Aspergillus nidulans ArfB Plays a Role in Endocytosis and Polarized Growth

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Filamentous fungi undergo polarized growth throughout most of their life cycles. The Spitzenkörper is an apical organelle composed primarily of vesicles that is unique to filamentous fungi and is likely to act as a vesicle supply center for tip growth. Vesicle assembly and trafficking are therefore important for hyphal growth. ADP ribosylation factors (Arfs), a group of small GTPase proteins, play an important role in nucleating vesicle assembly. Little is known about the role of Arfs in filamentous hyphal growth. We found that Aspergillus nidulans is predicted to encode six Arf family proteins. Analysis of protein sequence alignments suggests that A. nidulans ArfB shares similarity with ARF6 of Homo sapiens and Arf3p of Saccharomyces cerevisiae. An arfB null allele (arfB disrupted by a transposon [arfB::Tn]) was characterized by extended isotropic growth of germinating conidia followed by cell lysis or multiple, random germ tube emergence, consistent with a failure to establish polarity. The mutant germ tubes and hyphae that do form initially meander abnormally off of the axis of polarity and frequently exhibit dichotomous branching at cell apices, consistent with a defect in polarity maintenance. FM4-64 staining of the arfB::Tn strain revealed that another phenotypic characteristic seen for arfB::Tn is a reduction and delay in endocytosis. ArfB is myristoylated at its N terminus. Green fluorescent protein-tagged ArfB (ArfB::GFP) localizes to the plasma membrane and endomembranes and mutation (ArfB<sup>G25A</sup>::GFP) of the N-terminal myristoylation motif disperses the protein to the cytoplasm rather than to the membranes. These results demonstrate that ArfB functions in endocytosis to play important roles in polarity establishment during isotropic growth and polarity maintenance during hyphal extension.

Polarized growth is a biological process observed for many different eukaryotic cell systems, including pollen tube formation (50), neuronal cell development (7), and fungal hyphal growth (5). Filamentous fungi are characterized by extensive polarized growth throughout most of their life cycles. This unique reliance on hyphal morphogenesis makes filamentous fungi an attractive system to study polarized growth. Invasive hyphal growth is an important strategy for animal and plant pathogenic fungi to infect and penetrate their hosts. Therefore, understanding hyphal growth will be critical to the development of therapeutic or antifungal agents.

Girbardt reported the existence of the Spitzenkörper (apical body) at growing hyphae and proposed that the Spitzenkörper coincided with the polarized growth site (18). The Spitzenkörper consists of two populations of vesicles, apical vesicles (79 to 90 nm in diameter) and microvesicles (30 to 40 nm in diameter). Microvesicles possess chitin synthesis activity and have been called “chitosomes” (4, 6). The postulated vesicle supply center thought to be necessary to explain hyphal tip growth (19, 20) may correspond with the Spitzenkörper. This has driven the hypothesis that vesicle assembly and trafficking play key roles in polarized growth in filamentous growth.

ADP ribosylation factors (Arfs) are small GTPase proteins that function in vesicle assembly and trafficking (34, 36). In Saccharomyces cerevisiae, Arf3 is involved in polarity establishment during budding and an arf3 mutant shows a random budding phenotype (16, 22). The Arf3 protein localizes to cell membranes and predominantly to sites of polarized growth (22). Arf3 also genetically interacts with actin cable and actin patch components, including profilin (Pfy1), Arp2/3 complex protein (Las17p and Vrp1), formins (Bni1 and Bnr1), Arf-GTP exchange factor (Gea1), and actin-organizing proteins (Syp1 and Bud6) (30). Another function of Arf3p in yeast is in membrane trafficking during endocytosis (9, 10). Arf6, the human homolog of S. cerevisiae Arf3p (ScArf3p), is also involved in endocytosis, cytokinesis, and actin cytoskeleton organization (reviewed in reference 17).

Shaw et al. reported that a temperature-sensitive N-myristoyltransferase (NMT) mutant displayed a polarity maintenance defect in Aspergillus nidulans (43). Myristate from myristoyl-coenzyme A (CoA) is covalently attached to the secondary glycine of target proteins by NMT increasing in hydrophobicity (24). Myristoylated protein more readily associates with membranes or takes part in hydrophobic protein-protein interactions (24, 41). Many Arfs, including ScArf3p, are N myristoylated (2, 35). We hypothesized that Arf proteins may be the direct connection between N myristoylation and polarized cell morphogenesis in filamentous fungi. Indeed, in S. cerevisiae, N-myristoylated Arfs, including Arf1p, Arf2p,
Arf3p, and Arl1p, have been shown to be associated with polarized budding, in which Arf1p, Arf2p, and Arl1p are involved in a secretion pathway through the Golgi network (40, 46). These secretory vesicles may be destined for the Spitzenkörper in filamentous fungi.

