Ultrastructural plasma membrane asymmetries in tension and curvature promote yeast cell fusion

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Cell–cell fusion is central for sexual reproduction, and generally involves gametes of different shapes and sizes. In walled fission yeast *Schizosaccharomyces pombe*, the fusion of *h*− and *h*+ isogametes requires the fusion focus, an actin structure that concentrates glucanase-containing vesicles for cell wall digestion. Here, we present a quantitative correlative light and electron microscopy (CLEM) tomographic dataset of the fusion site, which reveals the fusion focus ultrastructure. Unexpectedly, gametes show marked asymmetries: a taut, convex plasma membrane of *h*− cells progressively protrudes into a more slack, wavy plasma membrane of *h*+ cells. Asymmetries are relaxed upon fusion, with observations of ramified fusion pores. *h*+ cells have a higher exo-/endocytosis ratio than *h*− cells, and local reduction in exocytosis strongly diminishes membrane waviness. Reciprocally, turgor pressure reduction specifically in *h*− cells impedes their protrusions into *h*+ cells and delays cell fusion. We hypothesize that asymmetric membrane conformations, due to differential turgor pressure and exocytosis/endocytosis ratios between mating types, favor cell–cell fusion.

Introduction

Cell–cell fusion is a fundamental process underlying sexual reproduction and development. Plasma membrane (PM) merging for cell fusion requires a number of preparatory steps. These include the formation of cell adhesion, the remodeling of extracellular material to allow membrane contact, and significant force to overcome repulsive membrane and hydration charges. The actin cytoskeleton can generate this force and allow the engagement of membrane fusogenic machineries (Chan et al., 2020; Shilagardi et al., 2013). Fusogenic proteins, only identified in some organisms and cell types, then likely drive the formation of fusion pore(s) (Brukman et al., 2019; Hernández and Podbilewicz, 2017). However, fusion pore formation and expansion during cell–cell fusion have not been directly observed in any system.

Though fusion processes and machineries are extraordinarily diverse, the two cells engaging in fusion are asymmetric in the vast majority of cases. Fertilization largely happens between gametes of widely differing shape and size. Even *Chlamydomonas reinhardtii* gametes, with very similar size and shape, show distinct mating structures, only one of them extending an actin-rich protrusion (Goodenough et al., 1982). The fusion of somatic cells also exhibits asymmetries, for instance during muscle development in *Drosophila*, where actin assembly in the myoblast powers podosome protrusion into the myotube, which responds through a myosin II–dependent cortical tension increase to promote fusion (Kim and Chen, 2019; Kim et al., 2015; Sens et al., 2010). Similar asymmetries have been described in other types of somatic cell fusion (Brukman et al., 2019). By contrast, cell fusion during sexual reproduction in yeast is thought to happen between two isogametes, i.e., gametes of similar morphology.

The differences between yeast gametes are defined by the genetic information in the active mating type locus, which encodes distinct pheromone and receptor genes. Pheromone-dependent communication underlies sexual differentiation and the formation of cell pairs (Merlini et al., 2013). Fission yeast cells are naturally homothallic, i.e., they can switch between mating types, making the population mating-competent when faced with nitrogen starvation. Cells can also be fixed in one or other heterothallic mating type, *h*− and *h*+, which can mate when mixed together. The two pheromones have distinct biochemical qualities and modes of secretion, and also exhibit slightly different functions and evolutionary constraints, suggesting some functional diversification between mating types (Merlini et al., 2013; Seike et al., 2019a; Seike et al., 2019b). However, very few additional genes are differentially expressed (Mata and Bähler, 2006; Xue-Franzén et al., 2006), and the cell fusion process itself is thought to happen between two symmetric cells.

The cell wall (CW) encasing yeast cells needs to be remodeled for cell morphogenesis. For cell growth, remodeling happens at

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https://doi.org/10.1083/jcb.202103142
the cell projection (also known as shmoo) tip, with turgor pressure providing the critical force (Attilgan et al., 2015; Minc et al., 2009). For cell fusion, the CW is locally digested to allow membrane contact. This requires an actin-based structure called the fusion focus, assembled by the formin Fus1, which serves to concentrate the myosin V–dependent delivery of secretory vesicles containing hydrolytic enzymes (Dudin et al., 2015). Turgor pressure is hypothesized to then help bring membranes in contact (Martin, 2016). Interestingly, we had previously described an asymmetry between mating types in the formation of the fusion focus, which is stabilized earlier in $h^-$ than $h^+$ cells (Dudin et al., 2015). While advanced live-cell imaging studies have revealed significant molecular and mechanistic information on the assembly of the fusion focus (Billault-Chaumartin and Martin, 2019; Dudin et al., 2015; Dudin et al., 2017; Dudin et al., 2016; Merlini et al., 2018), how the two cells shape their PMs for merging and how the initial fusion pore expands are not understood.

Here, we used correlative light microscopy (LM) and electron tomography to study cell fusion in Schizosaccharomyces pombe at the ultrastructural level. Our data show the organization of the fusion focus as a vesicle-rich structure that excludes other organelles. We describe all fusion stages, including pairs with membrane contact, and small and large fusion pores, indicating inhomogeneous pore expansion. Unexpectedly, the two pre-fusion gametes exhibit asymmetries in membrane tension and curvature, with $h^-$ cells protruding into $h^+$ cells. We trace the origin of these asymmetries to different ratios of endo- and exocytic activities and different turgor pressure between mating types, respectively, and propose that these asymmetries promote efficient cell–cell fusion.

Results

Ultrastructure of the yeast fusion site

We used correlative light and electron microscopy (CLEM) to study the ultrastructure of the actin fusion focus in homothallic (h90) WT strains expressing Fus1-sfGFP and Myo52-tdTomato. Using LM, we selected cell pairs exhibiting a Fus1 and Myo52 fusion focus signal, which allowed restriction of our analysis between ~1 h before and 15 min after cell fusion, corresponding to the fusion focus lifetime (Dudin et al., 2015). We subsequently identified these pairs in transmission EM (TEM) and acquired tilt series to generate a dataset of 124 3D tomographic reconstructions (Fig. S1 A). Based on the minimal distance between the two PMs, we assigned each pair to one of four stages along the fusion process (Fig. 1, A–D). We distinguish three pre-fusion stages: two early stages where cells are in contact through their CW, with average minimal PM distance of 142 ± 50 nm for the first stage (far CW contact; 13 tomograms; Fig. 1 A), and 49 ± 26 nm for the second (close CW contact; 50 tomograms; Fig. 1 B and Video 1), where CWs have merged; and a third stage (PM contact; 11 tomograms; Fig. 1 C), with an average minimal PM distance of 5 ± 4 nm, where some regions of CWs are fully digested and the PMs appear in contact. The fourth stage is after fusion, where both cytosols are continuous. Among post-fusion pairs, the vast majority showed a single large fusion pore (43 tomograms; Fig. 1 D and Video 2). A further seven tomograms, difficult to classify, showed local loss of density in the CW suggestive of continuity between partner cells, and are described in more detail below. The distribution of tomograms according to minimal PM distances is shown in Fig. 1 E. While the relative frequency of tomograms in the close CW contact, PM contact, and small pore connections may be taken as time proxy for the progression of the fusion process, the earliest (far CW contact) and latest stages (large fusion pore) are underestimated, as we did not systematically acquire all tomograms for these stages, which could be readily identified from an initial TEM image at high magnification. Expansion of the fusion pore also likely progresses further after disassembly of the fusion focus.

