Spatial focalization of pheromone/MAPK signaling triggers commitment to cell–cell fusion

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Cell fusion is universal in eukaryotes for fertilization and development, but what signals this process is unknown. Here, we show in Schizosaccharomyces pombe that fusion does not require a dedicated signal but is triggered by spatial focalization of the same pheromone–GPCR (G-protein-coupled receptor)–MAPK signaling cascade that drives earlier mating events. Autocrine cells expressing the receptor for their own pheromone trigger fusion attempts independently of cell–cell contact by concentrating pheromone release at the fusion focus, a dynamic actin aster underlying the secretion of cell wall hydrolases. Pheromone receptor and MAPK cascade are similarly enriched at the fusion focus, concomitant with fusion commitment in wild-type mating pairs. This focalization promotes cell fusion by immobilizing the fusion focus, thus driving local cell wall dissolution. We propose that fusion commitment is imposed by a local increase in MAPK concentration at the fusion focus, driven by a positive feedback between fusion focus formation and focalization of pheromone release and perception.

[Keywords: cell–cell fusion; fertilization; MAP kinase cascade; G-protein-coupled receptor signaling; pheromone; formin; fission yeast Schizosaccharomyces pombe]

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Cell fusion is a widespread process that occurs in a large variety of cell types. In somatic cells, it serves to shape tissues and organs such as muscles, bones, or the placenta in metazoans or form interconnected mycelia in fungi (Read et al. 2010). In gametes, it underlies the fertilization of all sexually reproducing species. In pathogenic fungi, this process is often tightly linked with virulence (Morrow and Fraser 2009; Ene and Bennett 2014). Cell fusion may also play important roles in regeneration and cancer (Lu and Kang 2009; Lluis and Cosma 2010). In all events of cell–cell fusion, three major conceptual steps can be defined: First, signaling occurs between the two partner cells to induce cell differentiation. Second, the cells polarize toward each other for cell–cell adhesion. Third, a fusion machinery is assembled at the cell–cell contact site for membrane merging. This machinery may be composed of two main parts: a specific actin structure that promotes membrane juxtaposition (Abmayr and Pavlath 2012) and fusogenic proteins that drive membrane fusion (Aguilar et al. 2013). In walled cells, such as fungal or plant cells, the process is further complicated because the cell wall first needs to be locally degraded while preserving cell integrity. The mechanisms underlying the cellular decision to fuse are unknown.

Conceptually, cell fusion may be initiated by diverse signals: Cell fusion may be induced upon physical engagement of transmembrane proteins, although, in walled cells, this would require proteins spanning the thickness of the cell wall. The signal may be of mechanical nature, leading to mechanical sensing of cells in contact. Support for this idea comes from the observation that imbalance in the osmotic pressure between two partner cells prevents cell fusion in Saccharomyces cerevisiae (Philips and Herskowitz 1997). Alternatively, it may be mediated by a chemical signal between partner cells at short range. The finding that S. cerevisiae cells expressing reduced levels of a-factor pheromone are specifically fusion-defective (Brizzard et al. 1996) suggests that pheromones may form such chemical signals. However, addition of exogenous pheromone to cells unable to secrete it does not restore fusion ability [Michaelis and Herskowitz 1988; Kjaerullff et al. 1994; Seike et al. 2013]. Individual cells exposed to even saturating pheromone levels also do not lyse (which would result from a fusion attempt without a partner cell), suggesting that the decision to fuse requires more than a simple step increase in pheromone signaling.

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We investigated fusion commitment in the sexual life cycle of the fission yeast *Schizosaccharomyces pombe*. Here, fusion occurs between two haploid cells of opposite mating types, known as M (h+\textsuperscript{−}) and P (h+\textsuperscript{+}), to form a diploid zygote. Mating is induced by nitrogen starvation and relies on pheromone-dependent communication (Merlini et al. 2013): The P cell secretes P factor, a 23-amino-acid peptide that activates its cognate receptor, Mam2, on the M cell (Kitamura and Shimoda 1991; Imai and Yamamoto 1994). Similarly, the M cell produces M factor, a short lipid-modified peptide that is transported outside M cells by a dedicated ABC-type transporter, Mam1, and sensed by the Map3 receptor on the P cell (Davey 1992; Tanaka et al. 1993; Christensen et al. 1997). Both receptors belong to the G-protein-coupled receptor (GPCR) family and signal through the same Ga subunit, Gpa1, and a conserved MAPK cascade to amplify the transcriptional mating program initially induced by nitrogen starvation (Obara et al. 1991; Hughes et al. 1993; Neiman et al. 1993; Xu et al. 1994). This leads to, among other things, the increased expression of pheromones, Mam1 transporter, pheromone receptors, and formin Fus1, which is essential for cell fusion. Beyond transcriptional control, pheromone signaling is also interpreted locally to induce polarized growth: Pheromone secretion machineries and receptor-coupled Ga coaccumulate at polarized cortical zones that dynamically explore the cell periphery and become stabilized by pheromone stimulation (Bendezu and Martin 2013; Merlini et al. 2016). Thus, facing zones in partner cells stabilize each other, leading to cell–cell pairing. In the ensuing polarized growth that brings cells into physical contact, pheromone receptors become enriched at the tip of the cellular extension (shmoo) toward the partner cell, as also observed in other fungi (Ayscough and Drubin 1998; Hirota et al. 2001; Merlini et al. 2016). Whether pheromone–MAPK signaling also directly regulates cell fusion is not known.