In this study we found six Arf family proteins in *A. nidulans*. Three, namely, AN1126.3, AN5020.3, and AN5912.3 (Arfa, ArfB, and ArfA, respectively), are predicted to be myristoylated (31). Overexpression of ArfA has been shown to suppress the swoF myristoylation mutant (32). Here ArfB is investigated further, since it is the putative ortholog of Arf3p in *S. cerevisiae* associated with polarized bud site selection (22) and actin organization (11, 30). We show that ArfB is important for the establishment and maintenance of hyphal polarity, possibly by directing endocytosis in *A. nidulans*, and that this role requires N myristoylation of the protein.

### MATERIALS AND METHODS

**Strains, growth conditions, plasmids, media, and reagents.** *A. nidulans* strains and plasmids used in this study are listed in Tables 1 and 2, respectively. The plasmids were maintained in *Escherichia coli* XLI-Blue (Stratagene, La Jolla, CA) and manipulated as previously described (42). E. coli DB3.1 was used for the propagation of plasmids containing the *ccdB* box. E. coli C118 was used for the propagation of the transposon donor plasmid containing the *R6K* replication origin. Complete and minimal media for *A. nidulans* were prepared as described previously (27). All chemicals for media, buffer, and supplements were purchased from Sigma (St. Louis, MO) unless otherwise indicated.

**Disruption of arfB by a transposon.** The *arfB* gene was PCR amplified from 500 bp upstream of the start codon and 451 bp downstream of the stop codon by use of the following primers: GarfBF (5′-CACCATGGCCGGCTCGTGGGCG-3′) and GarfBR (5′-GGGGTACCGGCTCGTGGGCG-3′). The amplicon was cloned into the pGEM-T Easy vector (Promega, Madison, WI) to produce pGEM-arfA. A transposon insertion mutagenesis strategy (37) was used to disrupt the *arfB* gene. This transposon contains the *Neurospora crassa* pyr-4*°* gene as a selectable marker. In-frame insertion of the transposon was verified by PCR. The exact insertion site was determined by sequencing with primers N and S from the *GPS-1* kit (New England Biolabs Inc., Ipswich, MA). pArfBTn2, carrying the transposon in the first predicted intron of the *arfB* (after 139 bp from the start of the gene) in the vector pGEM-arfB, was linearized with *SphI* (New England Biolabs Inc., Ipswich, MA), which cuts the plasmid outside of the *arfB* gene. The linearized plasmid was transformed using previously described methods (43) into *A. nidulans* strain TN02A7 (Table 1). This strain is transformed into AXL18, an NMT mutant (43), to observe the localization of *arfB*::GFP in *A. nidulans* and a different strain of *A. nidulans* (44). The linearized plasmid was transformed into BTN1 (Table 1) and separated on a precast 12% sodium dodecyl sulfate-polyacrylamide gel containing 2.2 M of formaldehyde. Standard procedures for Southern and Northern blotting were performed as previously described (31).

**MALDI-TOF MS.** Proteins for matrix-assisted laser desorption ionization–time of flight mass spectrometry (MALDI-TOF MS) analysis were prepared by immunoprecipitation. From a total protein preparation prepared as previously described (31), ArfB::GFP proteins were selectively bound to the monoclonal anti-GFP antibody (Santa Cruz Biotechnology, Inc., Santa Cruz, CA) and eluted directly onto the sample plate by use of 2 μl of α-cyanohydroxy-

### TABLE 1. Strains used in this study

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<th>Strain name</th>
<th>Description</th>
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<td>A773</td>
<td><em>pyrG89 w3 A</em> pyroA4 veA1</td>
<td>FGSC</td>
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<tr>
<td>A4</td>
<td>Wild type</td>
<td>FGSC</td>
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<td>TN02A7</td>
<td><em>pyrG89 pyroA4 riboB2 nkuAargarB</em></td>
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<td><em>arfB::GFP-pyr-4 veA1</em></td>
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<td>53</td>
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<td><em>arfB::Tc::pyr-4 actA::GFP-argB</em></td>
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**TABLE 2. Plasmids used in this study**

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<td>Promega</td>
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<td><em>arfB</em> in pGEM-T Easy vector Entry vector for gateway cloning</td>
<td>This study</td>
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<td>Invitrogen</td>
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<td><em>pyr-4</em> in pGEM-T Easy vector Gateway destination vector with <em>argB</em> gene</td>
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<td>pMT-sGFP</td>
<td>Gateway destination vector with <em>pyr-4</em> gene</td>
<td>51</td>
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<td>pSL-sGFP</td>
<td>Gateway destination vector with <em>pyr-4</em> gene</td>
<td>This study</td>
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</tr>
<tr>
<td>pArfB2A::GFP-pyr4</td>
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<td>This study</td>
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**Green fluorescent protein (GFP) tagging of ArfB and ArfB**2A