Fusion foci appeared as accumulations of dense vesicles, as expected from the localization of Myo52 and secretory vesicle markers in LM images (Billault-Chaumartin and Martin, 2019; Dudin et al., 2015). We analyzed the density and size of the vesicles from 61 randomly chosen tomograms in our dataset at the cells’ contact (see Materials and methods). Vesicle density did not change along stages (Fig. 1 F) with an average of 0.65 ± 0.22 vesicles/10^6 nm^3. Vesicle size measurements revealed two populations: the dense vesicles mentioned above with an average diameter of ~60 nm, and a less abundant population with an average diameter of ~30 nm. The larger vesicles are likely secretory (as confirmed below), whereas the smaller ones may be endocytic, as dimensions are similar to previous measurements in Saccharomyces cerevisiae (Fig. 1 G; Kukulski et al., 2012a). Both vesicle populations were present throughout the different stages, with a somewhat lower frequency of the large vesicles in early stages (Fig. S1 B). Fusion foci also showed linear structures, likely representing actin filaments, predicted from the activity of the formin Fus1, which colocalized with Myo52 in LM (Fig. 2, Video 1, and Video 2). While only 51% of tomograms revealed linear filament, which likely reflects the difficulty in preserving and detecting actin structures with this technique, the best-quality ones showed a large number of filaments positioned between vesicles and close to the PM. Actin filaments were observed in pre- and post-fusion cell pairs and could reach lengths >0.5 μm. Fusion foci were largely devoid of ribosomes and other organelles, though these were present in other parts of the tomograms (Fig. 1, A–D; and Fig. 2). Microtubules were rarely observed (12% cells), and more frequently after fusion (27% cells; Fig. S1 C). Interestingly, 52% cells showed organelles organized in sheets or reticulated with similar dense appearance as secretory vesicles (Fig. S1 D). Though we have not confirmed their identity, this is suggestive of Golgi or endosomal compartments, which may serve for local vesicle production. We conclude that fusion foci are dense vesicle- and actin-rich compartments that cannot easily be penetrated by other large cellular components.

Shape of the fusion pore(s)

To better understand the membrane merging process, we examined the shape of fusion pores. We identified a single cell pair with an apparent small pore (of diameter ~45 nm) entirely enclosed in the section (Fig. 3 A). A filament can be seen traversing the pore. Seven tomograms showed a local loss of density in the CW at the interface between cells, suggestive of continuity between the two cells’ cytosols (Fig. 3 B). However, the inner and
outer membrane leaflet could not be unambiguously resolved from the CW density to conclusively distinguish between local CW digestion and PM contact, or possible fusion intermediates or even fully formed tiny pores. In three cases, these connections between partner cells were entirely included in the tomographic volume. In the four other cases, they extended to one or both section edges, with one tomogram displaying several such connections, suggesting that membrane merging can initiate at multiple positions (Fig. 3 C, Fig. S2 A, and Video 3). However, as these connections extend to the edge of the tomographic volume, we cannot know with certainty whether they initiated independently or represent extensions of a single opening.

Figure 1. Correlative light-electron analysis of the yeast fusion site. (A–D) Virtual z-slices through tomograms of cell pairs during mating as identified by LM (see Fig. S1). Examples of different stages: far CW contact (A), close CW contact (B), PM contact (C), and after fusion (D). (E) Tomogram distribution according to minimal distance between PMs. (F) Vesicle density (n = 20 cells in far CW contact; n = 60 CW contact close; n = 10 PM contact; n = 32 after fusion, including 8 with local loss of CW density). P = 0.1293 (one-way ANOVA test). (G) Vesicle size distribution in cells as in F (n = 122 cells and 3,521 vesicles). Bin width = 5 nm. Blue asterisk = cortical smooth ER; green asterisk = rough ER; yellow asterisk = dense sheet or reticulated organelle; M = mitochondrion; V = vacuole; L = lipid droplet; N = nucleus. Scale bars, 100 nm.
Among pairs with a larger fusion pore, two classes retained our attention, as they exhibited nonspherical pore edges. Eight tomograms showed pore edges that extended into irregular extensions, such that at some cross-section of the tomographic volume, they appeared as the fusion intermediates described above (Fig. 3 D; Fig. S2, B and D; Video 4; and Video 5). Six further tomograms showed a large pore that split into two openings separated by a strand of CW (Fig. 3 E, Fig. S2 C, and Video 6). These observations suggest two non–mutually exclusive interpretations. First, initial fusion may occur at several locations, forming several small pores that fuse together to form a nonspherical larger pore (Fig. 3 F, i). Second, fusion may initiate at a single location, but pore expansion is limited by CW degradation and occurs nonhomogeneously, leading to finger-like extensions of the pore periphery (Fig. 3 F, ii). In either case, the small number of tomograms showing PM contact and small pores suggests that membrane fusion and initial pore expansion are rapid relative to CW digestion.

PM shape changes during the fusion process
To learn more about possible biophysical changes for PM merging, we focused our attention on the conformation of the PM in prefusion cell pairs. While yeast gametes are considered isogametes, we identified two morphological asymmetries at the ultrastructural level. First, the PM at the shmoo tip exhibited different levels of tension: its appearance ranged from taut, forming a smooth arc along the CW (smooth PM [sPM]), to slack/floppy, forming waves (wavy PM [wPM]). In most prefusion pairs, one cell had wPM and the other sPM (60%; 44/74), but other configurations (35% or 26/74 with both cells sPM; 5% or 4/74 with both wPM) were also observed (Fig. 4, A–C). The frequency of PM waviness, whether in one or both partner cells, increased along the fusion process and decreased after fusion (Fig. 4 D). Indeed, most post-fusion pairs showed sPM (84%; 36/43; Fig. S3 A).

Second, we found that the curvature of the zone of cell–cell contact varied, ranging from convex (positive curvature) to flat
Figure 3. **Shape of the fusion pore.** (A–E) Virtual z-slices through tomograms of cell pairs, showing one small pore entirely enclosed in the tomographic volume (A) and four cases of apparent connection between cells, with local loss of density in the CW, some of which may represent fusion intermediates (B). The CW is colored in purple. Zoomed views are shown at the bottom. (C–E) Four virtual z-slices through tomograms and model of the PM outer leaflet. Z-slice positions from left to right are indicated by purple lines on the model, where the z-slice intersects the green PM outline at a perpendicular angle (see Video 3, Video 4, and Video 6). (C) Several connections, similar to the ones shown in B. (D and E) Large pores with an irregular edge. In D, the larger pore at z = 84 nm is prolonged into a connection as in B at z = 18 and 4 nm. In E, a large pore at z = 5 nm splits into two openings separated by a strand of CW at higher z levels. The CW is colored in purple. Uncolored images are shown in Fig. S2, A–C. (F) Two non–mutually exclusive interpretations of fusion pore formation: fusion may initiate at several locations, forming pores that fuse together to form a nonspherical larger pore (i); fusion may initiate at a single location, but pore expansion occurs nonhomogeneously, limited by CW degradation (ii). Scale bars, 100 nm.
Figure 4. PM shape changes in preparation for fusion. (A–C) Virtual z-slices through representative tomograms of mating cells showing sPM versus wPM (the predominant situation; A), sPM in both (B), and wPM in both cells (C). Segmentation models of the PM outer leaflet and schematic drawings are shown at the right. (D) Percentage of cells with wPM along fusion stages (n = 26 far CW contact cells; n = 100 close CW contact; n = 22 PM contact and n = 86 after fusion). (E–G) Virtual slices through representative tomograms of mating cells showing symmetric (convex versus convex; E) and asymmetric PM curvatures (convex versus flat [F] and convex versus concave [G]). Segmentation models of the PM outer leaflet and schematic drawings are shown at the right. (H) Percentage of asymmetric mating pairs along fusion stages (n = 13 far CW contact pairs; n = 50 close CW contact; n = 11 PM contact and n = 43 after fusion). (I) Percentage of cells with sPM or wPM in prefusion pairs showing a convex–concave configuration, showing that low PM tension is associated with the concave state. n = 20 pairs. Scale bars, 100 nm.
to even concave (negative curvature). We called the situation symmetric when both partner cells showed similar positive or null curvature (Fig. 4 E), and asymmetric when the cells showed distinct curvature (Fig. 4 F and G). Asymmetry was observed in 46% (34/74) of prefusion cell pairs. In these pairs, one cell is convex while the other is either flat (Fig. 4 F) or concave (Fig. 4 G), such that one cell protrudes into the other (see also Fig. S3). Several protruding cells also showed unresolved membrane invaginations on either side of the protrusion, of unknown origin (Fig. S3, B–D). The frequency of asymmetric pairs, especially convex versus concave, increased along the fusion process, with a maximum >80% at the PM contact stage, and decreased after fusion (Fig. 4 H).