Cell fusion in fungi necessitates digestion of the cell wall to bring the plasma membranes into contact. In fission yeast cells, this is achieved through assembly of the actin fusion focus, a Fus1-dependent aster of linear actin cables that localizes the myosin V-dependent delivery of cell wall glucanases to the incipient fusion site (Dudin et al. 2015). Time-lapse imaging of the fusion focus revealed that this structure forms only in paired cells in contact, suggesting the existence of a signal at close range to trigger fusion focus assembly. Indeed, as fission yeast cells are under strong internal turgor pressure, estimated to be equivalent to that in a racing bike tire (~\textasciitilde 8 atm) [Minc et al. 2009], a fusion focus assembled too early or at the wrong place would lead to osmotic shock and cell lysis.

Here, we establish that fusion commitment is triggered by spatial focalization of the known pheromone–GPCR–MAPK signaling cascade. We show that pheromone signaling is both necessary and sufficient to induce cell fusion independently of physical cell contact and that the fusion signal is encoded not in the absolute amounts of pheromone but in its spatial organization. Concentration of pheromone release, GPCR, and MAPK signaling at the actin fusion focus results from a positive feedback loop between pheromone signaling and the actin fusion focus and induces cell fusion commitment by stabilizing the focus for local cell wall digestion. Thus, spatial reorganization of the signaling cascade underlies the decision to fuse.

### Results

#### A diffusible signal is necessary for cell fusion

Fission yeast cells efficiently form pairs and fuse when placed in a monolayer on solid medium [Dudin et al. 2015; Vjestica et al. 2016]. Similarly, homothallic yeast cells loaded in a flow chamber successfully formed zygotes in the absence of fluid flow but failed to do so in the presence of flow, which likely perturbed the formation of pheromone gradients [data not shown]. To test whether the flow directly perturbed fusion in addition to earlier events during mating, we let cell pairs engage in the fusion process in the absence of flow, forming fusion foci characterized by the focal localization of type V myosin Myo52 [Dudin et al. 2015], before transiently flowing fresh medium for 10 min [Fig. 1A]. Cell pairs engaged in fusion exhibited one of two distinct behaviors: Some were unperturbed by the flow and retained their fusion foci, defining a committed stage undisturbed by external flow [Fig. 1B, Supplemental Movie S1]. These cells rapidly fused together [Fig. 1C]. Others were uncommitted and disassembled their fusion foci, reassembling them and completing fusion with significant delay after the flow had been arrested [Fig. 1B,C, Supplemental Movie S1].

Four lines of evidence indicate that the committed pairs are more advanced in the fusion process than the uncommitted ones. First, committed cells fused faster than the average of untreated cells, indicating that commitment occurs in the later stage of the fusion process [Fig. 1C]. Second, the distance between the partner cell fusion foci was significantly smaller in the committed pairs, indicating closer cell–cell engagement [Fig. 1D]. Third, the fusion foci were more static in the committed pairs, as observed for late fusion pairs [Fig. 1E; Dudin et al. 2015]. Fourth, the second type V myosin Myo51, which accumulates at the fusion focus late in the process [Doyle et al. 2009; Dudin et al. 2015], was detected in both cells only in committed pairs [Supplemental Fig. S1]. We conclude that fusion consists of two successive stages: an early, uncommitted stage that requires an external diffusible signal and a late, committed stage. We hypothesized that pheromones represent this diffusible signal for fusion.

#### Autocrine cells attempt fusion in the absence of a partner cell

Any instructive fusion signal should carry sufficient information to induce the fusion process. We obtained direct evidence that pheromone signaling is sufficient to induce cell fusion by constructing autocrine cells. We replaced the coding region of the P-factor receptor Mam2 with that of the M-factor receptor Map3 at endogenous *mam2* genomic locus in M cells, yielding cells that
respond to the self-produced M factor [Fig. 2A]. During exponential growth, autocrine M cells [h\textsuperscript{−}mam2Δ::map3] were indistinguishable from wild-type M cells, as judged by cell length, cell width, septum position, growth assays, and localization of polarity factors such as type V myosin Myo52 and Cdc42 scaffold Scd2 (Supplemental Fig. S2A–E). This is consistent with the previous observations that expression of pheromones and their receptors is induced by starvation (Kitamura and Shimoda 1991; Tanaka et al. 1993; Kjaerulff et al. 1994). In contrast, upon nitrogen starvation, these autocrine cells spontaneously activated pheromone signaling, as reported previously (Tanaka et al. 1993): They stopped dividing and expressed pheromone-responsive gene products such as the pheromone transporter Mam1, the pheromone-dependent formin Fus1, and the replaced pheromone receptor Map3 (Supplemental Fig. S2F; Petersen et al. 1995; Christensen et al. 1997). Autocrine cells also exhibited strong morphological responses characteristic of sexual differentiation, forming multiple successive projections, leading to aberrantly shaped cells (Fig. 2B, Supplemental Movie S2) as well as at sites of lysis [Fig. 2D, Supplemental Movie S2]. We note that Myo52 spots associated with lysis were stable for a significantly longer time preceding the lysis event than at other times [Fig. 2E]. The Myo52 spot colocalized with the formin Fus1 as well as with Exg3 and Eng2, two glucanases that localize at the fusion focus and are needed to degrade the cell wall during fusion (Fig. 2F; Dudin et al. 2015), suggesting that these led to local cell wall digestion. In agreement with this, osmotic stabilization by addition of 1.2 M sorbitol in the medium efficiently suppressed cell lysis [Fig. 2C]. Finally, deletion of fus1 prevented formation of the Myo52 focus and completely suppressed cell lysis [Fig. 2C,G; data not shown]. We conclude that autocrine M cells assemble a fusion focus-like structure and attempt fusion in the absence of a partner cell, leading to cell lysis.