The intact *arfB* gene was amplified from wild-type strain A4 genomic DNA with the following primers: GarfBF (5′-CACCATGGCCGGCTCGTGGGCG-3′) and GarfBR (5′-GGGGTACCGGCTCGTGGGCG-3′). To generate nonmyristoylated ArfB, a site-directed mutation was introduced by using primer mGarfBF (5′-GACATGGCCGGCTCGAGGACAT-3′) instead of GarfBF, which changed the secondary glycine into alanine (G2A; the site of this change is underlined). The secondary glycine is critical for myristoylation (33). The *arfB* and *arfB*2A amplimers were cloned into the gateway entry vector pENTR (Invitrogen, Carlsbad, CA) to produce pArfB-entry and pArfB2A-entry, respectively.

To construct a GFP expression vector with the pyr-4 marker gene, pMt-sGFP (51) was partially digested with NotI (New England Biolabs Inc., Ipswich, MA) and the vector fragment without the 1.8-kb fragment containing the *arfB* gene was ligated with a 1.6-kb fragment containing the pyr-4 gene from pGEM-pyr4 to produce pSL-sGFP (Table 2).

The *ccdB* box in *pSL-sGFP* was exchanged with *arfB* or *arfB*2A in the entry vectors mentioned above through the use of a LR Clonase reaction following the manufacturer’s instructions (Invitrogen, Carlsbad, CA) to generate pArfB::GFP-pyr4 or pArfB2A::GFP-pyr4, respectively. Each vector was transformed into A773 (Table 1) or TN02A7. pArfB::GFP-pyr4 was also transformed into AXLI8, an NMT mutant (43), to observe the localization of *arfB*::GFP in the absence of N myristoylation.

**Southern, Northern, and Western blotting.** The intact *arfB* gene amplified with primers GarfBF and GarfBR was used as the probe for Southern and Northern blotting. The probe was labeled by using Prime-It II, a random-primer labeling kit (Stratagene, La Jolla, CA), by using [γ-32P]ATP. For Southern blotting, the wild-type strain (TN02A7) and BTN1 was extracted after 24 h of incubation in liquid culture by using Trizol reagent (Invitrogen, Carlsbad, CA) as indicated by the instruction of the manufacturers. The RNA was separated on a 1% agarose gel containing 2.2 M of formaldehyde. Standard procedures for Southern and Northern blotting were performed as previously described (42).

**Monoclonal anti-GFP antibody.** (Santa Cruz Biotechnology, Inc., Santa Cruz, CA) was used for Western blotting for GFP-tagged proteins. Total proteins were extracted from WB17 and MB13 (arfB::GFP and arfB2A::GFP, respectively, Table 1) and separated on a precast 12% sodium dodecyl sulfate-polyacrylamide gel electrophoresis gel (Bio-Rad, Hercules, CA). Protein extraction and Western blotting were performed as previously described (31).

**MALDI-TOF MS.** Proteins for matrix-assisted laser desorption ionization–time of flight mass spectrometry (MALDI-TOF MS) analysis were prepared by immunoprecipitation. From a total protein preparation prepared as previously described (31), ArfB::GFP proteins were selectively bound to the monoclonal anti-GFP antibody (see above) and the antibody-protein complexes were precipitated with protein A in the ImmunoPure immobilized protein A system (Pierce Biotechnology, Inc., Rockford, IL) following the company’s instructions. Sodium dodecyl sulfate gel electrophoresis (29) of the precipitated proteins was performed and the gels were stained with Bio-Safe Coomassie blue (Bio-Rad, Hercules, CA). A band corresponding to the *ArfB::GFP* protein was excised from the gel and subjected to reduction, alkylation with iodoacetamide, trypsin digestion (Promega, Madison, WI), and extraction according to established methods (45). The extracted peptides were purified using C18 ZipTip (Millipore) and eluted directly onto the sample plate by use of 2 μl of α-cyanohydroxy-

Downloaded from http://arc.asm.org/ on May 1, 2016 by guest
cinamic acid solution (6 mg/ml in 50% [vol/vol] acetonitrile containing 0.1%
trifluoroacetic acid). Mass spectra were collected in reflectron positive-ion mode
on an Axima CFR MALDI-TOF MS (Shimadzu/Kratos, Manchester, United
Kingdom). The peptide masses were calibrated internally with trypsin fragments.
Average mass differences between predicted and measured masses were between
20 and 300 ppm.