Finally, we found that PM waviness and overall curvature were highly correlated in prefusion cells: 80% of concave cells were wavy, and 95% of convex cells were smooth (Fig. 4, F, G, and I; and Fig. S3). Together, these data show that partner cells become highly asymmetric during the fusion process: one cell exhibits a taut sPM and protrudes into its partner, which exhibits a slack wPM. This asymmetry is relaxed after fusion.

**PM asymmetries are mating type–specific**

The asymmetry of prefusion partner cells suggests that each morphology may be associated with a specific mating type. To address this hypothesis, we repeated our CLEM-tomography approach using WT heterothallic strains (h+ and h−), of which only one was labeled with mCherry-D4H, a PM-localized sterol biosensor (Marek et al., 2020), to identify mating type, and with Fus1-sfGFP to identify pairs in fusion. We generated 22 tomos (15 with h+ labeled, 7 with h− labeled) at the CW contact stage. As in the h90 strain, we observed PM asymmetries in waviness and curvature: 73% of pairs (16/22) showed sPM versus wPM, and 63% (14/22) showed asymmetric curvature (Fig. S4 A). Importantly, cell morphology and mating type were correlated: 73% (16/22) of h+ cells, but only 18% (4/22) of h− cells, had wPM (Fig. 5 A and Fig. S4 B). Furthermore, in pairs with asymmetric curvature, 79% (11/14) of the convex cells were h− (Fig. 5 B and Fig. S4 C). We conclude that cell asymmetries are largely conditioned by mating type, in which the h− cell protrudes into an h+ cell with wPM.

**Cell type–specific balance of exo- and endocytosis**

The taut versus floppy PM appearance in h− and h+ cells suggests a possible change in the balance of exo- and endocytosis in the two cell types. We hypothesized that PM waves result from reduced endocytic and/or increased exocytotic rates, leading to an “excess” of PM. To test this hypothesis, we first used LM to measure the relative levels of endo- and exocytotic vesicle-associated proteins. The endocytic vesicle-associated Fimbrin (Fim1) in pairs labeled by a Myo52-tdTomato focus was sufficiently internal to unambiguously distinguish the signal from each cell. This allowed to compare Fim1-sfGFP levels in h+ and h− partners at the same fusion stage, thus eliminating potential stage-specific variations. To distinguish mating types, h+ cells further expressed cytosolic mTag-BFP2. We observed a 33 ± 12% reduction in Fim1-sfGFP levels in h+ cells (Fig. 5 C), indicating a reduced amount of endocytosis in h+ cells.

To estimate secretion, we used Myo52-tdTomato, which associates with secretory vesicles. Because the fluorescence signals from the two partner cells overlap, we analyzed Myo52-tdTomato expressed in a single mating type crossed with the other, unlabeled mating type. We observed a roughly twofold increase in Myo52-tdTomato fluorescence in h+ cells (Fig. 5, D–F). These cells also expressed Fim1-sfGFP, which reproduced the reduced endocytic signal in h− cells observed above, thus excluding artifacts due to stage-specific or experimental signal variation. Similar results were obtained using the secretory vesicle-associated exocyst subunit Exo84, which showed a 1.5-fold signal increase in h+ cells (Fig. S4 D). Thus, h+ cells show higher levels of exocytic and lower levels of endocytic signal than h− cells.

We reasoned that such a difference in vesicular trafficking between mating types should also be observed in the amount and size of vesicles detected by EM. Indeed, in the labeled h+ × h− cell pairs, h+ cells showed a 30% higher vesicle density, with an increased number of 60-nm, secretory vesicles, while the 30-nm vesicle population was slightly reduced compared with h− cells (Fig. 5 G). In our larger h90 tomogram dataset, cell type identity is unknown, but may be inferred in pairs showing clear PM asymmetries, i.e., a convex cell with sPM (inferred to be h−) mating with a concave partner with wPM (inferred to be h+). In this analysis, we obtained an ~23% increase in vesicle density in the concave/wPM, with a larger population of 60-nm, secretory vesicles and slight reduction in the 30-nm population (Fig. 5 H). Together, this set of data suggests that the difference in PM waviness between h+ and h− cells is a consequence of altered balance between endocytosis and exocytosis in the two cell types.

**PM waviness requires local exocytic activity**

To test the idea that membrane waviness is a consequence of excess exocytosis over endocytosis, we aimed to reduce local exocytosis. Deletion of Fus1, necessary for the assembly of the actin fusion focus, is predicted to prevent secretory vesicle accumulation, and thus reduce secretion locally. Because cell fusion completely fails when both partner cells lack fus1, we performed CLEM in heterothallic crosses between unlabeled Fus1−Δ cells and WT cells expressing Fus1-sfGFP and mCherry-D4H, allowing identification of pairs in fusion and mating types, respectively (n = 27 tomograms for h+ fus1Δ, Fig. 6 A, Fig. S4 E, and Fig. S5 A; n = 20 tomograms for h− fus1Δ, Fig. 6 B, Fig. S4 E, and Fig. S5 B). The density of vesicles was strongly reduced in the fus1Δ cells, whether this was h+ or h−, while WT h+ cells showed slightly higher vesicle density than h− cells, as in previous analysis (Fig. 6, A–C). The reduced vesicle density in fus1Δ is due to a strong decrease of the 60-nm vesicles, confirming that these are secretory (Fig. 6 D). We also searched for linear structures, which we presumed above to be actin filaments: in the 24 tomos in which we could identify filaments in the WT cell, only eight fus1Δ cells showed filaments near the cell–cell contact region. These results confirm at the ultrastructural level that Fus1 is necessary for formation of the actin fusion focus and the concentration of secretory vesicles. By LM, Myo52 and Exo84 also showed strong signal reduction in fus1Δ (Fig. 6 E and Fig. S4 D). Curiously, the endocytic vesicle marker Fim1-sfGFP signal...
was also slightly reduced in \textit{fus1}Δ (Fig. 6 F). Neither Myo52 nor Fim1 levels were significantly different between mating types in \textit{fus1}Δ cells (Fig. 6, E and F). Thus, local secretion at the cell–cell contact site is decreased in \textit{fus1}Δ cells, and this likely has consequences for endocytic activity.

Interestingly, the reduction in local secretion correlates with a strong loss of the wPM phenotype (Fig. 6, A and B; Fig. S4, E and F; and Fig. S5): only 4% of \textit{h+} \textit{fus1}Δ cells (1/27) showed wPM, whereas 73% (16/22) and 20% (4/20) of \textit{h+} WT showed wPM in WT × WT and WT × \textit{fus1}Δ crosses, respectively. The frequency