This attempted fusion upon autocrine signal activation represents a complete fusion response. Indeed, two autocrine M cells were occasionally able to fuse with each other. This mostly happened shortly after cell division, with the two sister cells re-fusing together (Fig. 2H; Dudin et al. 2015). While these events were infrequent, their existence demonstrates that autocrine M cells mount a genuine fusion response able to go to completion.

In summary, these data establish that the signal to trigger cell fusion does not rely strictly on cell–cell contact and can be elicited by simple autocrine activation of pheromone signaling. We infer that paracrine pheromone signaling in the normal situation of cell pair engagement also triggers fusion.
Focalized pheromone release serves as fusion signal

Addition of synthetic pheromone to heterothallic cells has been shown to promote cell cycle arrest, initiation of the sexual transcriptional program, and cell polarization (Davey and Nielsen 1994; Imai and Yamamoto 1994; Petersen et al. 1995; Christensen et al. 1997; Bendezu and Martin 2013). However, in contrast to the autocrine situation presented above, in either P or M cells exposed to very high concentrations of synthetic M or P factor, respectively, we did not observe fusion focus assembly or extensive cell lysis even upon deletion of the proteases that normally degrade these pheromones (Fig. 3A; Supplemental Fig. S3). Using time-lapse microscopy, we found that these cells transiently concentrated the Myo52 signal, but such foci were not maintained over time (L Merlini, unpubl.). One reason for this difference may be that the spatial organization of the pheromone signal is distinct in the two cases.

While pheromones are homogeneously distributed upon exogenous addition, local release in the autocrine situation may allow for a sharp, graded pheromone distribution. Although direct visualization of M factor is not possible, we found that, in autocrine M cells, the M-factor transporter Mam1 was focalized to a single sharp dot, which colocalized with the...
fusion focus (60 of 80 cells) and depended on fus1, suggesting that M factor is locally released at this location (Fig. 3B). Thus, autocrine activation of pheromone signaling promotes the focal organization of pheromone release.

We tested the importance of focal pheromone release to promote fusion attempts in autocrine M cells by homogenization of pheromone distribution. Autocrine M cells in suspension underwent lysis as described on solid medium above, but the number of lysed cells was reduced by strong agitation (Fig. 3C). Similarly, autocrine M cells placed in flow chambers lysed in the absence of flow but did not upon continuous flow of fresh medium [Fig. 3C]. Remarkably, addition of a high concentration of M factor on solid medium also strongly reduced the occurrence of lysis [Fig. 3D], indicating that the local...
pheromone gradient rather than the absolute pheromone concentration acts as fusion signal.

**Pheromone signaling is focalized at the fusion focus**

In addition to the Mam1 transporter, the M-factor receptor Map3 was highly enriched at the fusion focus of autocrine M cells labeled with Myo52-tdTomato (41 of 83 cells). Similarly, the receptor-coupled Gα Gpa1 localized at the fusion focus (19 of 54 cells). In fus1Δ autocrine cells, Map3 and Gpa1 were broadly distributed at the plasma membrane [Fig. 3B]. In contrast, neither pheromone receptor (the P-factor receptor Mam2 in M cells or the M-factor receptor Map3 in P cells) was focalized in heterothallic cells exposed to a high concentration of synthetic pheromones [Supplemental Fig. S3].

This focalized organization of pheromone signaling was also observed in wild-type mating pairs: Mam1, Map3, and Gpa1 all accumulated at the fusion focus in a fus1Δ-dependent manner [Fig. 3E]. Time-lapse examination of Map3 showed that signaling focalization occurs late in the fusion process: Whereas Myo52 focalized 75 min ± 15 min before cell–cell fusion, Map3-GFP colocalized as a dot only 43 min ± 13 min before fusion [Fig. 3F,G]. Moreover, we found that committed cells as defined in flow cell experiments [see Fig. 1] showed focalization of the pheromone receptor Map3, whereas no focalization of Map3 was observed in uncommitted cells [Fig. 3H]. We conclude that pheromone signaling becomes highly spatially constrained at the fusion focus and that this correlates with commitment to cell fusion.

