion phenotypes that we observe: strongest in fus3 KSS1, intermediate in FUS3 KSS1 and fus3 kss1, and weakest in FUS3 kss1. The postulated MAPK is unlikely to be either MPK1 or HOG1, two other MAPKs present in yeast, because deletions of these genes do not block haploid invasive growth. Alternatively, there may not be a third MAPK involved in haploid invasive growth. Instead, Kss1p could promote, and Fus3p antagonize, the Ste7p-dependent Ste12p activation required for haploid invasive growth.

Elements of a single MAP kinase pathway control two developmental events in the same cell type

Our finding that components of the pheromone response signal transduction pathway are required for both mating and haploid invasive growth demonstrates that a single MAP kinase cascade can specify two different developmental outcomes in a haploid cell. The target of this cascade, the transcription factor Ste12p, is required for both processes, presumably to regulate the expression of genes specific to each developmental program. Because mating-specific genes are not induced detectably during haploid invasive growth, Ste12p must have at least two functional states: one that controls the transcription of mating-specific genes, and another that controls the transcription of genes specific to haploid invasive growth. Each Ste12p state might have a distinct phosphorylation pattern that determines DNA-binding specificity, association with other transcription factors, or both.

How do haploid cells distinguish between two different stimuli that activate the same signal transduction pathway and initiate the developmental program specific to each primary signal? In one model, accessory proteins are activated that direct the association of common components of the pathway with their appropriate targets (Fig. 8). For example, in the response to mating pheromone, Ste5p assembles Ste11p, Ste7p, and Fus3p into a protein complex that modifies Ste12p, shifting it to a state that activates the transcription of mating-specific genes (Choi et al. 1994; Kranz et al. 1994; Marcus et al. 1994). According to this model, upon induction of the haploid invasive growth pathway, a scaffolding protein analogous to Ste5p mediates the assembly of Ste11p and Ste7p with a MAPK distinct from Fus3p. The protein complex modifies Ste12p, shifting it to a state that activates the transcription of genes specific to haploid invasive growth (Fig. 8). This explains why the pheromone receptors, G protein, FUS3, and STE5 are not necessary for haploid invasive growth, whereas STE20, STE7, STE11, and STE12 are essential. Other models that do not require a MAPK are consistent with these data.

A paradigm for generating diversity from a signal transduction pathway

Our observation that elements of a single MAP kinase pathway control two distinct developmental sequences could emerge as a general theme. MAP kinase pathways have been implicated in development and differentiation in many organisms. In mammalian cells, a wide array of growth factors and cytokines can activate a MAP/ERK kinase cascade that stimulates proliferation or differentiation (Blumer and Johnson 1994). In D. melanogaster, a dominant activated allele of a single MAPK gene disrupts the development of several tissues (Brunner et al. 1994), although in this case, each of these processes occurs in distinct cell types at different times during development.

There are precedents for two distinct signals acting through a single MAP kinase pathway in the same cell type. For example, in cardiac myocytes both endothelin [ET-1] and acidic fibroblast growth factor [aFGF] activate a MAPK, and induce hypertrophy in these cells (Bogoyevitch et al. 1994). ET-1 acts through G protein-coupled serpentine receptors and phospholipase Cβ, whereas aFGF activation is mediated by a family of tyrosine kinase receptors. These two agonists act through disparate primary signaling events, but both pathways converge on the MAPK to mediate the same physiologic response. Perhaps the closest parallel to the dual functionality of the MAP kinase pathway in S. cerevisiae is the behavior of PC12 cells in response to epidermal...
growth factor (EGF) and nerve growth factor (NGF) (Traverse et al. 1992). EGF induces transient ERK/MAPK activation and cell division in these cells. In contrast, NGF induces prolonged activation and nuclear translocation of ERK/MAPK, causing the same cells to arrest growth and initiate neuronal differentiation. The molecular sequence of events underlying this differential MAPK activation has not yet been determined. Identification of the primary signals and the putative novel MAPK required for yeast filamentous growth should help to elucidate the mechanism for achieving diversity from a single cascade.

Materials and methods

Strains and growth conditions

All yeast strains used in this study are described in Table 1 and are congenic to the Σ1278b genetic background (Grenson et al. 1966; Liu et al. 1993). Standard yeast culture medium was prepared essentially as described (Sherman et al. 1986). YPD plates (2% yeast extract (Difco), 4% Bacto peptone (Difco), 2% glucose, 2% Bacto-agar (Difco), and 0.03% L-tryptophan) were allowed to cool, polymerize, and dry for 3 days at room temperature before use. Low ammonia medium (SLAD) for scoring pseudohyphal growth was prepared as described (Gimeno and Fink 1994). Strains scored for pseudohyphal filament formation were streaked on SLAD plates and observed after 24, 48, and 72 hr of growth at 30°C. Long cells that had penetrated the agar were examined after washing the cells off of the agar surface with deionized water.

Cells in exponential growth phase in liquid were prepared by incubating cells in YPD medium on a roller drum at 30°C until the culture reached an OD_{600} of 0.6. Cells in exponential growth phase on plates were prepared by plating 3 ml of cells at an OD_{600} of 0.6 (see above) on a YPD plate and incubating them for an additional 8–10 hr at 30°C.

