Eisosome Organization in the Filamentous Ascomycete

Aspergillus nidulans

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Eisosomes are subcortical organelles implicated in endocytosis and have hitherto been described only in Saccharomyces cerevisiae. They comprise two homologue proteins, Pil1 and Lsp1, which colocalize with the transmembrane protein Sur7. These proteins are universally conserved in the ascomycetes. We identify in Aspergillus nidulans (and in all members of the subphylum Pezizomycotina) two homologues of Pil1/Lsp1, PilA and PilB, originating from a duplication independent from that extant in the subphylum Saccharomycotina. In the aspergilli there are several Sur7-like proteins in each species, including one strict Sur7 orthologue (SurG in A. nidulans). In A. nidulans conidiospores, but not in hyphae, the three proteins colocalize at the cell cortex and form tightly packed punctate structures that appear different from the clearly distinct eisosome patches observed in S. cerevisiae. These structures are assembled late during the maturation of conidia. In mycelia, punctate structures are present, but they are composed only of PilA, while PilB is diffused in the cytoplasm and SurG is located in vacuoles and endosomes. Deletion of each of the genes does not lead to any obvious growth phenotype, except for moderate resistance to itraconazole. We could not find any obvious association between mycelial (PilA) eisosome-like structures and endocytosis. PilA and SurG are necessary for conidial eisosome organization in ways that differ from those for their S. cerevisiae homologues. These data illustrate that conservation of eisosomal proteins within the ascomycetes is accompanied by a striking functional divergence.

In mammalian cells and in Saccharomyces cerevisiae there is cogent evidence that membrane proteins are organized in discrete domains. In the latter organism, some transporters, such as Can1p, Tat2p, and Fur1p, are organized in discrete domains on the plasma membrane. This specific domain has been named MCC, for membrane compartment occupied by Can1p on the plasma membrane. This specific domain has been characterized to be sites of lipid and protein endocytosis (53), but MCC and eisosomes are quite elusive. They were initially characterized to be sites of lipid and protein endocytosis (53), but this function is by no means certain (18). In Candida albicans, Sur7 (CaSur7) has additional roles in cell wall synthesis, actin cytoskeleton organization, and septin localization (3, 4), while in S. cerevisiae, sur7 mutants show a diminished efficiency of sporulation (54). Eisosomes are synthesized de novo in the bud during cell division (28). The Pil1/Lsp1 cytoplasmic components of the eisosome, together with the membrane protein Sur7, are conserved throughout the ascomycetes (3) (see below).

Recent work on eisosomal proteins has mainly dealt with two issues. First, both Pil1 and Lsp1 are phosphorylated. Phosphorylation is mediated by a pair of redundant kinases, Pkh1p and Pkh2p. The evidence linking phosphorylation with eisosome assembly and disassembly is, however, contradictory (25, 53). The Pkh1/2 kinases are conserved throughout the eukaryotes and strictly conserved in the ascomycetes, with one putative orthologue of the Pkh1/2 pair being present in all Aspergillus genomes sequenced. The second avenue of research is the identification of eisosome-associated proteins, including those necessary for eisosome assembly. Two recent reports have characterized a second conserved protein with four transmembrane domains, Nce102p, to be essential for eisosome assembly (15, 43). The striking conservation in all available ascomycete genomes of proteins involved in eisosome structure or assembly posits an interesting paradox. No obvious macroscopic growth phenotype is seen in S. cerevisiae cells from which PIl1, LSP1, or SUR7 is deleted in mutants with single, double, or triple mutations (52). The only report dealing with eisosomal proteins in an organism other than S. cerevisiae concerns the Sur7 homologue of Candida albicans. CaSur7 is organized in punctate eisosome-like structures. At
variance with S. cerevisiae, a deletion of the cognate gene results in a clear growth phenotype (see above) which resembles that resulting from the inhibition of β-glucan synthesis (2).

Model filamentous ascomycetes, such as Neurospora crassa, Aspergillus nidulans, Sordaria macrospora, and Podostroma antarctica, together with a host of plant and animal pathogens and an increasing number of opportunistic human pathogens, belong to the subphylum Pezizomycotina, which may have diverged from the subphylum Saccharomycotina (such as S. cerevisiae and C. albicans) between 650 million and more than 1,000 million years ago (34). All these organisms are characterized by a highly polarized growth pattern and developmental processes which include the alternation of asexual and sexual cycles involving two different types of spores: conidia and ascospores. Differently from the Saccharomycotina (see below), the sexual cycle involves a dikaryotic stage and a transient diploid, which never divides as such but which enters meiosis as soon as it is formed (see reference 55 and references therein).

A. nidulans is arguably, among the Pezizomycotina, the organism where membrane proteins and endocytosis have been better studied. In this organism, a number of transporters driven by their physiological promoters have been visualized in the cell membrane (UapC [48], PmrB [45], UapA and AzaA [35], AgtA [5], MstE [14], FcyB [51], UreA [M. Sanguinetti and A. Ramón, personal communication], and FdrD [G. Diillañas and F. Borbols, personal communication]). Load-and-chase experiments have shown that when the dye FM4-64 is endocytosed, it first appears in cortical punctate structures (36). Active research concerning membrane proteins (see reference 12 for a review), determination and maintenance of polarity during development (including the possible involvement of spinophilin [23], which may be involved in signaling eisosomal protein phosphorylation in S. cerevisiae), and ongoing recent work on endocytosis (1, 5, 6, 16, 36, 40) make A. nidulans an obvious model within the Pezizomycotina with which to study eisosomal presence, structure, and function. A description of the presence and fate of these organelles during asexual development is presented below.

(Proceed with the text as appropriate.)

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MATERIALS AND METHODS

Media and growth conditions. Minimal medium (MM) and complete medium (CM) as well as the growth conditions for A. nidulans were described by Cove (10), pH 5.5 and 8.2 MM were made using MM without salt solution and with the addition of 25 mM NaH2PO4 and 25 mM Na2HPO4, respectively. When necessary, supplements were added at the adequate concentrations (http://www.gla.ac.uk/acad/bils/molgen/aspergillus/index.html). Glucose (1%) was used throughout as a carbon source. Urea (5 mM) or ammonium L-(5-mg/ml stock solution), incubated at 25°C for 30 min, washed twice with phosphate-buffered saline (PBS), and observed under a phase-contrast fluorescent microscope with a 4′,6-diamidino-2-phenylindole filter (excitation, 365/10 nm; UV-1A filter; model DM 400 digital microscope and model BA 400 light microscope [Leica]).

Coverslips with germinated conidia were placed of plastic covers, covered with 0.1 ml of 1/1,000 dilution of CMAC (5-mg/ml stock solution), incubated at 25°C for 30 min, washed twice with phosphate-buffered saline (PBS), and observed under a phase-contrast fluorescent microscope with a 4′,6-diamidino-2-phenylindole filter (excitation, 365/10 nm; UV-1A filter; model DM 400 digital microscope and model BA 400 light microscope [Leica]).