Microscopy. GFP protein and FM4-64-stained live cells were visualized in
germlings grown in custom-made aluminum chambers designed for live-cell
imaging of fungal cells (28). For time-lapse imaging, an Olympus BX51 micro-
scope (Olympus America Inc., Melville, NY) was outfitted with a Prior shutter to
limit phototoxicity to the cells. Simple PCI software (version 5.3.1.081004;
Compix Inc., Imaging Systems, Cranberry Township, PA) was used to acquire
images of growth at 30-second intervals. To observe the endocytosis process in
cells, wild-type and BTN1 cells were grown on a coverslip submerged in liquid
complete medium. After 24 h of incubation, fresh medium containing 12.5
M
H9262
of FM4-64 was replaced with the growing medium lacking the dye. The coverslip
with cells was overlaid on the aluminum chamber. A narrow-band G-excitation
filter with 530 to 550 nm of excitation and 590 nm of emission (Olympus America
Inc., Melville, NY) was used to obtain FM4-64 fluorescent images. Microscopic
observation for GFP and all image processing were conducted as previously
described (44, 52).

RESULTS

ArfB is an ARF6 family protein in A. nidulans. A. nidulans
and S. cerevisiae encode six Arf family proteins (ArfA
[AN1126.3], ArfB [AN5020.3], ArlA [AN5912.3], AN3934.3,
AN0411.3, and AN0634.3 for A. nidulans) and Arf1p, Arf2p,
Arf3p, Arl1p, Sar1p, and Arl3p for S. cerevisiae (see Fig. S1 in
the supplemental material). We found that ArfB (AN5020.3) is
a putative ortholog of ScArf3p, which was previously shown to
be involved in the selection of bud sites (22) through interactions
with proteins involved in actin assembly (30).

A. nidulans ArfB (AnArfB) and ScArf3p are further grouped into the
ARF6 family of proteins (or class III Arfs), named after human
ARF6 (17). The AnArfB, ScArf3p, and Homo sapiens
ARF6 (HsARF6) protein sequences were aligned (Fig. 1A). AnArfB
shares 59% amino acid sequence identity with ScArf3p and
72% identity with HsARF6. ScArf3p shares 59% similarity
with HsARF6. The AnArfB, ScArf3p, and HsARF6 proteins
have a conserved N-terminal myristoylation site and a con-
served GTP binding and hydrolysis site for their GTPase ac-
tivity (Fig. 1A). The
arfB
gene is located on chromosome III
from bp 507,503 to bp 508,272 on contig 84 (of release 3 of
the
A. nidulans
genome database [http://www.broad.mit.edu/annotation/genome/aspergillus_group/MultiHome.html]). The
arfB
gene is 770 bp in length and is predicted to contain four
introns at the following positions: 94 to 153 bp, 312 to 363 bp,
511 to 561 bp, and 674 to 725 bp. ARF6 family proteins were
retrieved from filamentous fungal genome databases including
protein data for A. nidulans, N. crassa, Magnaporthe grisea,
Fusarium graminearum, Candida albicans, Cryptococcus neofo-
mans, and Ustilago maydis in addition to S. cerevisiae. All of
these proteins contain the consensus Met-Gly-x-x-Ser/Thr motif
(established in reference 24) at their N termini (Fig. 1B), suggesting that N myristoylation of ARF6 family proteins is important for their function.

**Disruption of arfB by a transposon insertion.** To investigate the role of arfB in *A. nidulans*, we disrupted the gene. The arfB disruption cassette was constructed in the pGEM-T Easy vector (see Materials and Methods). The transposon contains the N. crassa *pyr-4* gene as a nutritional selection marker. Sequencing verified the transposon insertion site in the predicted first intron of the arfB gene (after 139 bp from the start of the gene) (Fig. 2A), resulting in a truncation upstream of the GTP binding and hydrolysis sites. The intact arfB gene was replaced by transformation in the *nakA* deletion strain (TN02A7) (Fig. 2B). A single insertion was verified by PCR (data not shown). Southern blot analysis showed that the arfB disruptant (BTN1) has an 8.8-kb band that is upshifted relative to the approximately 4.8 kb of the control strain. This indicates that the BTN1 strain contains a single copy of the arfB::Tn cassette replaced the native arfB locus in the BTN1 strain. (D) Northern blot analysis shows that the transposon insertion in the predicted first intron disrupted the expression of the arfB gene. (E) BTN1 colonies develop markedly slower than do those of the wild type.

Since the transposon is in a predicted intron, we examined the expression of the arfB gene in the control strain and the arfB::Tn strain. A Northern blot (Fig. 2D) shows that the control strain expressed intact mRNA of arfB; from arfB::Tn, however, the arfB mRNAs were below the detection limit (Fig. 2D), indicating that the transposon insertion disrupts the expression of the arfB. We therefore conclude that arfB::Tn is a null allele.