**Loss of pheromone receptor focalization correlates with fusion defect**

To test the role of pheromone signaling focalization in cell fusion, we strove to perturb signaling focalization without disturbing the formation of the fusion focus or earlier events of cell differentiation and polarization. We reasoned that the focal distribution of the transmembrane receptor would be disturbed by interfering with its endocytic retrieval, likely necessary to counter the effects of lateral diffusion at the plasma membrane. Indeed, a Map3 allele with a truncation of its C-terminal cytoplasmic tail [Map3dn9-GFP, previously shown to be endocytosis-deficient] [Hirota et al. 2001] decorated the plasma membrane broadly and was not enriched at the Myo52-labeled fusion focus, whether it was expressed as the sole Map3 copy in autocrine M cells or mating P cells [Fig. 4A,B]. Remarkably, the map3dn9 truncation completely blocked cell lysis in autocrine M cells [Fig. 4C]. However, these cells, like their map3Δ counterparts, activated pheromone signaling: They stopped dividing, expressed pheromone-responsive transcripts [data not shown], and exhibited growth projections characteristic of sexual differentiation, which, however, occurred only at cell poles [Fig. 4B]. Consistently, map3dn9 mutant P cells were able to engage in mating pairs [Bendezu and Martin 2013] but were almost completely fusion-defective [Fig. 4D]. Thus, the endocytosis-deficient Map3dn9 allele largely separates early cell differentiation and polarization events from late ones depending on focalization. We note that Map3 C-terminal truncation may also affect other aspects of its regulation in addition to endocytosis. In both autocrine cells and cell pairs, the focalization of Myo52 indicates that map3dn9 mutant cells were able to form a fusion focus [Fig. 4A,B]. However, this focus was significantly more mobile than in wild-type cells [Fig. 4A,B,E; Supplemental Movie S4]. As a result, glucanases failed to become enriched at a focal point [Fig. 4H]. We conclude that pheromone receptor endocytosis is necessary for receptor concentration at the fusion focus and is required to stabilize the fusion focus.

We found a second condition in which the pheromone receptor is not enriched at the fusion focus by examining the phenotype of rgs1Δ cells. Rgs1 is a GTPase-activating protein for the receptor-coupled Gα Gpa1 previously characterized for its role in desensitization [Watson et al. 1999; Pereira and Jones 2001; Croft et al. 2013]. In this mutant, which mounts an exaggerated pheromone response, the Map3 receptor was not enriched at the fusion focus in both autocrine M cells and P cells engaged in cell pairs [Fig. 4F,G]. Mam1 and Gpa1 were also not enriched at the fusion site in these cells [Fig. 4G]. Similar to the case of map3dn9, rgs1Δ autocrine M cells did not lyse, and engaged rgs1Δ pairs failed to efficiently fuse [Fig. 4C,D; Bendezu and Martin 2013]. While these cells were able to assemble a fusion focus, this focus was transient, often breaking up in multiple dots and showing multiple cycles of formation and disappearance [Fig. 4A,B,E; Supplemental Movie S5]. Glucanases were also unfocalized in this mutant [Fig. 4H]. While the molecular mechanism by which Rgs1 influences Map3 focalization remains to be established, these observations further support the model that pheromone signaling local concentration stabilizes the fusion focus, which in turn allows the precise delivery of glucanases to pierce the cell wall for fusion. Because the focus was significantly less stable in rgs1Δ than in map3dn9, Rgs1 may also regulate focus stability in additional ways independent of Map3 focalization.

**Pheromone signaling stabilizes the fusion focus**

If spatial focalization of pheromone signaling promotes fusion by stabilizing the fusion focus, we reasoned that it might be possible to rescue the map3dn9 fusion defect by forcing the artificial recruitment of the endocytosis-defective receptor to the fusion focus. We used the tight binding between GFP and the GFP-binding protein [GBP] [Rothbauer et al. 2006, 2008] to recruit Map3dn9-GFP to Myo52-GFP-mCherry at the fusion focus [Fig. 5A]. Remarkably, this combination led to recruitment of Map3dn9-GFP to the fusion focus, stabilization of the focus, and a sevenfold increase in successful fusion [Fig. 5B,C,F; Supplemental Movie S6]. A sevenfold increase in cell lysis was also observed [Fig. 5C]. As expected, this rescue of cell fusion was abrogated upon fus1 deletion. In the presence of Myo52-GFP-mCherry, concentration of the Map3dn9-GFP...
signal was observed in 76% of the mating pairs (Fig. 5D), but only 43% formed a tight focus (Fig. 5E). In other cells, the Map3dn9-GFP signal appeared fragmented in multiple dots, enriched over a wider zone (>0.5 µm), or not concentrated at all (Fig. 5B,E,F,H). Remarkably, we observed a very tight correlation between the appearance of Map3dn9-GFP as a tight focus, the observation of a stable Myo52 focus (Fig. 5F; Supplemental Movie S7), and the ability of cells to fuse: Only cells with a tight focalization of Map3dn9-GFP (but nearly all of these) fused efficiently with a partner (Fig. 5G). This observation supports the idea that the principal reason for the observed fusion defect of map3dn9 cells is the failure of the truncated receptor to accumulate at the fusion focus. Cells with a fragmented Map3dn9-GFP signal frequently lysed, indicating deregulation of cell wall degradation (Fig. 5G). We conclude that tight localization of the pheromone receptor at the fusion focus is a key, necessary element to induce cell fusion.