Crosses between A. nidulans strains were carried out as described by Poncecorvo et al. (12). For selection, conidia were inoculated on solid CM or MM supplemented with the appropriate substrates and incubated at 25°C, 37°C, or 42°C for 2 to 4 days. To monitor PilA, PilB, and SurG mRNA steady-state levels in different developmental stages from ungerminated conidia to young mycelia, strains were grown on liquid MM containing 1% (wt/vol) glucose and 5 mM urca as the sole carbon and nitrogen sources, respectively, for 0, 4, 8, 12, 16, and 20 h at 25°C.

Fungal and bacterial strains. (i) A. nidulans strains. The different auxotrophic mutations of A. nidulans strains are compiled by A. J. Clutterbuck (http://www.gla.ac.uk/acad/bils/molgen/aspergillus/index.html). In particular, pianoB100, pabaA1, pabaB22, riboB2, pyroA4, pyrG99, and argB2 indicate auxotrophies for α-pantothenic acid, p-amino benzoic acid, riboflavin, pyridoxine hydrochloride, uracil/uridine, and l-arginine, respectively. The knuAΔ mutation results in a dramatically decreased frequency of heterologous integration events into the A. nidulans genome. The LO1516 (Table 1) strain expresses functional chimeric histone H1 molecules fused with the monomeric red fluorescent protein (mRFP). The VS125 (agtA::sgfp) strain expresses functional chimeric AgtA molecules fused with the green fluorescent protein (gSFP). These markers do not affect the localization of eisosomal proteins. All strains used in this work are listed in Table 1. In every case, MM indicates minimal medium supplemented with the requirements relevant to the strains used in the experiment. pabaA1 was used as the wild-type (wt) strain. The VS79 to VS81 and VS83 strains (p1439 pilB::sgfp and surG::sgfp pilC::mrfp, respectively; for complete genotypes, see Table 1) were isolated after transformation of protoplasts of the knuAΔ pyrG99 pyroA4 or knuAΔ pyrG99 riboB2 strain with the plasmid AgtA::AfpG (PYG gene of A. fumigatus), pilB::AfpG, surG::AfpG, or pilC::mrfp::AfpG (translational fusion cassettes; see below). The VS84 to VS86 (surG::AfpG) strain expresses functional chimeric proteins. For plasmid details, see below and Table 2.

(ii) Escherichia coli strains. The Escherichia coli strain used was DH5α.

Transformation methods. Transformation of E. coli was carried out as described by Sambrook and Russell (39). Transformation of A. nidulans is described by Tilburn et al. (47).

Plasmids. The PRG3 plasmid carries the radish 1S rRNA gene (11). The p1548 plasmid contains the riboB gene of Aspergillus fumigatus (44), which complements the riboB2 mutation of A. nidulans (kindly provided by M. Peñalva). The p1547 plasmid contains the pyroA gene of A. fumigatus (44), which complements the pyroA4 mutation of A. nidulans (kindly provided by M. Peñalva). The p1439 and p1491 plasmids contain a 5′ Gly-Ala (5GA) linker fused in frame with the gSFP and with mRFP, respectively, followed by the A. fumigatus pyrG2 gene (kindly provided by M. Peñalva) (44). DNA manipulations. Plasmid preparation from E. coli strains was carried out as described by Sambrook and Russell (39). DNA digestion was carried out as described by Sambrook and Russell (39). Total DNA extraction from A. nidulans is described by Lockington et al. (24). Southern blot analysis was carried out as described by Sambrook and Russell (39). The restriction enzymes used to monitor the deletion of surG, pilB, and pilA from the genomes of the VS84, VS85, and VS86 strains, respectively, were HindIII (p1439 pilA::AfpG), AfriboB, AfpyrG, and AfpyrA probes corresponding to...
the purified ∼1-kb PCR fragments were obtained using as template DNA the p1548, p1439, and p1547 plasmids, respectively, and the PilAΔ-Ribo F and PilAΔ-Ribo R, PilBα-PyrG and PilBα-PyrR, and SurGΔ-Pyro F and SurGΔ-Pyro R primer pairs, respectively (Table 3).

High-fidelity and long-fragment PCRs were carried out using an LA Taq kit (Takara). High-fidelity and small-fragment PCRs were carried out using platinum P6 polymerase (Invitrogen), while conventional PCRs were carried out using Taq polymerase (NEB). DNA bands were purified from agarose gels using a Wizard PCR Prep DNA purification system (Promega). The [32P]dCTP-surfG riboB using pyrG (VS80 pyrG89 agtA) strain by the VS125 pyrG89 argB2 surfG, and VS186 pyrG89 pilBα::AfpyroA surfG argB2 nkuA::Afribo+ riboB2 consequently to the 3′ amplified from the p1439 and p1491 plasmids. These fragments contain a 5GA ORF, followed by the 5GA linker, while the central part contains the pyroA gene of A. fumigatus amplified from the p1548 plasmid using primers SurGα-Pyro F and SurGα-Pyro R. The sequences 1,112 bp upstream and 1,014 bp downstream of the pilA ORF were amplified from genomic DNA of a wild-type strain (pasa1) using the P1 PilAα-P3 PilAα and P4 PilBα-P6 PilAα primer pairs, respectively. The sequences 1,112 bp upstream and 1,014 bp downstream of the pilB ORF were amplified from genomic DNA of a wild-type strain (pasa1) using the P1 PilBα-P3 PilBα and P4 PilBα-P6 PilBα primer pairs, respectively. The sequences 973 bp upstream and 1,073 bp downstream of the surG ORF were amplified from genomic DNA of a wild-type strain (pasa1) using the P1 SurGα-P3 SurGα and P4 SurGα-P6 SurGα primer pairs, respectively. The whole pilA, pilB, and surG deletion cassettes were used to transform the nkuAΔ-pyrG89 riboB2 and nkuAΔ pyroA4 strains were amplified using the P2 PilAα-P5 PilAα, P2 PilBα-P5 PilBα, and P2 SurGα-P5 SurGα primer pairs, respectively (Table 3). Selection of transformants was carried out on a medium containing minimal medium lacking riboflavin, uracil, uridine, or pyrimidine, as required for each replacement. The in-locus replacement of pilA, pilB, and surG, respectively, was confirmed by Southern blot analysis (data not shown). The deleted outcrossed strains were checked by PCR using specific primer pairs P2 PilAα-P5 PilAα for the pilA strain, P2 PilBα-P6 PilBα for the pilB strain, and P2 SurGα-P5 SurGα for the surG strain, respectively (Table 3).