Compared to the control strain, the arfB::Tn strain produced slow-growing colonies that are aconidial (Fig. 2E). The lack of conidiation was partially suppressed in the presence of an osmotic stabilizer (1.2 M sorbitol), though colonial growth remained limited (data not shown). This characteristic allowed the propagation of the arfB::Tn disrupted strain by conidia.

The arfB::Tn strain is defective in polarity establishment during spore germination and polarity maintenance during hyphal growth. The arfB::Tn strain displayed extended isotropic growth, multiple germ tube emergence, and dichotomous branching at hyphal apices (Fig. 3). Under identical conditions, the wild-type strain sent a primary germ tube following isotropic growth, multiple germ tube emergence, and dichotomous branching at hyphal apices (Fig. 3). Under identical conditions, the wild-type strain sent a primary germ tube following isotropic growth, multiple germ tube emergence, and dichotomous branching at hyphal apices (Fig. 3).

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FIG. 3. Phenotype of the arfB::Tn strain. (A) Microscopic phenotypes of the wild-type (WT) and arfB::Tn strains. A site where multiple germ tubes emerge from the same conidium is indicated by a white arrowhead. (B) Time-lapse images of wild-type and BTN1 cells. The white arrows indicate septation sites, the black arrowhead indicates a septum, and the black arrows indicate simultaneous septation sites for the arfB::Tn strain. (C) Polarized hyphal growth of the arfB::Tn strain. Symbols: *, abnormal hyphal growth of the arfB::Tn strain; **, wild-type-like hyphal growth of the arfB::Tn strain. (D) Combined Z-stack images from a spinning-disc confocal microscope of the wild-type and arfB::Tn strains. Multiple nuclei in the isotropic, apolar conidium are observed for the arfB::Tn strain. Germlings of the wild-type and arfB::Tn strains were stained with calcofluor to visualize cell walls and with Hoechst 33258 to visualize nuclei. Numerical aperture (NA), 1.4 with oil. Bars = 20 μm.
Supplemental material). (i) Large isotropically swollen conidia lyse (35.3% ± 3.7%; n = 607). (ii) Large isotropically swollen conidia send a germ tube, followed by the loss of polarity of the germ tube (17.2% ± 4.4%). These apolar germ tubes can later recover polarity to produce additional germ tubes. (iii) Multiple germ tubes emerge simultaneously from the swollen conidium (47.5% ± 3.7%). Hyphae often dichotomously split at their apices. Multiple simultaneous septation events were observed in dichotomously branching hyphae (Fig. 3B). Initial hyphal growth is consistently irregular. Hyphae meandered and produce irregularly lobed cells (Fig. 3C). Interestingly, hyphae later (after 60 to 80 μm of initial hyphal growth) appear almost like wild-type hyphae, suggesting that another gene may function to compensate for the absence of arfB or that the gene functions primarily during the initial stages of hyphal growth. Germlings of the wild-type and arfB::Tn strains were stained with calcofluor to visualize cell walls and with Hoechst 33258 to visualize nuclei (Fig. 3D). Multiple nuclei in cell compartments of the arfB::Tn strain indicated no cell cycle-associated phenotypes.

The Spitzenkörper of the wild-type strain (A4) and that of the arfB::Tn strain were examined. The Spitzenkörper is a filamentous fungus-specific “organelle” composed of a cloud of secretory vesicles and involved in hyphal growth (20). We visualized the Spitzenkörper by staining with FM4-64 (13). The wild-type strain has an intact Spitzenkörper at the hyphal apex (Fig. 4A); however, the arfB::Tn strain lacks an intact FM4-64-stained Spitzenkörper (Fig. 4B). It is possible that trafficking of the dye in the arfB mutant caused insufficient staining of the Spitzenkörper or that intact ArfB is required to assemble a wild-type Spitzenkörper.

ArfB is involved in endocytosis. To verify the function of ArfB in endocytosis in A. nidulans, we stained live cells of the wild-type and arfB::Tn strains with FM4-64 and tracked the uptake of the dye over time. In the wild-type strain, similar to previous observations (38), the dye appeared on the plasma membrane within 10 min after treatment. After 45 min, the dye was taken up by cells and localized to endomembranes, which may be mature endosomes or vacuoles (Fig. 5A and B). In the arfB::Tn strain, dye uptake was delayed. After 30 min of treatment, the dye remained on the plasma membrane. Even after 60 min or more of treatment, the labeling of endomembranes was diffuse in the arfB::Tn strain in comparison with what was seen for the wild type (Fig. 5B).