Figure 4. Loss of pheromone receptor focalization leads to severe fusion defects. (A) Time lapse of Myo52-tdTomato in map3dn9-GFP (top) and rgs1Δ (bottom) mating pairs. Note the displacement of the Myo52 signal relative to the static reference line. (Top) Note the stable Myo52 signal in the h+ partner, which is phenotypically wild type, as it does not express map3dn9. (B) Time lapse of Myo52-tdTomato in map3dn9-GFP (top) and rgs1Δ (bottom) autocrine M cells. (C) Cell lysis of wild-type, fus1Δ, map3dn9Δ, and rgs1Δ autocrine M cells. N > 1000. [***] P < 3 × 10⁻⁵, t-test. (D) Fusion efficiency of wild-type, fus1Δ, map3dn9Δ, and rgs1Δ homothallic pairs. n > 300. [***] P < 6 × 10⁻⁴, t-test. (E) Mean Myo52-tdTomato displacement per 2-min time point over 30 min in homothallic wild-type and map3dn9Δ and rgs1Δ mutants. n = 10. [**] P < 0.005, t-test map3dn9Δ; [***] P < 2 × 10⁻⁵, t-test rgs1Δ. (F) In the absence of Rgs1, the Map3 pheromone receptor is unable to focalize in autocrine M cells. (G) Broad weak Mam1-sfGFP, Map3-GFP, and Gpa1-mCherry localization in rgs1Δ cell pairs. (H) Cell wall glucanases Exg3-sfGFP and Eng2-sfGFP focalize in wild-type but not in map3dn9Δ or rgs1Δ pairs (arrowheads). Bars, 2 µm.

Focalization of Byr1 MAP2K leads to fusion commitment

Pheromone signal transduction involves a dedicated MAPK cascade composed of the MAP3K Byr2, MAP2K Byr1, and MAPK Spk1. Consistent with focalization of pheromone receptors and Gpa1, Byr2, Byr1, and Spk1 were all enriched at the fusion focus of wild-type mating pairs in a Fus1-dependent manner (Fig. 6A,B). As Byr1-sfGFP exhibited the strongest signal of the three, it was used to probe MAPK cascade localization in further experiments, which revealed a perfect correlation between pheromone receptor and Byr1 focalization: Byr1 was enriched at the fusion focus in autocrine M cells, which form a fusion focus containing Map3 (Fig. 6B); it localized to the shmoo tip of M and P cells exposed to homogenous synthetic pheromones but was not focalized, consistent with the absence of a fusion focus in these cells (Supplemental Fig. S3); similarly, it was present but unfocalized at the shmoo tip of map3dn9 and

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rgsΔ mutant cells [Fig. 6C]. Moreover, similar to Map3, Byr1 focalization occurred late in the fusion process, 42.5 min ± 15.5 min before fusion (Fig. 6D, compare with Fig. 3G). Interestingly, a constitutively active allele of Byr1, Byr1DD, accumulated at the shmoo tip but failed to focalize, and byr1DD mutants did not form a fusion focus and were strongly fusion-defective (Supplemental Fig. S4). Thus, focalization of Byr1 MAP2K and thus likely the whole MAPK cascade strongly correlates with fusion attempts.

To test whether MAPK focalization is the key event to induce fusion, we forced Byr1-GFP recruitment at the fusion focus in fusion-deficient strains carrying a GBP-tagged Myo52. We first generated an untagged map3dn9 allele, which was stably integrated as an extra copy in the genome. Cells carrying this allele showed phenotypes similar to, but somewhat less severe than, map3dn9-GFP mutant cells, exhibiting a mobile fusion focus and failing to efficiently fuse [Fig. 6E and data not shown]. As map3dn9 was present in addition to endogenous map3, this allele is dominant-negative. Remarkably, forced recruitment of Byr1-GFP to the unstable fusion focus of map3dn9 and rgsΔ mutant cells led to stabilization of the fusion focus, which showed increased lifetimes in kymographs and reduced mobility by quantifying instantaneous displacement (Fig. 6E–G; Supplemental Movies S8, S9). This led to a 2.5-fold to threefold increase in fusion/lysis compared with cells lacking Myo52-GBP-mCherry (Fig. 6E,F,H; Supplemental Movies S8, S9). Lysis was suppressed by addition of 1.2 M sorbitol or by deleting fus1, indicating that these cells were indeed attempting untimely fusion [Fig. 6H]. Finally, even in heterothallic M cells exposed to synthetic P factor in which fusion foci are only transiently observed, the forced recruitment of Byr1-GFP to Myo52GBP-mCherry induced the formation of a stable focus and a significant level of lysis (Fig. 6I,J). In contrast, forced recruitment of Byr1DD-GFP, which never forms a fusion focus, to Myo52-GBP did not lead to formation of a fusion focus [data not shown]. We conclude that recruitment of the MAP2K Byr1—and thus likely the whole pheromone–MAPK cascade at the fusion focus—is critical to consolidate the focus and promote cell fusion.
Discussion

Here, we addressed how cells make the decision to fuse. While this decision is expected to be coupled with the formation of a cell pair, we established that it requires neither cell–cell contact nor a signal distinct from that inducing previous events. Instead, it relies on diffusible pheromones, which also signal the earlier stages of cell differentiation and cell polarization. Importantly, we showed that the fusion information is coded in their spatially focalized distribution. This focalized organization results from a positive feedback loop between pheromone signaling and the actin fusion focus, which underlies the local concentration of pheromone release and signal activation, leading to the concentration at the fusion site of the MAPK cascade. In turn, this local concentration of the MAPK module drives fusion commitment by stabilizing the actin focus.