Figure 5. Mating type–specific PM asymmetries and vesicle distributions. (A and B) Representative examples of WT cell pairs with one mating type labeled showing wPM versus sPM (A), and convex versus concave PM curvature (B). Top left LM images show Fus1 in green and cell identity in red, based on mCherry-D4H expression. Bottom left TEM pictures show the same mating pair. A virtual z-slice of the tomogram shows PM shapes, segmented with Imod. (C) Average time-lapse projection of Fim1-sfGFP (green) and Myo52-tdTomato (red) expressed in both cells. Cytosolic mTag-BFP2 (blue) is expressed only in \textit{h+} cells. Paired Fim1 fluorescence intensities at the shmoo tips are shown on the right (\(n = 31\) pairs in two experiments; \(P < 0.0001\); paired \(t\) test). Individual data, mean ± SD, are shown. (D) Average time-lapse projection of Fim1-sfGFP (green) and Myo52-tdTomato (red) present only in one cell type (left: \textit{h+}; right: \textit{h−}) mated with unlabeled partner. (E) Fluorescence profile of Fim1 and Myo52 along the shmoo tips of cells as in D (\(n = 13\) \textit{h+} and 13 \textit{h−} cells). (F) Fim1 and Myo52 mean fluorescence intensity of the 5 maximum pixels around each peak from profiles as in E (\(n = 28\) \textit{h+} and 29 \textit{h−} cells from two experiments for Fim1 and \(n = 58\) \textit{h+} and 52 \textit{h−} cells from four experiments for Myo52; \(P\) values from Mann–Whitney tests). (G) Vesicles paired density and size distribution in prefusion WT \textit{h+}/\textit{h−} cell pairs. \(n = 14\) tomograms (264 vesicles for \textit{h−}; 350 for \textit{h+}). \(P = 0.0003\); paired \(t\) test. (H) Vesicle paired density and size distribution in prefusion WT h90 cells classified according to PM curvature and waviness. \(n = 20\) tomograms (669 vesicles for sPM/convex; 830 for wPM/concave cells), \(P < 0.0001\); paired \(t\) test. Scale bars are 5 µm in LM (1 µm for insets), 10 µm in TEM, and 100 nm in tomogram slices.
Figure 6. Local exocytosis underlies PM waviness. (A and B) Representative example of WT crossed with fus1Δ cells. The top left LM pictures show Fus1-sfGFP (green) and mCherry-D4H (red) expressed in the WT cell. Bottom left TEM pictures show the same mating pair. The virtual tomogram slices show PM shapes, segmented with Imod. Most fus1Δ cells, whether h+ (A) or h− (B), exhibit sPM. Scale bars are 5 µm in LM, 10 µm in TEM, and 100 nm in tomogram slices.

(C) Vesicle density in WT and fus1Δ prefusion cells. n = 40 cells, P < 0.0001, unpaired t test. Magenta dots are h+ cells; green dots are h− cells. (D) Vesicle size distribution from C (n = 566 vesicles for WT, 297 for fus1Δ). Bin width = 5. (E) Myo52-tdTomato average time-lapse projection fluorescence profiles along shmoo tips of h+ and h− fus1Δ cells crossed to unlabeled WT (n = 13 h+ WT, 12 h− WT, 9 h+ fus1Δ, 13 h− fus1Δ). Right: Myo52 mean fluorescence intensity of the 5 maximum pixels from profiles as on the left (n = 51 h+ fus1Δ, 45 h− fus1Δ cells from four experiments; P values are from Mann–Whitney tests). WT values are identical to Fig. 5, E and F. (F) Fim1-sfGFP average time-lapse projection fluorescence profiles along h+ and h− fus1Δ shmoo tips (n = 13 h+ WT, 12 h− WT, 9 h+ fus1Δ, 9 h− fus1Δ). Right: Fim1 mean fluorescence intensity of 5 maximum pixels from each peak of profiles as on the left (n = 28 h+ WT, 29 h− WT, 25 h+ fus1Δ, 24 h− fus1Δ)
was similarly reduced in h− cells (1/20 versus 4/22). These results are in line with the view that a higher exo- to endocytosis ratio, particularly prominent in h+ cells, is the cause of the PM waviness.

Regarding PM curvature in fus1Δ crosses, the contact region between cells was wider than in WT crosses with PMs of partner cells parallel over large surfaces (Fig. 6, A and B; and Fig. S5), consistent with a fusion delay. When the h− was WT, it was the convex one in 92% (11/12) of asymmetric pairs and could protrude into the h+ fus1Δ cell (6/12). When the h− was fus1Δ, we recovered an equal portion of pairs with a convex h− or h+ partner (6 and 6), though the asymmetry was overall poor. This suggests that the Fus1-dependent fusion focus is also important for the cell protrusive activity.

Previous measurements of fusion efficiencies in fus1Δ × WT crosses suggested that the fusion defect is more pronounced when fus1 is deleted in the h+ cell (Dudin et al., 2015). Indeed, while cell pair formation was not impaired upon fus deletion in either cell type, fusion efficiency was significantly more compromised when the h+ cell was fus1Δ (Fig. 6 G). Because the main consequence of the loss of focalized secretion in fus1Δ is the inability to concentrate glucanases for CW digestion, this finding is in line with the idea that the h+ cells exhibit a stronger secretion activity. It also this raises the possibility that loss of the wPM more directly contributes to the enhanced fusion failure in h− WT × h+ fus1Δ cell pairs.

**Cell protrusion requires strong turgor pressure**

A second hypothesis for the origin of PM asymmetries is that they may result from differences in turgor pressure in the two cell types. Specifically, higher turgor in the h− cell may enhance PM tension and facilitate protrusion into the h+ cell.

To start testing this hypothesis, we made use of the gpd1Δ mutant. Gpd1 is a glycerol-3-phosphate dehydrogenase necessary for glycerol production to restore turgor pressure in non-osmotic conditions (Bernal et al., 2014; Ohmiya et al., 1995). Cells lacking gpd1 exhibit reduced growth force when challenged in hyper-osmotic conditions (Minc et al., 2009). We repeated our ultrastructural studies in heterothallic crosses of WT expressing Fus1-sfGFP and mCherry-D4H to unlabeled gpd1Δ, generating 32 tomograms for h− gpd1Δ × h+ WT and 10 tomograms for h− WT × h+ gpd1Δ (Fig. 7, A and B; and Fig. S4 H). The frequency of PM waviness was not substantially altered in h+ gpd1Δ prefusion cells (63%; 5/8) and increased in h− cells (55%; 17/31; Fig. S4 I), and the waves were more prominent than in WT cells. LM confirmed that asymmetries in exo- and endocytosis were not altered in the gpd1Δ cells: the exocytic Myo52 signal was higher and the endocytic Fim1 signal lower in h+ cells, as in WT cells (Fig. 7, C and D). These data are consistent with a reduction in turgor pressure in gpd1Δ cells exacerbating the PM waviness phenotype.

Regarding PM curvature, cell pairs in which the h+ cell was gpd1Δ did not appear significantly different from WT. While our dataset is small, it showed the same tendencies as for WT pairs: both far CW contact cases were symmetric, and of the six close CW contact cases (PM distance = 47.5 ± 30.7 nm), two showed slight asymmetry, and three had the h− cell protrude into a concave mutant h+ (Fig. S4 J). Thus, further decrease of turgor pressure in h+ does not alter, and may even accentuate, the asymmetry between prefusion partner cells.

By contrast, when the h− cell was gpd1Δ, most cell pairs showed a symmetric (16/31) or only slightly asymmetric (8/31) curvature configuration. The remaining pairs showed a clearer asymmetry (7/31), but we did not find any example of a cell protruding into its partner. Furthermore, among all asymmetric situations, both mating types had the same probability of adopting the convex shape (7 h− versus 8 h+; Fig. S4 J and Fig. 7 B). The vast majority of cell pairs showed substantial amount of CW between cells, with only five tomograms with minimal PM distance between cells <100 nm. However, the h− gpd1Δ × h+ WT pairs did not look like the WT far CW contact class, as the CWs of the two cells had merged together over a large surface area, leading to extended contact zones between cells. This suggests that the fusion process is delayed in these cell pairs. While fusion delay may reduce the chances of observing asymmetric configurations, we note that the pairs with cells closer than 100 nm, including one with PM contact, were also symmetric or only very slightly asymmetric. These data suggest that reduction in turgor pressure in h− cells prevents protrusion into the partner cell and leads to a more symmetric arrangement between partners.

To further investigate the importance of cell pair asymmetry on fusion success, we quantified mating and fusion efficiencies of gpd1Δ cells by LM. Deletion of gpd1 in one or both mating types did not affect the ability of cells to pair together but caused opposite effects on fusion efficiency (Fig. 7, E and F): when the h− cell lacked gpd1, which we showed above results in symmetric configurations, fusion was delayed; when the gpd1Δ was deleted in the h− cell, which preserves or accentuates asymmetry, fusion occurred faster than in WT pairs. Importantly, when both partner cells lacked gpd1, fusion efficiency was restored to WT levels, indicating that the changes in fusion efficiency observed upon deleting gpd1 in only one partner cell reflect the difference in turgor pressure between the two cells rather than an absolute requirement of gpd1 in either cell type. We conclude that asymmetries in turgor pressure between cell types promote cell fusion.