Actin patches are thought to be sites of endocytosis (26). Therefore, we also observed actin patches in the wild-type and arfB::Tn strains. Our lab has previously constructed ActA::GFP (actin) (53) in which actin localized to cortical patches. To obtain an arfB::Tn strain expressing ActA::GFP, we crossed ASUAGA4 (actA::GFP) to BTN1 (arfB::Tn). Progeny displaying the arfB::Tn phenotype were examined for the expression of ActA::GFP. In wild-type cells, actin patches were concentrated at growing hyphal apices (Fig. 6A). In the arfB::Tn cells, however, the actin patches were dispersed randomly throughout apolar cells, while in polarized cells the actin patches were enriched at hyphal apices (Fig. 6B), indicating...
that the localization of actin patches at the hyphal apex was delayed in the mutant.

**ArfB::GFP localization.** To observe the localization of ArfB, expression constructs driven by the alcohol-inducible *alcA* promoter (14) were assembled for ArfB::GFP and ArfB*G2A*::GFP. These constructs were ectopically inserted in the wild-type A773 strain. Western blotting with monoclonal anti-GFP antibody showed a single band from total proteins of WB17 and MB13 that were expressed at comparable levels (see Fig. S2 in the supplemental material), indicating that the GFP localization reported below was not caused by free GFP. ArfB::GFP localized to the plasma membrane, to septa, and to endomembranes, which may be endoplasmic reticulum, vacuoles, or endosomes (Fig. 6C and E). ArfB::GFP could be endocytosed to internal membranes through membrane recycling. It is not surprising that membrane proteins are internalized through endocytosis (15, 21). It is also possible that the endomembrane localization could be caused by the *alcA*-driven overexpression of ArfB::GFP protein. ArfB::GFP was not enriched in the plasma membranes at the hyphal tip (Fig. 6D). Plasmolysis of labeled cells indicated that ArfB::GFP is not on the septum wall but rather on the plasma membrane at the septum (Fig. 6F). Because septa have plasma membrane on both sides of the wall, they were brighter than membrane along other regions of the hyphae. ArfB::mRFP (monomeric red fluorescent protein) showed a similar localization pattern (data not shown). Four different strains with genotypes of *alc(p):arfB:GFP arfB::Tn* or *alc(p):arfB mRFP arfB::Tn* showed wild-type hyphal growth in the presence of ArfB::GFP or ArfB::mRFP expression under *alcA*-inducing conditions, demonstrating that both ArfB::GFP and ArfB::mRFP are functional (see Fig. S3 in the supplemental material).

To elucidate the role of N myristoylation in the localization of ArfB, we constructed a nonmyristoylated form of ArfB by mutating the secondary glycine to alanine (G2A). ArfB*G2A*::GFP localized to the cytoplasm (Fig. 6C). We also expressed ArfB::GFP in the *swoFI* NMT mutant (43), where its localization was cytoplasmic (Fig. 6C). Multiple independent transformants with the *arfB::sGFP* (12 transformants) and *arfB*G2A*::sGFP* (5 transformants) constructs displayed the same localization pattern (data not shown). Thus, the elimination of the myristoylation motif completely disrupts the localization of ArfB. This result demonstrates that N myristoylation of ArfB is required for the proper localization of the protein.

**ArfB::GFP is N myristoylated.** We overexpressed ArfB::GFP from wild-type strains. After total protein extraction, the GFP-tagged protein was selectively purified by immunoprecipitation using monoclonal anti-GFP antibody (Santa Cruz Biotechnology, Inc., Santa Cruz, CA). The immunoprecipitated ArfB::GFP was approximately 47 kDa (Fig. 6A). The band was differentiated from the heavy chain (50 kDa) of the GFP antibody (data not shown). The band was excised from polyacrylamide gel; subjected to reduction, alkylation, and trypsin digestion; and analyzed by MALDI-TOF MS.

N-terminal myristoylation occurs at the secondary glycine after cleavage of the leading methionine (25). Therefore, various isoforms of ArfB::GFP could be generated. Theoretical monoisotopic masses for ArfB::GFP are summarized in Table 3. Each of the identified peaks showed the expected isotope patterns characteristic of peptides (data not shown). In the case of the wild-type ArfB::GFP, a peak was observed at the *m/z* value of 744.6550 (Fig. 7). This value is in good agreement with the theoretical value of the amino-terminal peptide of ArfB with the addition of myristoyl-CoA (744.6682) with the methionine absent. A leading methionine is cleaved and secondary glycine is myristoylated, as demonstrated previously (24). This result demonstrates that ArfB is myristoylated.

**DISCUSSION**

One of the long-term goals of our lab is to better understand how N myristoylation plays a role in polarized cell morphogenesis in *A. nidulans*. Indeed, the temperature-sensitive
**swoF1** mutant, containing a mutation in the NMT, displayed abnormal polarized growth at the restrictive temperature (43). When grown in submerged culture at restrictive temperature, the **swoF1** mutant first grows isotropically and then sends a germ tube that is similar to the wild-type germ tube. Immediately after this germ tube emerges, hyphal growth ceases and the tip swells isotropically. Previously we found that NMT regulates 26S proteasome activity during early polarized morphogenesis and established the myristoylome of *A. nidulans* with 41 predicted myristoylated proteins (31). We have also demonstrated that overexpression of ArfA partially suppresses the polarity phenotype of the **swoF** myristoylation mutant (32).