How does focalization of pheromone signaling happen?

The autocrine M cells show that the coexpression of a pheromone receptor and its ligand in the same cell is sufficient to drive the formation of a single focus, indicative
of a positive feedback motif [Fig. 7]. We propose that this positive feedback relies on two key elements: (1) the signaling machinery, which includes the M-factor receptor Map3, its ligand transported by the Mam1 transporter, and the MAPK cascade, and (2) the actin fusion focus, consisting principally of the formin Fus1 and type V myosins.

Two main reactions underlie this positive feedback. One reaction involves Fus1 assembling the actin cables along which Map3 and Mam1, both transmembrane proteins, are delivered to the plasma membrane in exocytic vesicles. At the plasma membrane, endocytosis retrieves Map3 [Hirot a et al. 2001], thus opposing lateral diffusion and ensuring Map3 enrichment at the site of delivery. The Mam1 transporter is likely also retrieved through endocytosis, as has been demonstrated for its budding yeast homolog, Ste6 [Kolling and Hollenberg 1994]. This leads to the activation of Map3 with maximal ligand concentration at or close to sites of exocytic delivery and consequent recruitment of the MAPK module at these sites. It is noteworthy that spatial restriction of both ligand and receptor is required, indicating that local graded pheromone distribution rather than its absolute concentration significantly contributes to focalization of the downstream signal. This also suggests that cells may have the ability to measure the steepness of the pheromone profile. The second reaction consists of the focalization of the MAPK module promoting the stabilization of the actin fusion focus. The molecular details of this reaction remain to be defined. It may be conceptually similar to how G-protein and MAPK polarization constrain the polarity patch for gradient tracking by forming a positive feedback on Cdc42 GTPase activation [Hegemann et al. 2015; McClure et al. 2015]. Alternatively, the MAPK signal may act more directly on Fus1 activation or recruitment or involve other elements of the fusion focus. In the autocrine situation, the positive feedback motif created by these two reactions underlies a symmetry-breaking event leading to the formation of a single constrained actin focus, which sharpens the localization of the signaling machinery, which in turn further constrains the focus.

In the natural situation involving the pairing of cells of distinct mating types, the only likely difference is that the pheromone ligand for receptor activation is delivered by the partner cell [Fig. 7]. In this case, the site of pheromone release from one cell defines the site of receptor activation and thus actin fusion focus stabilization in the second cell, which in turn promotes the local release of pheromone to sharpen the site of signaling in the first one. Thus, the feedback system is established upon bilateral cell pairing, ensuring that the cells do not focalize their signaling machinery in the absence of a mating partner.

In summary, at its core, pheromone signaling focalization relies on a positive feedback loop, similar to other symmetry-breaking systems [Motegi and Seydoux 2013; Yi et al. 2013; Martin 2015] in which Fus1 promotes the delivery of the receptor and its activator, and the activated receptor promotes Fus1 function.

Spatial focalization of MAPK signaling as a cellular decision

One notable and perhaps unexpected result is that forced recruitment of either pheromone receptor or the downstream MAP2K to the fusion focus induces fusion attempts. Because both pheromone receptor and Byr1 MAP2K recruitments have the same phenotypic consequence of stabilizing the fusion focus, it is unlikely that this is caused simply by tethering the fusion focus to the endocytosis-deficient receptor at the plasma membrane or forcing a kinase–substrate contact. Rather, in conjunction with the observation that pheromone signaling focalization coincides with the commitment of wild-type mating pairs to fusion, this result suggests that the cellular decision to fuse is triggered by the spatial concentration of MAPK signaling.

Mechanisms that underlie cellular decisions must typically convert a graded signal [here pheromones] into an all-or-none response [here the decision to fuse]. Classical work on the MAPK cascade has shown that it can function as a switch that converts a low-level signal into a maximal response.
signal [Ferrell 1996]. However, the MAPK module is highly versatile and can be modulated in various cell types and situations to generate distinct graded or switch-like outputs [Inder et al. 2008; English et al. 2015]. For instance, the MAPK pheromone pathway in S. cerevisiae exhibits an essentially graded response to induce gene expression proportional to pheromone concentration [Poritz et al. 2001; Paliwal et al. 2007], but its role in inducing the formation of the mating projection is switch-like, a behavior dependent on the MAPK scaffold Ste5 [Hao et al. 2008; Malleshaiah et al. 2010]. The S. pombe pheromone MAPK module, like the mammalian Raf–MEK–ERK cascade, which can functionally replace it [Hughes et al. 1993], lacks a scaffold protein. Interestingly, synthetic biological approaches have shown that a simple concentration increase of sequential Raf–MEK–ERK kinases is sufficient to lower the activation threshold and enhance ultrasensitivity, leading to full activation [O’Shaughnessy et al. 2011]. This evidence leads us to propose that the local concentration of the pheromone MAPK cascade at the fusion site in S. pombe represents a natural situation in which the cell exploits the ultrasensitivity of the MAPK cascade to convert it to the fully active state, which, due to the spatially restricted location in which this takes place, drives an essentially irreversible decision to fuse.