Construction of pilA, pilB, and surG in-locus fusions. Cassettes containing the pilAαsgfp, pilBαsgfp, surGαsgfp, pilAαmrfp, and pilBαmrfp sequences were constructed by joining three different PCR fragments, as described by Szewczyk et al. (44), using the PilA, PilB, and SurG pairs of primers (Table 3). To construct the pilAαsgfp, pilBαsgfp, surGαsgfp, pilAαmrfp, and pilBαmrfp translational fusions, DNA fragments corresponding to the central part of the construction were amplified from the p1439 and p1491 plasmids. These fragments contain a 5GA linker fused in frame with the sgFP and the mRFP proteins, followed by the pyroA gene of A. fumigatus. The upstream fragmenting sequence of the sgfp and mrfp ORFS is a 1,128-bp fragment containing a part of the pilA ORF followed by the 5GA linker, while the downstream fragmenting sequence is a fragment containing a 1,014-bp fragment corresponding to the 3′ end of the pilA gene, just after the chain termination codon. The upstream fragmenting sequence of the mrfp ORF is a fragment containing 1,105 bp of the pilB ORF, followed by the 5GA linker, while the downstream fragmenting sequence is a fragment containing 1,112-bp fragment corresponding to the 3′ downstream region of the pilB gene, just after the chain termination codon. The upstream fragmenting sequence of the mrfp ORF is a 1,157-bp fragment of the surG ORF, followed by the 5GA linker, while the downstream fragmenting sequence is a fragment containing a 1,073-bp fragment corresponding to the 3′ end of the surG gene, just following the chain termina-

### TABLE 2. Plasmids used in this work

| Cloning vector | Description | Source, yr (reference) |
|----------------|-------------|-----------------------|
| pRG3          | pGEM with 18S rRNA gene | Delcasso-Tremouly et al., 1998 (11) |
| p1548         | AfriboB     | Szewczyk et al., 2006 (provided by M. Peñalva) (44) |
| p1547         | AfpyrA      | Szewczyk et al., 2006 (provided by M. Peñalva) (44) |
| p1439         | 5GA:sgfp-AfpyrG | Szewczyk et al., 2006 (provided by M. Peñalva) (44) |
| p1491         | 5GA:mrfp-AfpyrG | Szewczyk et al., 2006 (provided by M. Peñalva) (44) |

* See also Materials and Methods.
tion codon. All fragments were amplified from genomic DNA of a wild-type strain (paba1Δ) using the P1-P3 (PilA, PilB, or SurG) and P1-P4 (PilA, PilB, or SurG) primer pairs. The whole fusion cassettes were amplified using the P2-P5 (PilA, PilB, or SurG) pair of primers and were used to transform nkaApyrG98 ribo2 (pilA::sgfp, pilB::sgfp, surG::sgfp, and pilB::mrfp) or nkaApyrA4 (pilA::mrfp) strains. Transformants were selected on MM with ura as the sole nitrogen source and without uracil/uridine. The in–locus replacement of pilA, pilB, and surG with pilA::sgfp, pilB::sgfp, surG::sgfp, or pilB::mrfp was confirmed by Southern blot analysis. Genomic DNA was restricted with EcoRV (pilA::sgfp, pilA::mrfp), NcoI (pilB::sgfp), or PsI (surG::sgfp) and hybridized with the Arabidopsis, AlpyrG, and AlpyrA sequence fragments, respectively. These fragments were amplified from plasmids p1439, p1548, and p1547 using the primer pairs ribo F-Ribo R with “pilA-Ribo F with PilB-Ribo R,” “pilA-Ribo F with PilB-PyrG R,” and “pilB-PyrG F with SurG-Pyr R,” respectively (Tables 2 and 3). All transformants checked by Southern blot analysis contained single-copy sequence integrations at the pilA, pilB, and surG loci; and their sequences did not show any differences from those of the wild-type controls under any of the conditions tested (see Results).

RNA manipulations. Total RNA extraction from A. nidulans was carried out using the TRIzol reagent (Invitrogen), according to the manufacturer’s instructions. RNA was separated on glyoxal agarose gels, as described by Sambrook and Russell (39). The hybridization technique is described by Church and Gilbert (9). To monitor RNA loading, the radish 18S rRNA gene was used as the probe (11). The 15-kb EcoRI-EcoRI fragment purified from plasmid pR32, pilA, pilB, and surG mRNA steady-state levels were monitored by hybridization with probes corresponding to the purified 15-kb PCR fragments obtained using as the template DNA from a wild-type (paba1Δ) strain and the P1-P3 (PilA, PilB, and SurG, respectively) primer pairs.

Membrane protein extraction and Western blot analysis. Protein extracts were prepared as described by Kafasla et al. (21). In particular, mycelia grown for 16 h in MM containing 5 mM ura as the sole nitrogen source and supplemented with the appropriate auxotroph and cinnamidopine (0 h of growth) were harvested, ground in liquid nitrogen. All subsequent steps were carried out at 4°C. Conidia and mycelial powder was resuspended in 1.5 ml ice-cold extraction buffer (50 mM Tris- HCl, pH 7.4, 150 mM NaCl, 5 mM EDTA, pH 8.0) supplemented with a protease inhibitor cocktail (Sigma) and 0.2 mM phenylmethylsulphoxi-