**Discussion**

By using correlative light-electron tomography, we reveal ultrastructural details of the fission yeast cell fusion process. While mating yeast cells are considered isogametes, which cannot be distinguished morphologically, our study now reveals that these cells display asymmetries at the ultrastructural level:
Figure 7. Different turgor pressure between cells of a pair is required for fusion. (A and B) Representative example of h− WT cells crossed with h+ gpd1Δ cells. The top left LM pictures show Fus1-sfGFP (green) and mCherry-D4H (red) expressed in the WT cell. The bottom left TEM pictures show the same mating pair. The virtual tomogram slices show PM shapes, segmented with Imod. Scale bars are 5 µm in LM, 10 µm in TEM, and 100 nm in tomogram slices. (C and D) Myo52-mCherry and Fim1-sfGFP mean fluorescence intensities of the 5 maximum pixels around each peak from shmoo tip profiles of tagged gpd1Δ cells crossed to WT imaged as in Fig. 6, E and F (n = 22 h+, 25 h− gpd1Δ cells from two experiments; P values are from Mann–Whitney tests). (E) LM images of WT × gpd1Δ crosses at 7 h of nitrogen starvation. Fus1-sfGFP (green) and mCherry-D4H (red) are present only in WT cells. gpd1Δ cells are unlabeled. (F) Mating and fusion efficiencies over time (n ≥ 900 cells of each genotype from five to seven experiments; *, P < 0.05; **, P < 0.01 from unpaired t tests).
one cell, generally the h− partner, protrudes into the other cell, which usually exhibits a less tensed PM. Tomograms of cell pairs at or around the time of membrane merging further suggest how fusion pores form and expand during cell fusion of a walled species.

Organization of the fusion focus
The tomographic information reveals the fusion focus organization as a dense assembly of vesicles interspersed with linear filaments, largely excluding other organelles and ribosomes, in agreement with the localization of the formin Fus1, actin-associated proteins, myosin V Myo52, and other secretory vesicle-associated markers (Billault-Chaumartin and Martin, 2019; Dudin et al., 2015; Dudin et al., 2017). The filaments likely represent linear actin filaments, as the frequency of detection was reduced in cells lacking Fus1. We note that fus1Δ cells contain other actin nucleators, so complete absence of filaments is not expected. There are two sub-populations of vesicles. The lower-abundance, smaller-size vesicles may represent endocytic vesicles, but we cannot exclude that they represent a mixed population with other transport carriers. The most abundant population, with diameters averaging 60–70 nm, up to 100 nm, represents secretory vesicles, as it is strongly diminished in cells lacking Fus1, which concentrates the delivery of secretory markers at the cell growth projection (Dudin et al., 2015). Interestingly, the size of these vesicles is smaller than reported during mitotic growth, where the average diameter of secretory vesicles is 100 nm (Cheng et al., 2002; Wang et al., 2002; Wang et al., 2016). A similar observation was made in S. cerevisiae, where 60–70-nm vesicles accumulate at mating projections and fusion sites (Baba et al., 1989; Gammie et al., 1998), but 100-nm vesicles are reported in mutants of the secretory pathway during vegetative growth (Novick and Schekman, 1979; Walworth and Novick, 1987). Thus, secretory vesicles have a reduced size, and perhaps specific contents including CW digestive enzymes, during sexual reproduction.

Formation and expansion of the fusion pore
Examination of fusion pores revealed three notable features. First, the majority of fusion pores captured in our tomograms were large, extending beyond the tomographic volume, with only three pores fully captured within this volume. Thus, early stages of membrane fusion and pore expansion are rapid relative to other steps. However, the existence of mutants that exhibit transient fusion indicates that this step is not absolutely unidirectional (Vještica et al., 2018). Second, the edges of fusion pores often did not appear circular. In larger pores, this manifested by the presence of a strand of CW penetrating into the pore. In tomograms of a pore edge along the section plane, these edges were often irregular with finger-like extensions into small regions lacking CW densities. We also identified such regions at cell contacts without an obvious pore, suggesting that these represent early stages of fusion, though the resolution limits due to resin embedding do not allow unambiguously tracing both PM leaflets around these features. Third, one tomogram displayed at least two such connections distant from each other, suggesting that membrane fusion can initiate at several positions. While merging of several expanding fusion pores may be the cause of the irregular edges, a second non-mutually exclusive possibility is that the rate of CW digestion limits fusion pore expansion (Fig. 3F). As CW digestion depends on secreted enzymes (Dekker et al., 2004; Dudin et al., 2015; Dueñas-Santero et al., 2010; Martín-Cuadrado et al., 2003), secretion of individual vesicles may lead to nonhomogeneous enzyme release and cause degradation to occur at different rates around the pore edge.

Control of PM tension during yeast mating
A striking feature of prefusion cells is the wPM pattern. This pattern strongly correlates with local exocytosis. Indeed, PM waves are specifically observed at the site of cell–cell contact and are not seen in the lateral PM regions. The waves are shallower than Ω-shapes formed immediately upon vesicle–PM fusion, but the observation that they are almost completely abolished in fus1Δ cells argues that this pattern results from strong local secretion activity. In S. cerevisiae, undulating PM was also noted at the tips of mating projections (Baba et al., 1989), suggesting that this is a conserved feature during yeast sexual reproduction. The wavy pattern indicates reduced membrane tension, which does not instantaneously equilibrate with lateral membrane regions.

Interestingly, the occurrence of the wPM pattern increased through the fusion process and was more frequent in the h+ cell partner. This correlates with previous measurements showing increase in Myo52 levels in the course of fusion (Dudin et al., 2015) and with differences in the ratio of exo- to endocytic vesicles between cell types, where this ratio is higher in h+ than h− cells, as measured on both EM and LM. Exo- and endocytosis are strongly correlated and mutually influence each other through changes in membrane tension, complicating the dissection of the cause of the cell type-specific difference. Indeed, studies in several cell types have shown that high membrane tension stimulates exocytosis, which in turn leads to tension reduction (Wang and Galli, 2018). Conversely, endocytosis is inhibited by membrane tension, contributing to its increase (Joseph and Liu, 2020). We envisage two possible scenarios. First, h+ cells may have a higher secretion rate. Consistent with this view, the number of exocytic vesicles and associated markers is higher in h+ cells, and this higher signal depends on Fus1, suggesting that the cell type-associated difference is inherent to the fusion focus. Previous data had also shown a faster turnover of Myo52 in fluorescence recovery after photobleaching experiments in h+ cells (Dudin et al., 2015). However, this scenario does not explain the lower amount of endocytic vesicles in h+ cells. A second scenario is that the cell type-specific difference resides in h+ cells having a lower maximal endocytic capacity. This may cause a membrane “traffic jam,” where endocytosis cannot keep up with the exocytic capacity, leading to a local reduction of PM tension, manifested in PM waves, and a consequent decrease in the fusion of secretory vesicles with the PM, resulting in their accumulation. While further work will be required to distinguish between these possibilities, they both lead to the same overall conclusion that the balance between exo- and endocytosis is shifted toward higher exocytosis in h+ cells, which causes a local loss of PM tension at the cell front.
Control of cell protrusion by turgor pressure

Our analysis revealed a strong asymmetry in membrane curvature at the site of cell contact, which increases as the CW thins, and culminates shortly before membrane fusion, with one cell, generally the h− cell, protruding into its partner. The asymmetry is relaxed after fusion, as only few pairs with fusion pore showed asymmetry. Protruding h− cells also showed an unresolved membrane invagination on either side of the protrusion. While the cause of this structural feature is unknown, its frequency suggests it may be important to help the organization of the fusion site.