The exact role of NMT in polarized growth in systems including *S. cerevisiae*, filamentous fungi, and mammalian cells remains to be fully elucidated.

In filamentous fungi, the Spitzenkörper directs hyphal polarized growth (18). The Spitzenkörper consists of a cloud of vesicles and has been considered a vesicle supply center (19, 20). Since Arfs are involved in coated vesicle formation and trafficking and many Arfs are N myristoylated (2, 34–36), they could link N myristoylation to polarized growth in the filamentous fungi.

A comparative genome analysis of *A. nidulans* and *S. cerevisiae* revealed that both organisms encode six Arf family proteins (see Fig. S1 in the supplemental material). In a previous study using yeast, Arf3p has been shown to be involved in

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**TABLE 3. Predicted mass value of N-terminal tryptically digested fragment of ArfB::GFP**

<table>
<thead>
<tr>
<th>Mass value</th>
<th>Position</th>
<th>Modification(s)</th>
<th>Modified mass value</th>
<th>Peptide sequence</th>
<th>Protein</th>
</tr>
</thead>
<tbody>
<tr>
<td>665.33a</td>
<td>1–7</td>
<td>MSO</td>
<td>681.32</td>
<td>MGGSVSK</td>
<td>ArfB</td>
</tr>
<tr>
<td>1–7</td>
<td>MYRd</td>
<td>875.71</td>
<td>GGSVSK</td>
<td>ArfB</td>
<td></td>
</tr>
<tr>
<td>1–7</td>
<td>MSO + MYR</td>
<td>891.70</td>
<td>GGSVSK</td>
<td>ArfB</td>
<td></td>
</tr>
<tr>
<td>534.29b</td>
<td>2–7</td>
<td>MYR</td>
<td>744.67</td>
<td>GGSVSK</td>
<td>ArfB</td>
</tr>
</tbody>
</table>

*a* Theoretical mass value of unmodified N-terminal fragment of ArfB::GFP.

*b* Theoretical mass value of unmodified N-terminal fragment of ArfB::GFP after cleavage of methionine.

*c* MSO, addition of methionine sulfoxide (artificial modification).

*d* MYR, N myristoylation.
developing polarity during bud formation (22) and in actin cytoskeleton organization (30). ArfB (AN5020.3) of \textit{A. nidulans} shares close similarity with Arf3p (Fig. 1A) and both proteins have a conserved N-myristoylation motif. ArfB and Arf3p are further classified into the ARF6 protein family (named for human ARF6), and many filamentous fungal ARF6 family proteins contain a conserved consensus N-myristoylation motif (Fig. 1B). ArfB, therefore, may be one of the connections between N-myristoylation and polarized cell morphogenesis.

Disruption of the \textit{arfB} gene resulted in defects in polarized morphogenesis (Fig. 3). The ArfB disruptant displayed extended isotropic growth upon spore germination, leading to apolar cells of up to 20 \mu m in diameter. In contrast, wild-type conidia grow isotropically to no more than 5 \mu m of diameter before sending out a germ tube (Fig. 3B). The loss of polarity without cell cycle arrest (Fig. 3D) is similar to what is seen for other polarity-defective mutants examined in our lab (43, 44, 52). After the isotropic growth stage, approximately 48% of the \textit{arfB} mutant germlings sent germ tubes (Fig. 3A and B). Previously, it was observed that delivery of active Cdc42 to an artificial polar site triggers a spontaneous polarization without polarity determinants in \textit{S. cerevisiae} (55). Our data also show that \textit{A. nidulans} is likely to polarize spontaneously after the abnormal swellings without the function of ArfB through an as-yet-undetermined mechanism.

Arf3 participates in actin organization in \textit{S. cerevisiae} (30). We have shown that in the \textit{arfB} mutant, the actin cytoskeleton is not organized properly in apolar cells but that the apical concentration of actin patches is restored in polarized cells. Therefore, random actin distribution at the initial stage of polarity establishment could cause the random polarization resulting in multiple germ tube emergence sites.

The \textit{arfB} mutant also displayed abnormal hyphal morphologies in the early stages of hyphal growth (hyphal lengths of less than 60 \mu m), in which the mutant often meandered off the long axis of polarity and branched dichotomously at hyphal apices, which may result from the loss of the ability to maintain polarity during hyphal growth (Fig. 3C). Hyphal growth to lengths of greater than 60 to 80 \mu m from conidia displayed almost wild-type-like morphology, suggesting that the cell may mobilize another mechanism to compensate for the loss of ArfB, perhaps by upregulating one of the other five Arfs encoded by \textit{A. nidulans}.