Similar spatial rearrangements may control cellular decisions in other cell types. The spatial organization of signaling molecules is generally thought to be important for signal transduction [Kiel and Serrano 2012]. For instance, during adaptive immunity, activation of the T-cell receptor is thought to result from its spatial segregation from bulky inactivating phosphatases that are sterically excluded from the zone of receptor engagement with the antigen-presenting cell [Davis and van der Merwe 2006; James and Vale 2012]. Similarly, distinct subcellular localization of GPCR-mediated signaling and the MAPK cascade can drive distinct cellular responses in various cell types [Harding et al. 2005; Inder et al. 2008; West and Hanyaloglu 2015], although it is not known to what extent spatial changes are harnessed to modulate the signal. It will be interesting to probe whether other cells spatially reorganize their signaling pathways to change the system’s output.

Mechanistic function of fusion focus stabilization

How does spatial stabilization of the fusion focus promote cell fusion? One critical role of the fusion focus is to promote the local delivery of glucanases that catalyze the hydrolysis of the bonds in glucan polymers to locally dissolve the cell wall, whereas glucan synthases are more broadly distributed [Dudin et al. 2015]. In the absence of the fusion focus, in fus1Δ pairs, both enzymatic activities occupy similarly broad distributions, leading to cell wall remodeling for polarized growth but not cell wall dissolution [Dudin et al. 2015]. Glucanases are likely also broadly distributed in byr1ΔΔ mutants, in which localization fails. Similarly, in map3ΔΔ or rgs1Δ mutant pairs that display an unstable fusion focus, no sharp glucanase localization can be detected. We hypothesize that glucanases are still locally released at the fusion focus, but their [broad or nearly undetectable] localization reflects the time-averaged consequence of secretion from a motile site, which is similar to the localization resulting from secretion over a broad region. Thus, in the absence of fusion focus stabilization, the cell wall cannot be pierced, and cells instead continue to grow. In contrast, immobilization of the fusion focus leads to local imbalance in favor of hydrolytic activity such that the cell wall is pierced for fusion.

Timing of the decision to fuse

Because yeast cell growth is driven by turgor pressure, the cell wall is essential to preserve cell integrity. Therefore, the decision to stabilize the fusion focus to locally digest the cell wall has to be precisely coordinated with the formation of cell pairs. Indeed, untimely fusion focus stabilization in autocrine cells or upon forced signaling focalization leads to cell lysis because cell wall digestion occurs in the absence of an attached partner cell. Interestingly, ~60% of autocrine M cells do not lyse, at least within the time frame of our experiments, whereas wild-type mating pairs almost always fuse. Thus, while focalization of pheromone signaling serves as a fusion signal, the zone of cell–cell contact may offer a favorable context for this signal, for instance, by constraining diffusion in the cell wall to two dimensions, thus promoting a local increase in pheromone levels, as proposed previously for cell wall remodeling enzymes [Huberman and Murray 2014]. Alternatively, cells may possess built-in break mechanisms that antagonize the positive feedback for focalization, which are alleviated upon cell–cell contact. Future work should establish how the decision to fuse is taken at the appropriate time.

Materials and methods

Strains, media, and growth conditions

Strains used in this study are listed in Supplemental Table S1. For assessing cells during exponential growth, cells were grown in rich medium [YE], Edinburgh minimal medium [EMM], or minimal sporulation medium with nitrogen [MSL + N] supplemented with amino acids as required. For assessing cells during the mating process, liquid or agar minimal sporulation medium without nitrogen [MSL – N] was used [Egel et al. 1994; Vjestica et al. 2016]. All live-cell imaging was performed on MSL – N agarose pads [Vjestica et al. 2016] or in microfluidics chambers [described below]. Mating assays were performed as in Vjestica et al. [2016].

Gene tagging was performed at endogenous genomic loci at the 3’ end, yielding C-terminally tagged proteins as described [Bähler et al. 1998] and was confirmed by diagnostic PCR for both sides of the gene insertion. Tagging with sfGFP was performed as in Dudin et al. [2015]. The map3-GFP tag was obtained from Hirota et al. [2015]. We also generated a map3–sfGFP, in which sfGFP was inserted at the 3’ end of the map3 ORF and likely disrupted the nested gene in the opposite direction without obvious ill effects. This strain was used only in Supplemental Figure S3, for which the result was also confirmed with a map3-GFP strain. Tagging with GBP-mCherry was performed at the C termini of endogenous genomic loci.