We explored the hypothesis that higher turgor pressure in the protrusive cell may cause the protrusion. Indeed, deletion of Gpd1 in h− cells largely prevents their protrusion into h+ cells and yields more symmetrical fusion pairs. By contrast, its deletion in h+ cells does not prevent h− cell protrusion. During vegetative growth, deletion of Gpd1 reduces the growth force in hyper-osmotic conditions (Minc et al., 2009). During sexual reproduction, our results suggest that gpdΔ cells have reduced growth force even without external hyper-osmotic challenge. This is consistent with Gpd1 expression, which is induced upon osmotic shock (Alba et al., 1995; Shiozaki and Russell, 1996; Wilkinson et al., 1996), but also increases about sixfold during mating (Mata and Bähler, 2006). Strong turgor is not the only requirement for protrusion, as disruption of the fusion focus in fusΔ also interferes with formation of the protrusion in h− cells. This suggests a role for the actin cytoskeleton in protrusion formation, which we hypothesize is indirect through delivery of digestive enzymes that weaken the CW. Thus, the fusion focus and higher turgor pressure in the h− cell are both required for its asymmetric protrusion into h+ cells.

Regulation of turgor pressure may be generally important to control the fusion of walled cells. In S. cerevisiae, a mutual down-regulation between the high osmolarity glycerol (HOG)–MAPK signaling and the pheromone-MAPK signaling has been described (Hall et al., 1996; Patterson et al., 2010; Vaga et al., 2014; Yamamoto et al., 2010). The pheromone–MAPK pathway also indirectly promotes HOG-MAPK signaling, by stimulating glycerol release through the glycerol efflux aquaglyceroporin Fps1 and consequent HOG-dependent glycerol synthesis to compensate for turgor loss (Balatnas et al., 2013). Interestingly, deletion of Fps1 in one, but not both, partner cell causes an increase in fusion intermediates, indicating a block or delay in fusion, though deletion of Gpd1 had no effect in this organism (Philips and Herskowitz, 1997). However, different from the situation in S. pombe, deletion of fps1 in either cell type leads to similar changes in cell contact site curvature, as observed by LM (Smith et al., 2017). Cytosolic protrusions of one cell into the other observed in the pmΔ mutant deficient in PM fusion are also reported to happen with equal frequency in either direction (Heiman and Walter, 2000). Thus, whereas our data indicate that S. pombe mating partners have different turgor pressures, the turgor pressure may be more equal between partner cells in S. cerevisiae.

Origin of fission yeast gamete asymmetry

An open question is how the asymmetry between cells comes about. Our analysis shows that ultrastructural asymmetry is linked with cell type: h+ cells have a higher exo- to endocytic ratio leading to low membrane tension, and h− cells likely exhibit increased turgor pressure, which drives protrusion, likely also contributing to membrane tension increase. These cell type-linked differences are also visible at the functional level, as gpdΔ is detrimental only in h− cells. Thus, at least part of the reason for the asymmetry between the two cells is linked to their identity.

At the transcriptional level, there are very few differences between mating types. Only 16 genes are reportedly differentially expressed, most coding for agglutinins, pheromones, processing enzymes, and receptors (Mata and Bähler, 2006; Xue-Franzén et al., 2006). Cell type–specific differences may reside in the different pheromones and receptors: though acting through the same signaling pathway, they may elicit quantitatively different signals that alter crosstalk with the osmosensing pathway. Alternatively, turgor pressure differences may be linked to one uncharacterized small gene (SPAC5656.03), which is expressed specifically in h+ cells (Mata and Bähler, 2006; Xue-Franzén et al., 2006) and promotes growth on glycerol, like gpd1 (Malecki et al., 2016). We note that gpd1 expression itself, though induced during mating, is unlikely to be the key cell type–specific difference, as deletion of gpd1 in both partner cells restores WT-like fusion kinetics and thus likely functional asymmetries.

Our analysis of heterothallic WT cells also revealed three cases of asymmetry reversal, where the h+ cell was protruding into the h− partner, indicating that asymmetry is not completely hardwired. One possibility is that rapid fluctuations in the rates of exo/endocytosis around a higher average in the h+ cell underlie the presence of both wPM and sPM patterns in both cell types. Another possibility is that the cell type–imposed asymmetry is small and can be inverted by stochastic events. Because asymmetries increase during the course of fusion, small initial asymmetries may be amplified by cell interaction through either increase in signaling or mechanical signals upon contact with the partner cell.

Function of membrane asymmetries

Cell fusion is more strongly compromised by gpd1 deletion in the h− cell and fus1 deletion in the h+ cell. These situations reduce the asymmetries observed in WT cells by preventing protrusion and PM waviness, respectively. This raises the question of a causal link between loss of asymmetry and fusion defects. Though PM waves may help initial membrane contact and merging, it is difficult to make a firm functional inference between membrane waves and fusion efficiency. Indeed, fus1 deletion or any other attempt to reduce local secretion to lessen PM waviness will also likely compromise the secretion of CW digestive enzymes. The functional importance of cell protrusion is easier to make, as gpdΔ causes fusion delays but only when solely deleted in the h− cell. In addition, gpd1 deletion only in the h+ cell, which maintains or exacerbates the protrusion asymmetry, leads to faster cell fusion. Thus, the different morphologies of the two cells are likely to promote efficient cell–cell fusion.

The asymmetry of yeast cell fusion recalls similar asymmetry observed in the fusion of Drosophila myoblasts, where fusion–competent myoblasts form a podosome that protrudes into the

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https://doi.org/10.1083/jcb.202103142
myotube (Sens et al., 2010). Podosome-like structures are also observed in other types of somatic cell fusion (Brukman et al., 2019). Although the underlying protrusive force generation may be different (Martin, 2016), the similar configurations suggest that evolution of asymmetry is beneficial to the success of cell fusion. Future work should test whether local membrane curvature in wPM helps destabilize the bilayer and lower the energy required to initiate membrane merging. Cell protrusion promoted by turgor pressure and the fusion focus, which is initially countered by the intervening CW, may, upon complete CW removal, help overcome repulsive forces to bring the cells’ PM close enough for merging. While the identity of the fusogenic machinery is still unknown, these ultrastructural PM features suggest that biophysical changes in membrane tension and curvature are important contributors to the membrane fusion process.

Materials and methods

Strains and growth conditions

S. pombe strains used in this study are listed in Table S1. Heterothallic h+ and h− strains and homothallic (h90) strains, able to switch mating types, were grown in minimum sporulation media (MSL) supplemented with nitrogen (+N). Mating assays were conducted on MSL without nitrogen (−N). Essentially as described (Vjestica et al., 2016), except for gpd1Δ strains, which were mated on malt extract (ME).

Strain construction

Strains were constructed using standard genetic manipulation of S. pombe either by tetrad dissection or transformation. For generation of fus1 deletion mutants, WT strains were transformed with a linearized deletion plasmid (based on the pFA6a-kanMX backbone) containing at least 400 bp of homology to fus1 gene flanking regions (pSM1966). For generation of gpd1 deletion mutants, the gpd1 ORF was replaced by ura4 ORF. This was achieved by PCR amplification of a fragment from strain NM204 (received from Fred Chang, University of California, San Francisco, San Francisco, California) carrying ura4+ ORF at the gpd1 locus, which was transformed and integrated in the genome of ura4-D18 strains by homologous recombination. For C-terminal tagging of myo52 with mCherry, strains were transformed with linearized fluorophore tagging plasmid containing at least 400 bp of homology to the C-terminal part of the gene, in-frame mCherry, kanMX resistance, and myo52 3’UTR (pSM2735). For tagging fim1 with sfGFP, a pFA6a-sfGFP-natMX plasmid (pSM1686) was used as a template for PCR-based targeted tagging, as described (Bähler et al., 1998).