It is evident that disruption of the \textit{arfB} gene resulted in deficiencies in bulk membrane internalization (Fig. 5) by endocytosis. The \textit{arfB} mutant displayed a significant delay of FM4-64 uptake, and the efficiency of the dye uptake was lower than that seen for the wild type (Fig. 5). In addition, actin patches, which are considered to be clathrin-mediated endocytosis sites (26), were disrupted in apolar \textit{arfB} mutant cells (Fig. 6A), suggesting that ArfB may be involved in clathrin-mediated endocytosis in \textit{A. nidulans}. The clathrins have been shown to be recruited by Arf (26). Therefore, ArfB may play a role in recruiting the clathrins to the endocytosis site. An additional phenotype of the \textit{arfB} mutant includes perturbation of endomembranes, which are less readily stained and have a diffuse appearance when perturbed (Fig. 5B). This phenomenon is similar to the perturbation of endomembranes in the human ARF6 mutant (39).

The polarization of exocytosis is generally accepted to be a key mechanism in polarized hyphal growth (3, 54). A role for endocytosis in the polarization of hyphae is, however, a relatively new concept. In pioneering work in the development of the endocytosis marker dye FM4-64 for filamentous fungi, Fischer-Parton and colleagues showed that endocytosis is an
ongoing process during hyphal growth (13). Steinberg has also addressed the role for endocytosis in the pathogenicity of Ustilago maydis (47). In three very recent studies, endocytosis has been incorporated into a model explaining hyphal growth (1, 48, 53). The endocytic markers FimA (53), AbpA, and SlaB (1) localize to a subapical collar in growing hyphae. Disruption of fimA resulted in a loss of endocytosis accompanied by severe cell polarity defects that produce phenotypes very similar to those for arfB:Tn as reported here (53). An apical tip growth apparatus model was recently proposed to combine both exocytic and endocytic mechanisms into a unified model for tip growth (48). An endocytic recycling model has also been proposed to explain the role of endocytosis in maintaining polarized membrane components at growing hyphal apices while recycling these components from the subapical region (53). This model accounts for the many varied hypotheses as to what the identity of polarized marker(s) is, including structural proteins, signaling proteins, ion channels, and lipid raft components. Takeshita and colleagues (49) have recently demonstrated the presence of TeaA and TeaR as cell end markers in A. nidulans. Could these proteins be maintained at the cell apex through endocytic recycling?

ArfB localized to plasma membranes and also appeared to localize to endomembranes (Fig. 6). This localization pattern needs to be interpreted with some caution, since the construct described here is both overexpressed via the alcA promoter and ectopically inserted. This localization pattern was, however, similar to that observed for human ARF6 (12, 39). In contrast, Arf3p in yeast is concentrated to the polarized budding site, though it localizes to peripheral membranes (22). The localization of ArfB to membranes depends on the myristoylation of the proteins. We have shown that mutation of the N-myristoylation motif changes the ArfB localization pattern to diffusion throughout the cytoplasm (Fig. 6). Similarly ArfB-GFP was diffuse in the swoF myristoylation mutant. Therefore, the loss of myristoylation of ArfB may explain in part the polarity defect previously reported for the NMT mutant (43).

ArfB localized to the plasma membrane and to endomembranes, but we did not observe it specifically concentrated to the hyphal apices. How then does ArfB take part in polarity establishment and maintenance? One model is that a protein which interacts with ArfB, such as its GDP/GTP exchange factor, concentrates to growing hyphal apices or its GTPase activation protein concentrates at sites away from the growing apex. A similar pattern has been observed for yeast where Bud1 (Rsr1), a ras-like GTPase, is required for bud site selection (8). The localization of Bud1p is cytosolic (23), but its GTPase activation protein, Bud2, localizes specifically to the bud sites (23).

In this paper, we have shown that ArfB plays a critical role in polarity establishment and maintenance. Hyphae tens of microns distal to the germ tube or branch initiation site grew as wild-type hyphal grow, suggesting that another gene may function to compensate for the absence of arfB or that the gene may function primarily during the initial stages of hyphal growth. At least one function of ArfB is in endocytosis, but whether its endocytosis function is through nucleating vesicle assembly or through participating in the organization of the actin cytoskeleton remains to be fully elucidated. N myristoylation of ArfB is critical for its proper localization. ArfB is only 1 of 41 proteins in the predicted myristoylome of A. nidulans (31). Therefore, further investigation will continue to elucidate the role of myristoylation in regulating cell development in A. nidulans.

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