| Name            | Sequence (5′–3′)                                                                 |
|-----------------|---------------------------------------------------------------------------------|
| PilΔ-Ribo F     | GCA GAA TAT CGG CTG TC CGC TCT AGA ACT AGT GGA TCC                               |
| PilΔ-Ribo R     | GCT CAT TCA GTG AGT GCT CG CCT CGA GGT CGG TAT CG                                |
| PilB-RyrG F     | CAC CTG TCT AGC TGAG AAC GTC TCA TGG TCA AAC AAT GCT CT T                       |
| PilB-RyrG R     | CAC AAA GCA CTA ATG ACC SGC CGG TCT AGA GAG GGC ACT GTG CAG                     |
| SurG-Pyro F     | GTC TAC TCG CCT CTC ATC AGA CGG TCT AGA ACT AGT GGA TCC                         |
| SurG-Pyro R     | GGT TAC ATG ACG CGA CAG CAG CAG CCT CGA GGT GTG CGA CGG TAT CG                  |
| P1 PilΔ        | CGA TCC TAG CTC TCA GGA TC                                                      |
| P3 PilΔ        | GGA TCC AGT TCT AGA GCG GAG ACC ACG CGA TAT TCT GC                              |
| P4 PilΔ        | CGA TAC CGT CGA CCT CGA GG GGA CGA CTC ACT GAA TGA GC                           |
| P6 PilΔ        | ACC AAC CTA GTG GAC GTG AC                                                      |
| P1 PilB        | CTC AGC TGA GTG ACT GTG AC                                                      |
| P3 PilB        | GAA GAG CAT GTG TCG AGG CTA ATG CAG AGG TGG TTA GTA GTG CT GT TGT G             |
| P4 PilB        | GCA TCA GTG CCT CTC ATC AGG GTG GGA ACT AAG                                     |
| P5 SurG        | CGT CCT CAT CCG TGT CTC                                                       |
| P6 SurG        | GAA GAC ACC GTG TCA ACC GTG GC                                                  |
| P1 SurG        | GCT CAT ATC ACC ACT GAT TAG                                                     |
| P3 SurG        | GGA TCC AGT TCT AGA GGA TCG TAT GAG GAG AGA GTA GAC                             |
| P4 SurG        | CGA TAC CGA CCT CGA GG TGC TGT CG TGT CG TCG CAA ATC                             |
| P6 SurG        | GAC TGC CAC ACC TCA GCT                                                        |
| P1 PilB        | TCA TCA CGG AAG ATC ATC                                                        |
| P3 PilB        | CAT GTG CAT GCT CAC AG                                                         |
| P2 SurG        | TCT GTG GAT GGA GGA ACT AAG                                                     |
| P5 SurG        | CGT CCT CAT CGG TGT CTC                                                        |
| PilA F         | GGG GAG CGC CCT CGA GAG GGT GTG TCA GCA CTT AGA CAA ATC                          |
| PilA R         | GCT CAT TCA GTG AGT GCT CG ACT AGA GGA GCC ACT GAT GC                           |
| PilA P1        | GCA TCG TAC ATC CTC TAT GCG                                                     |
| PilA P2        | CAC CAC TCT CCT CGA CCA AG                                                     |
| PilA P3        | CAG CGC CTG CAC CAG CTC C GAC AGG GAC ACC CCG CCG CCG CCG CCG                  |
| PilA P4        | GCA TCA GTG CCT CTC ATC AGA GCC GAG TGA TGT CG CTA GAG AGG TGG TGT CTG CTA      |
| PilA P5        | GCA TCC ATG ATG TCA CGA TAC                                                    |
| PilB F         | AAC ACC CTA GTG GAC GTG AC                                                      |
| PilB R         | ACG GAG TCG TAC GTC CCA TT GGA GGT GCA GTT CCA GGT CAC                         |
| PilB P1        | GCA TCG TAC ATC CTC TAT GCG                                                     |
| PilB P2        | CGT GCT GAT GGC AGA AAT GG                                                     |
| PilB P3        | CAG CGC CTG CAC CAG CTC C AAT GGG CAC TGC AAC TCG CT                             |
| PilB P4        | GCA TCA GTG CCT CTC ATC AGA AAC GAG TGA TTA GTG TTT GTT G                        |
| PilB P5        | CAT GTG CTC CAT GCT CAC AG                                                      |
| PilB P6        | TCG TCA GAG TAC TAG AGC                                                         |
| SurG F         | GCA ACA AGG AAA TCG CTC CCG GAG CTG GTGCAG GCG CTG GAG                           |
| SurG R         | GTG TAT AGC CGA CAG CAG CAC TGA GAG GGA CTA GTG CTA GAG                          |
| SurG P1        | CTT CAT CTT GCA ATC AGC TAC                                                    |
| SurG P2        | TCT AGT CTG TCA TCA TCA GAG                                                     |
| SurG P3        | GCC TGG GTT ACC ACC AGG GCC GTG TCT CGT GTT GAG                                  |
| SurG P4        | CAC GCA CCT GTG CTC CTC AGT CGG TCG CTA CAA TAC CTT GAG                         |
| SurG P5        | TGG CTG TCT GCC TCC CTC                                                        |
| SurG P6        | GAC TGC CAC ACC TCA CCT C                                                      |
fonyl fluoride (PMSF). A volume of ~200 µl sterile glass beads (diameter, 0.1 mm) in conidia extraction buffer was added, and the suspension was vortexed for about 2 min. After 10 min incubation on ice, unbroken cells and larger cell debris were removed by a short low-speed centrifugation (3,000 x g for 5 min). Tri-chloroacetic acid was added to a final volume of 5%, followed by 10 min incubation on ice. Total proteins were precipitated by a 5-min centrifugation at 13,000 x g, washed with 500 µl Tris base, and resuspended in extraction buffer. The sample protein concentration was measured by the method of Bradford (7a).

The sample proteins (20 µg) were fractionated on a 10% SDS-polyacrylamide gel and electroblotted (Mini Protean Tetra cell; Bio-Rad) onto a polyvinylidene difluoride (PVDF) membrane (Macherey-Nagel) for immunodetection. The membrane was treated with 2% nonfat dry milk, and immunodetection was performed using a primary mouse anti-GFP monoclonal antibody (Roche) and a secondary goat anti-mouse IgG horseradish peroxidase (HRP)-linked antibody (Cell Signaling). Blots were developed by the chemiluminescent method using an enhanced chemiluminescence reagent (Amersham Bioscience).

FM and LSCM. Ten milliliters of a suspension of 5 x 10^5 or 5 x 10^6 conidia/ml was inoculated onto sterile coverslips embedded into appropriate liquid culture media, incubated for 4 h or 16 h at 25°C, and observed by fluorescence microscopy (FM) and laser scanning confocal microscopy (CSLM), as described previously (45, 46). Two hundred microliters of a suspension of 5 x 10^6 conidia/ml was inoculated into sterile Eppendorf tubes and observed by FM and CSLM, as described previously (45, 46). To elicit AgtA-GFP endocytosis, ammonium tetrathionate was added to a final concentration of 10 mM at 30, 60, and 120 min before observation. Filipin was used at a concentration of 25 µg/ml at 37°C for 15 min before the observation. Samples were observed on an Axioplan Zeiss phase-contrast epifluorescent microscope with appropriate filters, and on the resulting images were acquired with a Zeiss MRC digital camera using AxioVision 4.0 (version 4.40.0) software. Images were then processed with Adobe Photoshop CS2 (version 9.0.2) software. Confocal scanning laser microscopy to examine PilA-sGFP, PilB-sGFP, SurG-sGFP, PilA-mRFP, PilB-mRFP, and HhoA-mRFP localization was carried out on an MRC 1024 confocal system (Laser Sharp, version 3.2 software [Bio-Rad]; zoom, x 2 to x 5; excitation, 488 nm/blue and 568 nm/yellow for sGFP and mRFP, respectively; samples at laser power 30; [PilA, SurG, and HhoA] and 30% [PilB]; Kalman filter; n = 5 to 6; 0.3- µm cut; iris, 7 to 8; crypton/argon laser; Nikon Diaphot 300 microscope with a x >60 [oil immersion] lens; emission filter, 522/DF35; lens reference, Plan Apo 60/1.40 oil immersion DM 160175, 60 DM/Ph4, 160/0.17 [Nikon, Japan].

Bioinformatic tools and databases. The databases consulted were http://www.lgcnet.net, through which most fungal genomes are accessible. Specifically, Aspergillus sequences were obtained through http://www.broadinstitute.org/annotation/genome/aspergillus_group/MultiHome.html. The yeast databases were consulted were http://www.genolevures.org/yeastgenomes.html, http://www.yeastgenomes.org/, http://www.gendb.org/gendb/pombe/, and http://www.candidagenome.org/. The Pneumocystis carinii database is http://pgp.cchmc.org/. Phylogenetic trees were constructed online using the different programs contained in http://www.phylogeny.fr/. Muscle alignments were carried out with http://ebi.ac.uk/Tools/muscle/index.html. T-Coffee alignments were carried out at http://www.phylogeny.fr/. Alignments were carried out with the Boxshade program (http://www.ch.embnet.org/software/BOX_form.html). Transmembrane protein topology were predicted with the HHMTOP (http://www.enzim.hu/hhtmtop/html/submit.html), PRED-TMR (http://athina.biol.uoa.gr/PRED-TMR2/input.html), TopPred (http://www.ch.embnet.org/software/TMPRED_form.html), and SPLIT4 (http://split.pmfst.hr/split/4) programs. Coiled coils were predicted at http://toolkit.tuebingen.mpg.de/coils and http://npsa-pbil.ibcp.fr/cgi-bin/npsa_automat1.pl?page=npsa_lupas.html.