Mating assays

For both LM and CLEM experiments, cells were first precultured overnight in MSL−N at 25°C, then diluted to OD600 = 0.025 into MSL−N at 30°C for 16 h. The amount of exponentially growing cells equivalent to OD600 of 3–5 (depending on the experiments but constant within the same experiment) was pelleted, washed in MSL−N by three rounds of centrifugation, and resuspended in 50–100 µl MSL−N. Cells were then placed in 10-µl drops on 2% agar MSL−N or 2% agar ME plates at 30°C to allow mating. Samples for CLEM were further processed after 5 h (see CLEM and tomography section below). Samples for time-lapse live imaging were transferred to MSL−N 2% agar pads after 4 h at 30°C, mounted on a microscope slide, covered with a coverslip, sealed with VALAP (a mixture of vaseline, lanolin and paraffin; Vjestica et al., 2016), incubated one more hour on the pad at 30°C, and imaged. Samples for experiments to measure mating and fusion efficiencies were placed on ME and maintained at 30°C for 5, 7, 9, 11, and 24 h, and cells were imaged at the indicated times after mounting them on slides and covered with coverslips.

LM

LM images in Fig. 7 E were obtained using wide-field microscopy performed on a DeltaVision platform (Cytiva) composed of a customized inverted microscope (IX-71; Olympus), a 100×/1.4 NA oil objective, a camera (CoolSNAP HQ2; Photometrics), or 4.2Mpx PrimeBSI sCMOS; Photometrics), and a color combined unit illuminator (Insight SSI 7; Social Science Insights). Images were acquired using softWoRx v4.1.2 software (Cytiva). Images shown are single-plane views.

Images presented in Fig. 5, C and D, or used to get data for Fig. 6, E and F; Fig. 7, C and D; and Fig. S4 D, are average projections of time-lapse videos acquired a maximum speed during 3 min using a spinning-disk microscope composed of an inverted microscope (DMi4000B; Leica) equipped with an HCX Plan Apochromat 100×/1.46 NA oil objective (PerkinElmer; including a real-time confocal scanning head [CSU22; Yokogawa Electric Corporation], solid-state laser lines, and an electron-multiplying charge coupled device camera [C9100; Hamamatsu Photonics]). Images were acquired using the Velocity software (PerkinElmer).

For CLEM, LM images for the WT h90 samples were acquired on a Nikon TE2000-E microscope equipped with a 100×/1.49 NA TIRF oil objective, a NEOL cScope DS-152Q-C00-F1 camera (Andor), and a Lambda DG-4 lamp (Sutter Instruments). Images were acquired using the NIS Elements software (Nikon). All other images were acquired on the DeltaVision platform described above, but with a UPlan Apochromat 60 ×/1.42 NA oil objective.

All LM images and time-lapse were acquired at room temperature (~22–23°C).

CLEM and tomography

CLEM was as described in Kukulski et al. (2012b), adapted for mating cells. Briefly, cells were grown for mating as described above. After washes to remove nitrogen, cells were added into MSL−N plates or ME plates for gpd1 deletion mutant strains. We allowed cells to mate for 5 h. A few microliters of MSL−N were pipetted onto the cells to form a thick slurry, which was pipetted onto a 3-mm-wide, 0.1-mm-deep specimen carrier (Wohlwend type A) closed with a flat lid (Wohlwend type B) for high-pressure freezing with a HPF100 (Leica Microsystems; for h90 samples) or a Wohlwend HPF Compact O2. The carrier sandwich was disassembled in liquid nitrogen before freeze substitution. High-pressure frozen samples were processed by freeze substitution and embedding in Lowicryl HM20 using the Leica AFS 2 robot as described (Kukulski et al., 2012b). 300-nm sections were cut with a diamond knife using a Leica Ultracut E or Ultracut UC7 ultramicrotome, collected in H2O, and picked...
Up on carbon-coated 200-mesh copper grids (AGS160; Agar Scientific). For LM, the grid was inverted onto a 1× PBS drop on a microscope coverslip, which was mounted onto a microscope slide and imaged as indicated above. The grid was then recovered, rinsed in H2O, and dried before post-staining with Reynolds lead citrate for 10 min. 15-nm protein A–coupled gold beads were adsorbed to the top of the section as fiducials for tomography. TEs were acquired on a FEI Tecnai 12 at 120 kV using a bottom mount FEI Eagle camera (4k×4k). Low-magnification images were acquired at 17.816-nm pixel size and high magnification at 1.205-nm pixel size. A few tomography acquisitions (including panel 3D) were performed on a TF20 microscope (FEI) in scanning TEM mode, with an axial brightfield detector, using a camera length of 200 mm and a 50-µm C2 aperture (Ader and Kukulski, 2017). For tomographic reconstruction of regions of interest, tilt series were acquired at 1.205-nm pixel size (Tecnai) or 1.1-nm pixel size (TF20) over a tilt range as large as possible up to ±60° at 1° increments using the Serial EM software (Mastronarde, 2005). The IMOD software package with gold fiducial alignment (Kremer et al., 1996; Mastronarde and Held, 2017) was used for tomogram reconstruction, segmentation, and presentation.

Quantification and statistical analysis

Minimal distance between PMs, and density and diameters of vesicles in tomograms were manually measured within a cylinder of diameter 1 µm centered at the center of the cells’ contact zone using IMOD software. Vesicle measurements were done for each cell separately in the half cylinder covering that cell.

Segmentation models of the PMs were done drawing the outer leaflets using IMOD software.

Waviness was assigned as follows: waves’ amplitude (A) and wavelength (λ) were measured using IMOD software. The presence of at least one wave with A ≥ 5 nm and 2A/λ ≥ 0.2 was set as a threshold for the wPM pattern.

Myo52-tdTomato or mCherry, Fim1-sfGFP, and Exo84-GFP fluorescence profiles at shmoo tips were obtained on average projections of time-lapse videos acquired at maximum speed during 3 min. Fluorescence intensity along a 7-pixel-wide segmented line was collected using the FIJI plot profile tool. Values were corrected for the external background. The curves were centered on the maximum pixel values of the Myo52 channel. Values in Fig. 5 F; Fig. 6 E, right; Fig. 6 F, right; Fig. 7, E and F; and Fig. S5 F were obtained from the average of the 5 pixels around the D’Agostino–Pearson normality test.

Online supplemental material

Fig. S1 shows examples of CLEM and ultrastructural features during mating. Fig. S2 shows the shape of the fusion pore. Fig. S3 shows FM asymmetries during cell fusion. Fig. S4 shows classification of tomograms from heterothallic crosses and additional data on distribution of exocytic vesicles. Fig. S5 shows morphology of WT × fus1 Δ crosses. Video 1 shows a tomogram and model of a cell pair prefusion. Video 2 shows a tomogram and model of a cell pair after fusion. Video 3 shows a tomogram and model of a cell pair with several connections. Video 4 shows a tomogram and model of a cell pair with an asymmetric pore. Video 5 shows a tomogram and model of a cell pair with an asymmetric pore. Video 6 shows a tomogram and model of a cell pair with a pore containing a strand of CW. Table S1 lists yeast strains used in this study.

Acknowledgments

We thank the University of Lausanne EM facility, in particular Jean Daraspe, for help with EM sample preparation and acquisition, Bruno Humbel for help with manipulation of the electron microscope, and Christel Genoud for discussions and suggestions. We thank the EM and LM facilities of the Medical Research Council Laboratory of Molecular Biology for microscopy support. We thank Fred Chang (University of California, San Francisco, San Francisco, California) for sharing a gepiΔ strain and Alejandro Meleo and Laura Merlìni for comments on the manuscript.

This work was funded by a European Research Council Consolidator grant (CellFusion) and a Swiss National Science Foundation grant (31003B_176396) to S.G. Martin, and a “poste de soutien” awarded to O. Muriel. W. Kukulski was supported by the Medical Research Council (MC_UP_1201/8). S.G. Martin also thanks the Herbette Foundation (University of Lausanne) for support of her sabbatical visit in the lab of W. Kukulski.

The authors declare no competing financial interests.

Author contributions: S.G. Martin prepared CLEM samples and reconstructed tomograms for the h90 cells with help from W. Kukulski. O. Muriel acquired EM data for all samples except for a few acquired by W. Kukulski, and prepared CLEM samples and reconstructed tomograms for all heterothallic crosses. O. Muriel performed all other experiments with technical support from L. Michon. O. Muriel analyzed the data together with S.G. Martin. O. Muriel and S.G. Martin prepared figures and wrote the manuscript, with comments from W. Kukulski. O. Muriel and S.G. Martin acquired funding.