Invasive growth assay

Strains were patched on YPD plates with a toothpick, with care being taken to avoid scratching the agar surface. Patches were allowed to grow at 30°C for 3 days and incubated at room temperature for an additional 2 days. For tetrad analysis, haploid cells were transferred from liquid cultures in a microtiter dish to a YPD plate. Because these samples started at a higher cell density than the patches described previously, invasive growth was scored after 2 days of growth at 30°C. Samples were photographed with a 35mm camera using Technical Pan (Kodak) film, and a gentle stream of deionized water was then used to rinse all the cells from the agar surface. The agar was never rubbed directly or otherwise disrupted. Microscopic analysis revealed that the cells remaining after washing were below the focal plane of the agar surface and inaccessible to a microdissection needle unless the needle pierced the agar. Plates were allowed to dry briefly and were then photographed again as described above. Cells that had penetrated the agar were excised with a toothpick for subsequent manipulations.

Filament formation was scored after 2 days of invasive growth at 30°C because the number of cells in the agar after 5 days was too high to distinguish individual filaments. Cells remaining in the agar after washing the plate were overlayed with a coverslip and scored by direct microscopic examination for the presence of filaments. For the purpose of this study, a filament is defined as a chain of cells attached to one another that projects in an ordered pattern away from the first cell of the chain.

Photomicroscopy

Preparations of cells in exponential growth were prepared as described above, placed on a slide with a coverslip, and compressed gently to ensure that all the cells in each clump were in the same focal plane. Cells undergoing invasive growth were cultured as described above and prepared for observation by excising the washed agar from the plate and placing it between two coverslips. Samples were viewed with Nomarski optics on a Zeiss Axioscop and photographed with a 35mm camera using Technical Pan (Kodak) film. To determine the axial ratio of cells, we prepared samples and photographed them as described above. Images were printed onto photographic paper and directly measured. The length was measured along the longest axis of each cell, and the width was measured at the midpoint of the longest axis (Gimeno et al. 1992). The mean length/width ratios are presented for each sample along with the standard error of the mean.

Bud scar staining

Invasive cells were prepared as described above and excised from washed plates with a toothpick. Cells in exponential growth phase were prepared as described above. Samples were forced through a 24-gauge needle 10 times to separate cells as much as possible and rinsed in water once before fixation. Cell suspensions were fixed at room temperature for 2 hr in 3.7% formaldehyde. Samples were rinsed twice in water and resuspended in 200 μl of a fresh stock of 1 mg/ml of calcifluor white (Fluorescent Brightener no. 28 F6259, Sigma) in water. Samples were stained at room temperature in the dark for 10 min and washed three times in water before observation.

Bud scars were visualized by fluorescence microscopy on a Zeiss Axioscop, and photographed with a 35mm camera using TMAX 400 (Kodak). Cells with 2–10 clear bud scars were divided into three bud scar distribution classes: axial, cells with all bud scars at one end of the cell immediately adjacent to one another; bipolar, cells with two or more bud scars with at least one scar at each end of the cell (the birth end and the free end); anomalous, cells with bud scar distributions other than axial or bipolar. Numbers in the tables represent the percent of cells in each class for a sample of n cells.

Time-lapse photomicroscopy

A plate of invading cells was prepared as described above, and a coverslip was placed over the cells remaining in the agar. Each plate was taped down on a Zeiss WL light microscope stage, maintained at 30°C, and photographed at 90-min intervals over a 14-hr period with a 35mm camera using Technical Pan film (Kodak). Budding was observed and classified according to the site where the first bud of a virgin mother emerged. None of the cells present on the plate at time zero was scored, as their birth end could not always be determined. Instead, buds that emerged after the first time point and subsequently initiated buds of their own were defined as virgin mothers. The birth end of a virgin mother was defined as the region adjacent to the site where she was attached to her mother, and the free end was defined as the end of the cell opposite the birth end (Freifelder 1960). The position of bud site emergence of these virgin mothers was determined by direct microscopic observation.

Exponential growth samples were prepared as described above and subjected to time-lapse analysis. In these preparations we
find that mother and daughter cells remain attached to one another even after being subjected to high shear forces. This site of attachment defines the birth end of each cell [Freifelder 1960]. However, because the majority of these cells are in large clumps, we could not always identify the first bud of a virgin mother and simply scored new buds as emerging from the free end or the birth end of the cell.

β-Galactosidase assays

β-Galactosidase assays were performed essentially as described [Trueheart et al. 1987]. Exponential growth phase samples were prepared as described above and split into two samples. These samples were spun, resuspended in 5 ml of YPD (pH 4) with or without 5 μM α-factor, and incubated at 30°C for an additional 2 hr before being harvested. Cells undergoing invasive growth were prepared as described above. Before washing the plate, cells on the agar surface were gently scraped away and harvested. Separate plates grown under the same conditions were washed on the agar surface were gently scraped away and harvested. All cells were washed with water once before being lysed to remove contaminating protein. β-Galactosidase activity was normalized to the total protein in each extract according to the equation: (OD420 x 1.7)/(0.0045 x mg/ml protein x extract volume x time).

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