RESULTS

Eisosomal core components in Aspergillus. A search of the available fungal databases reveals that homologues of PilA and Lsp1 are present in all the ascomycetes (properly, phylum Ascomycota). In the available genomes of all the Pezizomycotina, including the aspergilli, we found proteins belonging to two clades, which we call the PilA and the PilB clades. Very stringent conservation is seen within the PilA clade, while the proteins of the PilB clade show a higher rate of divergence. Proteins of the PilA clade show a central coiled-coil domain (residues 166 to 198 for PilA), similarly to Pil1 (52) and Lsp1, while all proteins of the PilB clade show the presence of an additional carboxy-terminal coiled-coil domain (residues 147 to 178 and 225 to 244 for PilB; data not shown). The PilA and PilB clades are represented in A. nidulans by the sequences with the designations ANID_05217.1 and ANID_03931.1, respectively. The duplication which generated PilA and PilB is possibly ancestral to the Pezizomycotina, as orthologues of ANID_05217.1 (75% identity) and ANID_03931.1 (56% identity) are present in Tuber melanosporum (R. Percudani, personal communication), a member of the Pezizales, which is a basal branch of the Pezizomycotina (42). The PilA clade clusters with the Pil1-like and Lsp1-like proteins of the subphylum Saccharomycotina, with the latter arising from an independent duplication event at the root of the Saccharomycotina. For clarity, the results for proteins from only two genomes (S. cerevisiae and Kluyveromyces lactis) of the latter subphylum, together with the results for proteins from some representative genomes of the Pezizomycotina, are shown in Fig. 1. A third duplication independent of those of the Saccharomycotina and Pezizomycotina is present in all the available genomes of the genus Schizosaccharomyces, represented in Fig. 1 by S. pombe. Two close homologues showing 64% to 42% identities with the two S. pombe proteins, are also present in Pneumocystis carinii, another member of the Taphrinomycotina (data not shown). The homologues present in the Taphrinomycotina are quite divergent from Pil1/PilA, but their position in the tree is consistent with the proposal that the ancestral Pil protein could have been Pil1/PilA-like. This topology, indicating the occurrence of three independent duplications, one for each subphylum, is maintained when all available genomes are included in trees constructed with a variety of algorithms (32, our unpublished data; A. Olivera-Couto and P. Aguilar, personal communication). The Meu14 protein of S. pombe, necessary for the second division of meiosis and the accurate formation of the forespore membrane (30), is clearly related to Pil1/PilA/Lsp1 (22% identity with both Pil1 and PilA, 24% identity with PilB) (31). When it is included in trees similar to that shown in Fig. 1, it appears as an outgroup to both the Pil1/Lsp1/PilA clade (including the S. pombe putative eisosomal proteins) and the PilB clade (data not shown) (32). The Sur7 transmembrane protein has been characterized as a component of eisosomes in both S. cerevisiae and C. albicans (see Introduction). Three putative members of the Sur7 family of proteins are present in A. nidulans, ANID_04615.1, ANID_01331.1, and ANID_05213.1. These proteins represent each of the three clades of Sur7-like proteins present in the aspergilli (see Fig. S1 in the supplemental material). ANID_04615.1 (to be called SurG, whose cognate gene is surG) is clearly the orthologue Sur7 protein of S. cerevisiae and C. albicans (showing 21% and 27% identities, respectively). The positions of the four transmembrane domains, typical of Sur7, are conserved in the three homologues, together with the sequence showing similarities with the Claudin-like domain (3).

The subcellular localization of eisosomal proteins is developmentally regulated. To determine the subcellular localization of PilA, PilB, and SurG, we used strains carrying the pilA-gfp and pilA-mrpf, pilB-gfp, and surG-gfp alleles replacing the corresponding resident genes, respectively. The GFP fusion strains behave like the wild type in relation to growth on complete medium in the presence or absence of itraconazole.
FIG. 1. Eisosomal proteins in the subphylum Pezizomycotina. Phylogenetic unrooted tree, including the Pil1 (YGR086C) and Lsp1 (YPL004C) proteins of S. cerevisiae; the two putative eisosomal proteins of the second member of the Saccharomycotina, Kluvyermices lactis (KLLA); a member of the Taphrinomycotina, S. pombe (SPAC3C7.02c, SPCC736.15); and the PilA and PilB homologues (see text) of a number of representative fungi of the Pezizomycotina. The proteins of A. nidulans (Eurotiomycetes) referred to are PilA (ANID_05217.1) and PilB (ANID_03931.1). Other species referred to are Botritis cinerea (BC) (Leiotiomycetes), Coccidioides immitis (CIGH; Eurotiomycetes), Magnaporthe grisea (MGG), Neurospora crassa (NCU; Sordariomycetes), and Stegnospora nodorum (SNOG; Dothideomycetes), followed by their systematic numbers in the relevant databases. Alignments were carried out with T-Coffee, curated with the Phylogeny site internal curation program (see Materials and Methods), and the tree was obtained with the maximum-likelihood program allRT-PhyML (4a; http://www.phylogeny.fr/version2.cgi) and redrawn after a Drawtree image. The digits at the nodes represent the results of the approximate likelihood ratio test and are nonparametric branch support values (http://atgc.lirmm.fr/alrt/). The PilA clade is clustered with all the homologues of the Saccharomycotina; in trees (data not shown) where many more species of Saccharomycotina and Pezizomycotina are included, the topology of a tight cluster of PilA homologues and a looser cluster of PilB homologues is maintained in a tree that was constructed with the same or other algorithms and that includes at least one species of every genus of the Pezizomycotina for which there is an available sequence. Arrows show the nodes corresponding to the three independent duplications that gave origin to the PilA and PilB clades of the Pezizomycotina, the Pil1 and Lsp1 clades of the Saccharomycotina, and the two clades of the Taphrinomycotina.

(see below). It was also checked that the temporal pattern of expression of the mRNAs corresponding to each of the gfp fusion proteins is identical to that seen for the wild-type genes (data not shown; see below).

Epifluorescence microscopy of ungerminated conidia (Fig. 2) showed PilA-GFP, PilB-GFP, and SurG-GFP localized as patches in the periphery of the spores. SurG-GFP is additionally localized perinuclearly. The perinuclear location of SurG is more clearly shown by confocal microscopy in a strain carrying both SurG-GFP and the H1 histone gene fused to mRFP, suggesting localization of SurG to the endoplasmic reticulum, similar to that shown by the endoplasmic reticulum chaperone ShrA (Fig. 2B) (13). Polar and equatorial sections of confocal microscopic images of ungerminated conidia showed that the PilA-GFP patches are not as clearly separated in A. nidulans as they are in S. cerevisiae (see Fig. S2 in the supplemental material).