Submitted: 22 March 2021
Revised: 15 July 2021
Accepted: 26 July 2021

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Figure S1. **Examples of CLEM and ultrastructural features during mating.** (A) LM images are GFP and tdTomato signals on 300-nm sections of resin-embedded yeast cells expressing Fus1-GFP and Myo52-tdTomato, placed on EM grids. Scale bar is 45 µm. Cells with fusion focus fluorescence signal are identified in low magnification (17.816-nm pixel size) TEM according to their position, preservation is tested at high magnification (1.205-nm pixel size) TEM, and tilt series are acquired at this same magnification and processed for tomographic 3D reconstruction. Scale bars are 10 µm in low magnification TEM and 100 nm in high magnification TEM and virtual z-slice through tomogram. BF, brightfield. (B) Size distribution of vesicles separated in the four stages: far CW contact (n = 20 cells; 616 vesicles), CW contact close (n = 60 cells; 1,865 vesicles), PM contact (n = 10 cells; 252 vesicles), and after fusion (n = 16; 788 vesicles). Bin width = 5 nm. (C and D) Virtual z-slices through electron tomograms of mating cells showing microtubule bundles crossing the fusion pore (C, yellow arrow) and organelles with similar density to secretory vesicles, organized in sheets or reticulated structures (D, blue arrows). Scale bars for EM images, 100 nm.
Figure S2. **Shape of the fusion pore.** (A–C) Original noncolored virtual z-slices through tomograms shown in Fig. 3, C and D. (D) Four virtual z-slices through a tomogram showing two large pores with irregular edges. These two pores are likely joined in a single pore outside the reconstructed tomographic volume. On the top line, the CW is colored in transparent purple to help visualization. Corresponding models of the outer leaflet of the PM are on the right. The position of each z-slice is indicated by the purple lines on the model, where the z-slice intersects the green PM outline at perpendicular angle. Z-slices are ordered from bottom (left edge of the model image) to top (right edge; see Video 3, Video 4, Video 5, and Video 6). Scale bars, 100 nm.
**Figure S3. PM asymmetries during cell fusion.** (A) Table classifying the set of tomograms from h90 WT cells used in this study, according to PM morphology along the four stages of fusion. (B–D) Virtual z-slices through tomograms of mating cells showing PM asymmetries (convex versus concave) in close CW contact stage (B), PM contact stage (C), and post-fusion stage (D). Blue arrows point to fusion pores; yellow arrows point to unresolved regions at the PM. Scale bars, 100 nm.

| h90                | N tomograms | PM layout                          | wPM both | sPM both | wPM vs sPM | Total |
|-------------------|-------------|------------------------------------|----------|----------|-------------|-------|
| far-CW-contact    | 13          | Convex vs. convex / flat vs. flat  | 0        | 9        | 2           | 11    |
|                   |             | Convex vs. flat                     | 0        | 1        | 1           | 2     |
|                   |             | Convex vs. concave                  | 0        | 0        | 0           | 0     |
| close-CW-contact  | 50          | Convex vs. convex / flat vs. flat  | 4        | 10       | 13          | 27    |
|                   |             | Convex vs. flat                     | 0        | 1        | 9           | 10    |
|                   |             | Convex vs. concave                  | 0        | 3        | 10          | 13    |
| PM-contact        | 11          | Convex vs. convex / flat vs. flat  | 0        | 0        | 2           | 2     |
|                   |             | Convex vs. flat                     | 0        | 0        | 2           | 2     |
|                   |             | Convex vs. concave                  | 0        | 1        | 6           | 7     |
| Post-fusion       | 43          | Convex vs. convex / flat vs. flat  | 1        | 32       | 4           | 37    |
|                   |             | Convex vs. flat                     | 0        | 0        | 0           | 0     |
|                   |             | Convex vs. concave                  | 2        | 4        | 0           | 6     |
Figure S4. Classification of tomograms from heterothallic crosses and additional data on distribution of exocytic vesicles. (A) Table classifying the set of tomograms from WT h+ × h− cell pairs used in this study, according to PM morphology. (B) Table showing the number of each mating type in WT h+ × h− crosses with wPM or sPM. (C) Table showing, among asymmetric WT mating pairs, the number of each mating type with convex or flat/concave shmoo tip conformation. (D) Left: Fluorescence profile of Exo84 (GFP) along the shmoo tips of labeled WT and fus1Δ cells crossed to unlabeled WT cells. Images were obtained from average projections of time-lapse videos acquired at a maximum speed during 3 min (n=16 h+ WT cells, 13 h− WT cells, 15 h+ fus1Δ cells, and 14 h− fus1Δ cells). Right: Exo84 mean fluorescence intensity of the 5 maximum pixels around peak from profiles as on the left (n=29 h+ WT cells, 24 h− WT cells, 26 h+ fus1Δ cells and 25 h− fus1Δ cells from two experiments; P values from Mann–Whitney tests). (E) Table classifying the set of tomograms from WT × fus1Δ crosses used in this study, according to PM morphology in the CW contact stage. (F) Tables showing the number of each mating type in WT × fus1Δ crosses with wPM or sPM. (G) Tables showing, among asymmetric WT × fus1Δ mating pairs, the number of each mating type with convex or flat/concave shmoo tip conformation. (H) Table classifying the set of tomograms from WT × gpd1Δ crosses used in this study, according to PM morphology in the CW contact stage. (I) Tables showing the number of each mating type in WT × gpd1Δ crosses with wPM or sPM. (J) Tables showing, among WT × gpd1Δ asymmetric mating pairs, the number of each mating type with convex or flat/concave shmoo tip conformation.
Video 1. **Tomogram and model of a cell pair prefusion.** The video shows all virtual z-sections of the tomographic reconstruction of a prefusion site at the close CW contact stage. The PM is segmented in cyan, vesicles are shown in pink (large vesicles) and purple (small vesicles), and filaments are segmented with a yellow line. The left cell shows a convex shape with a sPM. The cell on the right shows a slightly concave shape at the site of cell–cell contact with PM that exhibits an undulating pattern. Scale bar, 100 nm.

Figure S5. **Morphology of WT × *fus1Δ* crosses.** Virtual z-slices through tomograms of mating cells in CW contact stage for *h+ fus1Δ × h− WT* (A) and *h− fus1Δ × h+ WT* (B). Scale bars, 100 nm.
Video 2. **Tomogram and model of a cell pair after fusion.** The video shows all virtual z-sections of the tomographic reconstruction of a cell pair after fusion. The PM is segmented in cyan, vesicles are shown in pink (large vesicles) and purple (small vesicles), and filaments are segmented with a yellow line. Scale bar, 100 nm.

Video 3. **Tomogram and model of a cell pair with several connections.** The video shows all virtual z-sections and PM segmentation (green) of the tomographic reconstruction of a cell pair exhibiting at least two apparent connections between cells. This corresponds to the example shown in Fig. 3 C and Fig. S2 A. Scale bar, 100 nm.

Video 4. **Tomogram and model of a cell pair with an asymmetric pore.** The video shows all virtual z-sections and PM segmentation (green) of the tomographic reconstruction of a cell pair with a noncircular pore. This corresponds to the example shown in Fig. 3 D and Fig. S2 B. Scale bar, 100 nm.

Video 5. **Tomogram and model of a cell pair with an asymmetric pore.** The video shows all virtual z-sections and PM segmentation (green) of the tomographic reconstruction of a cell pair with a noncircular pore. This corresponds to the example shown in Fig. S2 D. Scale bar, 100 nm.

Video 6. **Tomogram and model of a cell pair with a pore containing a strand of CW.** The video shows all virtual z-sections and PM segmentation (green) of the tomographic reconstruction of a cell pair whose fusion pore is split in two by a strand of CW. This corresponds to the example shown in Fig. 3 E and Fig. S2 C. Scale bar, 100 nm.

**Table S1 is provided as a separate file online and lists yeast strains used in this study.**