Construction of plasmid for GBP-mCherry tagging was done as follows. First, a GBP-mCherry fragment was amplified from
To generate maps\textsuperscript{dso} sfGFP autocrine M cells, we truncated the last 228 nucleotides of the map3\textsuperscript{dso} coding region in the M-cell autocrine cells obtained above by integration of sfGFP, yield-}


ting cells × 100. For mating assays in microfluidic chambers, mat-

ing cells were prepared as described above but were not mounted

on agarose pads. Instead, cells were loaded into a microfluidic cell

chamber (CellASIC Corp., Y04C) with the ONIX (CellA-

sic Corp.) as an automated controller, allowing rapid medium

change. After the addition of the MSL \textsuperscript{−} agarose

pads, and cells were incubated

The DeltaVision platform (Applied Precision) described previous-

ly (Bendezu and Martin 2013) was used mainly for time-lapse im-

aging of a 4.6-μm z section, which is

**Microscopy and image analysis**

The DeltaVision platform (Applied Precision) described previously

by Bendezu and Martin (2013) was used mainly for time-lapse imaging

overnight or during microfluidic experiments (Figs. 1, 2D,E, 3A,B,D–F, 4F–H, 6B–J). **Supplemental Figs. S1, S2A–C.** To limit photobleaching, overnight movies were captured by OAI (optical axis integration) imaging of a 4.6-μm z section, which is

**Mating assays**

Mating assays, including treatment of autocrine M cells, were per-

formed as in Bendezu and Martin (2013), Dudin et al. (2015),

and Vjestica et al. (2016). Briefly, precultures of cells were grown

at 25°C to OD\textsubscript{600} = 0.4–1 in MSL + N. Cultures were then diluted

to OD\textsubscript{600} = 0.025 in MSL + N (for heterothallic crosses, cells were

mixed in equal parts) and grown for 18–20 h to OD\textsubscript{600} = 0.4–1 at

25°C in MSL + N. Cells were harvested by centrifugation, washed

three times with MSL + N, and, unless indicated otherwise,

mounted onto MSL – N 2% agarose pads that were incubated

for either 1 h at 25°C before imaging in overnight movies or over-

night at 18°C before imaging. Fusion efficiency was measured as

in Dudin et al. (2015). The percentage of lysis was calculated as

percentage lysis = number of lysing cells/total amount of shino-

cing cells × 100. For mating assays in microfluidic chambers, mat-

ting cells were prepared as described above but were not mounted

on agarose pads. Instead, cells were loaded into a microfluidic cell

culture chamber (CellASIC Corp., Y04C) with the ONIX (CellA-

sic Corp.) as an automated controller, allowing rapid medium

flow. Cells in MSL – N were loaded into the microfluidic cham-

ber and left for 2 h at 25°C in the absence of fluid flow. To tran-

siently perturb any diffusible gradient, fresh MSL – N medium

was flowed at a rate of 13 μL/h (5 psi) for 10 min. To obtain control

cells without flow, scotch tape was tightly fixed on the first six air

channels of the manifold of the corresponding chamber. For pher-

omone treatments, P-factor phenome was purchased from Pep-

nome and used from a stock solution of 1 mg/mL in methanol. M

factor was synthesized and purchased from Schaefer-N and used

from a stock solution of 2 mg/mL in methanol. Different concen-

trations of pheromones were directly added to the agarose pads.

Cells were then incubated overnight at 25°C prior to imaging.

Ethanol was used as a control.

Experiments on autocrine M cells were performed similarly to

normal homothallic strains. However, as the autocrine cells are

extremely sensitive to the absence of nitrogen, cells were usually

maintained in MSL + N at an OD = 600 < 0.6. Autocrine cells were

washed with MSL – N and placed directly on MSL – N agarose

pads. For experiments including external pheromones, M factor

was added directly in the agarose pads, and cells were incubated

for 16 h.
essentially a real time z sweep (Figs. 2D, 3A,F, 4F, 6B,F). Spinning-disk microscopy, also described previously [Bendezu and Martin 2013], was used mainly for quantitative analyses of fusion efficiency, cell lysis, cell size, and calcofluor staining as well as high temporal resolution and Z-stack maximal projection images [Figs. 2B,C,E, 3C–E, 4A–E, 5A–H, 6A,B]. Supplemental Figs. S2A–C, E, S3, S4. To measure the mobility of the fusion focus [Figs. 1E, 5D], the X and Y coordinates of the highest-intensity pixel of the Myo52-tdTomato signal were used to measure the instantaneous displacement between each time point (every 2 min) in each mating partner. When Myo52 signal was broad, we recorded the coordinates of the maximum fluorescence intensity. Displacement was calculated as \( \sqrt{[X_n - X_{n-1}]^2 + [Y_n - Y_{n-1}]^2} \). This was then averaged over the length of the time-lapse acquisition. Kymographs in Figures 5 and 6 were constructed in ImageJ version 1.47 [National Institutes of Health] by drawing a 3-pixel-wide line at the cell tip.

**Gene expression quantifications**

Fluorescence intensities of Mam1-sfGFP, Fus1-sfGFP, and Map3-sfGFP in Figure 2B were measured in ImageJ using a manually drawn line around the entire cells in sum projections of seven slices over 4-µm total depth. Background fluorescence for sfGFP was measured from similarly treated cells lacking sfGFP and was subtracted from the original measurements.

Figures were assembled with Adobe Photoshop CS5 and Adobe Illustrator CS5. All error bars are standard deviations. All experiments were done a minimum of three independent times, and statistical analysis was done across repeats of the same experiment.

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