No change in the localization of PilA or PilB is seen during the period of isotropic growth. The same stands for the peripheral localization of SurG; nevertheless, the perinuclear ring disappears before the emergence of the germination tube (this is better seen in Fig. 9 [see below]). All three proteins are restricted to the periphery of the conidial head opposite the emergent germ tube, while they have different destinations in young and older mycelia. PilA is present in discrete patches in young mycelia (16 h; Fig. 2) but also older and branching mycelia (20 and 24 h; data not shown). PilA spots in hyphae are not uniform in size and are not restricted to the periphery as they are in conidiaospores. This is shown in Fig. 3, where the brightest PilA spots are localized at the periphery of the hyphae, while smaller ones are present both at the interior of the hyphae and at its periphery. Furthermore, double labeling of PrnB (the major proline transporter) (41, 45) and PilA indicates that hyphal membrane-bound PilA punctate structures (which we can equate to eisosomes) are localized to the internal face of the membrane (Fig. 3E). In hyphae, PilB-GFP is seen at very low intensity, diffused fluorescence in the cytoplasm (Fig. 3A), the settings of the microscope being such that the difference in fluorescence between the ungerminated conidia and the germings is faithfully reflected. This suggests that the protein is downregulated after germination. PilB-GFP is excluded from the nuclei, as shown in a strain carrying both PilB-GFP and the gene encoding the H1 histone fused to mRFP (Fig. 3B). Finally, in hyphae SurG-GFP is confined to the vacuole and endosomes (Fig. 3A, black arrows) (some residual signal can be seen in the membrane of the conidial head), and this is confirmed by CMAC staining (data not shown).

Expression of pilA, pilB, and surG genes. The accumulation of the three eisosomal proteins in conidia led us to investigate whether this is correlated with transcription of the cognate genes, from resting conidia through young mycelia. The results presented in Fig. 4A show that both pilA and surG mRNAs accumulate more in resting conidia that at any other time point of germination tested. At variance with this, pilB mRNA is abundant in resting conidia and is not detectable within the sensitivity of the Northern blot at 4 h, 8 h, 12 h, and 16 h after the onset of germination. The transcription of the pilB homologue of A. fumigatus, Afu6g08320, is also clearly detected in
dormant conidia and is barely detectable at the onset of germination (22).

The fluorescence signal (Fig. 2) seen in germlings and mycelia for PilB (cytosolic) and SurG (vacuolar) could be due either to the presence of the intact fusion proteins, to degradation products conserving the proteins’ C termini, or simply to degradation-resistant GFP accumulating in the cytosol and vacuoles, respectively. Western blots (Fig. 4B) carried out with protein extracts of ungerminated conidia (0 h) or young mycelia (16 h) grown under the same conditions used to investigate intracellular localization show that bands corresponding to full-length PilA-GFP and PilB-GFP are present in both conidia and mycelia, while bands corresponding to full-length SurG-GFP are present in conidia and are faintly observed in mycelia. Molecular masses are indicated in kDa. The slower-migrating bands of PilA-GFP and PilB-GFP have approximate apparent molecular masses of 70 kDa and 80 kDa, respectively (calculated molecular masses, 67 kDa and 71.5 kDa, respectively). The PilA and PilB bands running just below them have apparent molecular masses of 62 and 72 kDa, respectively. We cannot conclude from these data if N-terminal proteolysis and/or other posttranslational modifications are responsible for the presence of these additional electrophoretic species, together with those seen between 48 and 27 kDa of the protein marker. The apparent molecular mass of SurG-GFP is about 48 kDa (calculated molecular mass, 54 kDa), a difference which is not surprising for a protein comprising a highly hydrophobic moiety. A band with an apparent molecular mass of 27 kDa corresponding to free GFP is seen only in strains carrying the SurG-GFP fusion. This, together with the down-regulation of the full-length SurG-GFP observed after 16 h of growth, demonstrates that the vacuolar fluorescence detected in the mycelia of SurG-GFP strains (Fig. 2A) mostly derives from free GFP and degradation intermediates.

**Eisosome proteins assemble during conidiogenesis.** The presence and colocalization (see below) of the three eisosomal proteins in ungerminated conidia led us to investigate their appearance during conidial development. Figure 5 shows that during conidiogenesis the three proteins are present in late, mature conidia (Fig. 5B and 5C for PilA-GFP, Fig. 5G for PilB-GFP, and Fig. 5H for SurG-GFP). PilA spots can be detected in the stalk of the conidiophore, the vesicle, metulae, and phialides (Fig. 5A). Differently from PilB and SurG, which are seen only in mature, older conidia on the conidiophore, the appearance of PilA in conidia budding from the phialides is quite variable from one conidiophore to the other and even within the same conidiophore. Figure 5D and F shows two extremes in which none and almost all of the emerging conidia express PilA-GFP, respectively. PilA spots can be detected in the stalk of the conidiophore, the vesicle, metulae, and phialides (Fig. 5A). Differently from PilB and SurG, which are seen only in mature, older conidia on the conidiophore, the appearance of PilA in conidia budding from the phialides is quite variable from one conidiophore to the other and even within the same conidiophore. Figure 5D and F shows two extremes in which none and almost all of the emerging conidia express PilA-GFP, respectively. While the micrograph in Fig. 5E shows an intermediate situation, with only some of the budding conidia expressing PilA-GFP. Control strains which did not include any GFP fusion protein do not show any con-
idiophore or conidiospore fluorescence under the same observation conditions (data not shown).

Colocalization of PilA, PilB, and SurG in quiescent conidia. PilA, PilB, and SurG are localized at the periphery of resting conidia. We thus investigated their colocalization. To this aim we constructed strains carrying PilA-mRFP and PilB-GFP or SurG-GFP. Figure 6 shows colocalization of PilA with PilB and colocalization of PilA with SurG in the periphery of the conidia; there is obviously no colocalization of PilA with the perinuclear fraction of SurG. The PilB fluorescence intensity more consistently follows the peaks of PilA fluorescence than those of SurG. It can be seen from both Fig. 2 and Fig. 6 that the localization pattern of SurG is less discontinuous and punctuate than that of PilA and PilB.

Phenotypic characterization of mutants with deletions. (i) Growth phenotypes. We have constructed strains with pilA, pilB, and surG deletions (see Materials and Methods). A strain with both pilA and pilB deletions was also constructed. No growth phenotype at 25°C, 37°C, and 42°C was seen for any of the strains with deletions on either complete or minimal medium supplemented with urea, ammonium, or nitrate as the sole nitrogen source (data not shown). Moreover, conidia from strains with a pilA, pilB, or surG deletion exhibited swelling and polarity establishment (time of germination tube appearance) indistinguishable from that of a wild-type strain at 25°C, 37°C, and 42°C. This was seen after incubation of conidia in minimal medium supplemented with urea as the sole nitrogen source for 2 h, 4 h, and 6 h under the conditions described above and observation of differential interference contrast (DIC) images using a phase-contrast fluorescent microscope (data not shown). As eisosomal proteins are localized in quiescent conidia, we investigated whether they are important for conidial survival. The conidial survival for strains with a pilA, pilB, or surG deletion was not significantly different from that for the

FIG. 3. (A) Confocal z-stack sections showing PilA-GFP in a wild-type strain. The strain was grown in the presence of 5 mM urea and 1% (wt/vol) glucose as the sole nitrogen and carbon sources, respectively, for 16 h at 25°C. (B to D) Inverted black and white fluorescence of the first z-stack section merged to the corresponding DIC. Bar, 5 μm (A to D). Note that PilA spots are not uniform in size and are not restricted at the periphery of mycelia. The largest PilA eisosomes (filled arrows in panel D) are localized at the periphery, while the smaller ones are at both the interior and the periphery (dashed arrows). (E) Subcellular localization of PilA and PrnB proteins in mycelia. Representative pictures from laser scanning confocal microscopy of strains expressing both PilA-mGFP and PrnB-GFP molecules in young mycelia (12 h) are shown. The upper right inset in the Merge panel shows a magnification of the boxed region. Strains were grown in the presence of 5 mM urea and 1% (wt/vol) glucose as the sole nitrogen and carbon sources, respectively, at 25°C. To induce prnB gene expression, 20 mM l-proline was added during the last 2 h of growth (45). Bar, 5 μm.
wild-type conidia after incubation of freshly harvested conidia for 4 h at 4°C (<100%), 25°C (<60 to 70%), and 45°C (<60 to 75%). Strains with deletions were checked for sensitivity to caffeine, ethanol, SDS, Congo red, CaCl₂, and the antifungal drugs caspofungin and itraconazole. Figure 7 shows that pilAΔ mutants, pilAΔ pilBΔ double mutants, and surGΔ mutants but not pilBΔ strains show moderate resistance to 15 μM itraconazole. The itraconazole resistance phenotype cosegregates in crosses with the genetic markers riboB and pyroA, used to interrupt the pilA and surG genes, respectively (see Materials and Methods; data not shown). No other phenotype of resistance or hypersensitivity was observed.

(ii) Cell localization patterns and processes. PilA (but not PilB or SurG) is present in punctate structures in mycelia. The presence of ammonium results in endocytosis in the germlings and mycelia of a number of transporters involved in the utilization of nitrogen sources (5, 35, 49). We have investigated whether the ammonium-elicited endocytosis of the dicarboxylic amino acid transporter AgtA (5) was affected in a strain expressing PilA, PilB, or SurG tagged with GFP proteins was fractionated on a 10% SDS-polyacrylamide gel, transferred to a PVDF membrane, and probed with a primary mouse anti-GFP monoclonal antibody and a secondary goat anti-mouse IgG HRP-linked antibody. Protein markers are indicated in kDa on the left. Equal loading was checked by Coomassie blue staining.

![Figure 4](image-url) FIG. 4. (A) Expression of the pilA, pilB, and surG genes in a wild-type strain, pilA, pilB, and surG transcript levels in ungerminated (0 h), swollen (4 h), and germinated (8 h) conidia and in young (12 h, 16 h) and older (20 h) mycelia are shown. Strains were grown in the presence of 5 mM urea and 1% (wt/vol) glucose as sole nitrogen and carbon sources, respectively, at 25°C. Steady-state 18S rRNA levels were used to monitor the amount of RNA loading in each lane. (B) Western blot analysis of the PilA, PilB, and SurG tagged proteins. Approximately 20 μg of total protein fractions of conidia (0 h) and young mycelia (16 h) derived from strains expressing PilA, PilB, or SurG tagged with GFP proteins was fractionated on a 10% SDS-polyacrylamide gel, transferred to a PVDF membrane, and probed with a primary mouse anti-GFP monoclonal antibody and a secondary goat anti-mouse IgG HRP-linked antibody. Protein markers are indicated in kDa on the left. Equal loading was checked by Coomassie blue staining.

We have checked if any of the strains with a deletion is affected by the distribution of filipin, a polyene antibiotic which selectively stains ergosterol and which can be detected by its blue fluorescence under UV light. No changes in filipin binding were seen in young mycelia of strains with the pilA and surG deletions (data not shown). In these strains, the strong filipin staining at the tip of the hyphae is not altered. However, preliminary results strongly suggest that the distribution of filipin staining is affected in ungerminated conidia by the deletion of either pilA or surG but not by the deletion of pilB (not shown).

(iii) Effects on eisosome assembly. We have checked the distribution of SurG-GFP and PilB-GFP in strains with the pilA deletion and PilA-GFP and PilB-GFP in strains with the surG deletion. Figure 9A shows that in pilAΔ ungerminated conidia (0 h), PilB-GFP patches become larger and more distinctly separated from each other. After 5 h of isotropic growth, PilB is distributed in the cytoplasm as it is in the wild type, with fewer patches persisting at the conidial periphery. In a pilAΔ background, ungerminated conidia (0 h) show a significant decrease of SurG-GFP peripheral patches, while the inner perinuclear ring remains intact, indicating that PilA is required for proper localization of SurG at the conidial periphery. In the absence of SurG (Fig. 9B), the peripheral targeting and distribution of PilB-GFP are drastically affected, with the protein being localized to a few bright clusters at the periphery of ungerminated conidia, which project into the cytoplasm, and to a diffuse cytoplasmic pool, which becomes more evident after 5 h of isotropic growth. At this time point, PilB-GFP clusters not only are peripheral but also appear in the cytoplasm. On the other hand, deletion of SurG does not affect the localization of PilA-mRFP.

**DISCUSSION**

Our own data (see the data in the supplemental material) and those of others (32; A. Olivera-Couto and P. Aguilar,
unpublished data) establish the universal presence of homologues of Pil1 and Lsp1 in the ascomycetes. The sequence divergence between the PilA and PilB clades of the Pezizomycotina, including differences in the conservation of putative phosphorylation sites (data not shown), and the presence of a second coiled-coil domain in PilB are associated with a different fate in the course of development in *A. nidulans*. While Pil1 and Lsp1 are integral components of eisosomes in all stages of the *S. cerevisiae* cell cycle, in *A. nidulans*, PilB shows colocalization with PilA only in conidia. The eisosomes of the *A. nidulans* conidiospore can be considered equivalent to those of *S. cerevisiae*, as they contain both Pil paralogues and SurG, the orthologue of Sur7 (see below). However, there are some intriguing differences between the *S. cerevisiae* cellular eisosomes and those of the conidiospores of *A. nidulans*. In *S. cerevisiae* there is about 1 eisosome/3 μm² of surface area, and they are well separated from each other at a distance of >0.5 μm (28). In the *A. nidulans* conidiospore, eisosomes are more tightly packed and seem to touch each other; thus, they cannot be easily counted (see Fig. S2 in the supplemental material). The peripheral localization of SurG does not match exactly the localization of PilA in conidia but shows a more continuous distribution (Fig. 6). The perinuclear ring of SurG has not been reported for either *S. cerevisiae* or *C. albicans*.

In *S. cerevisiae*, Pil1 but not Lsp1 is essential for proper eisosome assembly (28, 53). In contrast to *S. cerevisiae*, in *A. nidulans* the absence of PilA does not markedly affect the localization of its paralogue, PilB, in conidia. In the absence of PilA, PilB patches are still uniformly distributed at the conidial periphery, but they are larger and more widely spaced than in the wild type. In *S. cerevisiae*, Sur7 and Lsp1 are similarly affected by a pil1 deletion. In *A. nidulans*, the distribution of SurG in a pilA background (see Fig. 9 and Results section) is drastically altered, but the scanty SurG patches are very different from the large peripheral Sur7 clusters seen in pil1 strains. A deletion of SUR7 in *S. cerevisiae* has no effect on the localization of Pil1 and Lsp1. This also applies to strains with additional deletions from the two paralogues, which cluster with Sur7 in the phylogenetic tree (see Fig. S1 in the supplemental material). Similar results were obtained for the deletion of Sur7 in *C. albicans* (3). The situation is quite different in *A. nidulans*. While the distribution of PilA is not affected in

**FIG. 5.** Localization of PilA, PilB, and SurG proteins during conidial development. Representative wide-field fluorescence and DIC pictures of wild-type conidiophores expressing chimeric PilA-GFP (A to F), PilB-GFP (G), and SurG-GFP (H) proteins are shown. The Merge section of panel A shows in inverted black and white an enlarged portion of the same conidial head shown in the PilA-GFP and DIC panels in order to highlight the PilA-GFP spots present in metulae and a budding phialide. Expression of PilB-GFP (G) and SurG-GFP (H) during conidial development is also presented. Young PilA-GFP-expressing conidiophores are shown in panels A and D to F, and mature PilA-GFP-expressing conidiophores are shown in panels B and C. All strains were inoculated on microscope slides covered with MM containing 2% glycerol and 0.8% agarose and grown at 37°C for 2 days. Bars, 5 μm.
surGΔ strains, that of PilB is drastically affected. The distribution of PilB in surGΔ strains is partly reminiscent, in quiescent conidia, of the distribution of Lsp1 in pil1Δ strains. That SurG is necessary for PilB localization is also consistent with the localization of PilB to the cytoplasm of wild-type hyphae, where SurG is sequestered in the vacuole and endosomes and, consequently, may not be available to interact with PilB. Thus, the strong similarity of eisosomal components in S. cerevisiae cells and A. nidulans conidiospores is not correlated with the identical interactions leading to their assembly.

Eisosomes have been defined as portals of endocytosis (52). However, the data presented in this article imply that if they play a role in endocytosis in A. nidulans, this role is minor or limited to specific cargos, which remain unidentified. Neither the endocytosis of FM4-64 nor the ammonium-elicited endocytosis of the AgtA (dicarboxylic amino acid transporter [5]; data not shown) or PrnB (the major proline transporter [41]; data not shown) are visibly affected in mycelia by a deletion of pilA, encoding the only eisosomal protein present in discrete foci in hyphae. It should be noted that although the AgtA and PrnB proteins of A. nidulans and the Hxt2 protein of S. cerevisiae are transporters with a homogeneous distribution in the plasma membrane, upon endocytosis Hxt2 accumulates in discrete plasma membrane-associated foci (52), while the two A. nidulans proteins do not (5).

Endocytosis in A. nidulans has been investigated in mycelia (36), where it is involved in the cycling of transporters (5, 16, 35, 49) and in sensing the external pH (19, 38, 50), but has not been investigated in the process of conidial germination; and the possibility remains open that the yeast-like eisosomes of conidiospores have a role in early, germination-related endocytosis events. The endocytic pathway is nearly essential in A. nidulans. Deletion of components of the endosomal sorting complex required for transport III (ESCRT-III; Vps24 [38] and Vps20, Vps32, and Vsp36 [A. M. Calcagno-Pizzarelli and H. N. Arst, personal communication) results in a drastic impairment of growth. This contrasts with the absence of any obvious growth phenotype in surGΔ, pilAΔ, and pilBΔ strains. A possible explanation for this apparent paradox is that, as proposed for S. cerevisiae, endocytosis could proceed by two independent pathways, one of which would not involve eisosomes (18).

A vexing result, common to S. cerevisiae and A. nidulans, is the absence of any drastic phenotype in deletion mutants of the eisosomal soluble proteins (Pil1/Lsp1/PilA/PilB) or membrane proteins (Sur7/SurG). We did not observe any growth phenotype in any media or at any temperature tested for strains with pilA, pilB, pilA pilB, or surG deletions. Nevertheless it was
reported at a recent meeting that a deletion of ANID_04615.1 (surG) results in growth impairment on minimal medium with nitrate as the nitrogen source at 37°C and 42°C, an impairment that is seen only on complete medium at 42°C (8a; D. Chung and B. Shaw, personal communication). We did not observe this phenotype on minimal medium using a variety of nitrogen sources, including nitrate, or on complete medium at any temperature. The phenotype did not appear after outcrossing our strain with the deletion, which eliminates the possibility that our strain could carry a suppressor of the growth phenotype. We do not know the reason for this difference, which merits investigation. However, a growth phenotype was observed for strains with the surG deletion and the pilA deletion. This is the mild but clear resistance to itraconazole which cosegregates in crosses with each deletion. This is surprising, as deletions of SUR7 in Candida albicans result in marked hypersensitivity to fluconazole (3), another triazole antifungal agent to which Aspergillus species are tolerant (33). This resistance probably operates at the level of conidial survival or germination, and it may be in some way related to the mislocalization of filipin staining in conidia but not in mycelia both in strains with the surG deletion and in strains with the pilA deletion. Triazole drugs act by inhibiting the cytochrome P450-dependent conversion of lanosterol to ergosterol. Filipin is a polyene macrolide antifungal agent that selectively binds and stains ergosterol at the plasma membrane.

One of the surprising results from this work is that in A. nidulans, S. cerevisiae-like eisosomes assemble during conidial formation and disassemble during germination, resulting in germlings and mycelia in eisosome-like punctate structures comprising PilA and a novel cellular distribution (cytosolic and vacuolar, respectively) for PilB and SurG. As no specific conidial survival phenotype was found in strains with eisosomal component deletions, this developmentally regulated distribu-
tion remains baffling. EgID, a putative endoglucanase comprising an expansin-like domain, also localizes at the periphery of ungerminated conidia (7). It would be of interest to study whether this localization implies a topological or functional association with eisosomes. In mycelia, PaI and PaII, involved in pH sensing, have been reported to form membrane-associated punctate structures (8). The connection between PaI and these proteins is under investigation.

The work presented above is a first study of eisosomal structure and function in a model filamentous ascomycete. The universal presence of eisosomal proteins in the ascomycetes and probably in other fungal phyla (3, 32; Olivera-Couto and Aguilar, unpublished; C. Sczazdoczko, unpublished data), coupled with the paucity and diversity of the phenotypes of the deletion mutants observed in three different fungal species (S. cerevisiae, C. albicans, and A. nidulans) and the striking developmental pattern of eisosomal protein distribution seen in A. nidulans, continues to be an unsolved paradox. The role of the Meu14 protein, a relative of the Pil1/PilA proteins in the meiosis of S. pombe, suggests that eisosomal proteins may have acquired entirely new functions in different ascomycete phylogenetic groups. Eisosomes, as defined in S. cerevisiae, are present within the asexual cycle only in the conidia of A. nidulans (see also reference 29), positing the question of their developmental fate of other conserved proteins known to share the ascomycete the developmental fate of other conserved proteomes.

Thus, it would be of interest to explore in this filamentous (see also reference 29), positing the question of their association remains baffling. EglD, a putative endoglucanase, with an expansin like domain is localized in the conidial cell wall of Aspergillus nidulans. Fungal Genet. Biol. 45:839–850.

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