Contribution of Peroxisomes to Secondary Metabolism and Pathogenicity in the Fungal Plant Pathogen *Alternaria alternata* 

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The filamentous fungus *Alternaria alternata* includes seven pathogenic variants (pathotypes) which produce different host-selective toxins and cause diseases on different plants. The Japanese pear pathotype produces the host-selective AK-toxin, an epoxy-decatrienoic acid ester, and causes black spot of Japanese pear. Previously, we identified four genes, *AKT1*, *AKT2*, *AKT3*, and *AKTR*, involved in AK toxin biosynthesis. *AKT1*, *AKT2*, and *AKT3* encode enzyme proteins with peroxisomal targeting signal type 1 (PTS1)-like tripeptides, SKI, SKL, and PKL, respectively, at the C-terminal ends. In this study, we verified the peroxisome localization of Akt1, Akt2, and Akt3 by using strains expressing N-terminal green fluorescent protein (GFP)-tagged versions of the proteins. To assess the role of peroxisome function in AK-toxin production, we isolated Aa*PEX6*, which encodes a peroxin protein essential for peroxisome biogenesis, from the Japanese pear pathotype and made Aa*PEX6* disruption-containing transformants from a GFP-Akt1-expressing strain. The ΔAa*PEX6* mutant strains did not grow on fatty acid media because of a defect in fatty acid β-oxidation. The import of GFP-Akt1 into peroxisomes was impaired in the ΔAa*PEX6* mutant strains. These strains completely lost AK toxin production and pathogenicity on susceptible pear leaves. These data show that peroxisomes are essential for AK-toxin biosynthesis. The ΔAa*PEX6* mutant strains showed a marked reduction in the ability to cause lesions on leaves of a resistant pear cultivar with defense responses compromised by heat shock. This result suggests that peroxisome function is also required for plant invasion and tissue colonization in *A. alternata*. We also observed that mutation of Aa*PEX6* caused a marked reduction of conidiation.

Peroxisomes are single-membrane-bound organelles and have a wide range of metabolic functions, including β-oxidation of fatty acids, peroxide detoxification, and glyoxylate metabolism (60, 62). Peroxisome biogenesis has been extensively studied using yeast mutants, and a number of *PEX* genes that encode peroxins have been identified (48, 60, 62). Peroxins are proteins required for peroxisome biogenesis and division and for import of proteins into the peroxisome matrix (48, 60). Most peroxisome matrix proteins contain peroxisomal targeting signal type 1 (PTS1), the tripeptide sequence (S/A/C)-(K/R/H)-L, at the C terminus (9). Other matrix proteins have PTS2 sequences close to the N terminus; the consensus PTS2 sequence is (R/K)-(L/V/I)-X5-(H/Q)-(L/A) (29).

Peroxisomes are required for specific functions in filamentous fungi. In *Penicillium chrysogenum*, peroxisomes participate in penicillin biosynthesis: acyl coenzyme A (acyl-CoA); isopenicillin N acyltransferase, which catalyzes the final step of penicillin biosynthesis, localizes to peroxisomes (38, 39). In the cucumber anthracnose pathogen *Colletotrichum orbiculare* (synonym, *C. lagenarium*) and the rice blast pathogen *Magnaporthe oryzae* (synonym, *M. grisea*), peroxisome function is necessary for plant infection (6, 8, 25, 49, 65, 66). These pathogens produce specific infection structures, called appressoria, that are used to penetrate the host plant cuticle using mechanical force (61). Their dome-shaped appressoria are darkly pigmented with dihydroxynaphthalene melanin, which is essential for the function of the appressoria. In these pathogens, peroxisome function is involved in appressorium maturation with accumulation of melanin and initial host invasion via appressoria (6, 8, 25, 49, 65, 66).

The imperfect fungus *Alternaria alternata* is one of the most cosmopolitan fungal species and is generally saproxylic (28, 50, 58). This species, however, does include seven pathogenic variants (pathotypes) which produce different host-selective toxins and cause necrotic diseases on different plants (28, 58). Host-selective toxins produced by fungal plant pathogens are generally low-molecular-weight secondary metabolites and are critical determinants of host-specific pathogenicity or virulence in several plant-pathogen interactions (17, 28, 58, 68).

The Japanese pear pathotype of *A. alternata* produces AK-toxin (Fig. 1) and causes black spot on a narrow range of susceptible Japanese pear cultivars, including the commercially important cultivar Nijisseiki (41, 45). We previously isolated the gene cluster involved in AK-toxin biosynthesis from the Japanese pear pathotype and identified four genes, *AKT1*, *AKT2*, *AKT3*, and *AKTR* (55, 56). *AKTR* encodes a transcription regulator of the Zn(II)2Cys6 family (56). *AKT1*, *AKT2*, and *AKT3* are predicted to encode proteins with similarity to the carbonyl-activating enzymes, the estelase-lipase family enzymes, and the hydratase-isomerase family enzymes, respec-

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the strawberry and tangerine pathotypes of A. alternata. AK-toxins of the Japanese pear pathotype (41), AF-toxins of the strawberry pathotype (43), and ACT-toxins of the tangerine pathotype (27) have a moiety, EDA, in common.

Orthologs of AKT1, AKT2, AKT3, and AKT4 are present in the strawberry and tangerine pathotypes of A. alternata (15, 33, 35, 55, 56). The strawberry pathotype produces AF-toxin (Fig. 1) and causes Alternaria black spot of strawberry (31, 43). The tangerine pathotype produces ACT-toxin (Fig. 1) and causes black spot of citrus and mandarins, a disease that has not yet occurred in Japan (27). The toxins of Japanese pear, strawberry, and tangerine pathotypes have a 9,10-epoxy-8-hydroxy-9-methyl-decatrienoic acid (EDA) structural moiety in common (Fig. 1) (27, 41, 43). The AKT orthologs were isolated from the strawberry and tangerine pathotypes and named the AFT and ACTT genes, respectively (15, 33, 35). AKT, AFT, ACTT, and their respective orthologs show more than 90% nucleotide identity, suggesting that these genes encode the enzymes for the biosynthesis of the common precursor, EDA. The PTS1-like tripeptides are also conserved in the orthologs from the strawberry and tangerine pathotypes and named the same PTS1-like tripeptide, SKI, SKL, and PKL, respectively (55, 56). A PSORT II (40) analysis of the amino acid sequences of Akt1, Akt2, and Akt3 identified PTS1-like tripeptides in their C-terminal ends, suggesting that these enzymes are located in peroxisomes (55, 56).

Materials and Methods

Fungal strains and genomic library. Strain 15A of the Japanese pear pathotype of A. alternata and its transformants were used in this study and routinely maintained on potato dextrose agar (PDA; Difco, Detroit, MI). A genomic cosmid library of strain 15A has been previously described (26).

Plasmids. The plasmids used in this study are listed in Table 1. The integrative transformation vectors pSH75 (26), pII99 (44), and pSB116 were used for the transformation of A. alternata. The vectors pSH75 and pII99 carry hph and nptII, respectively, fused to the Aspergillus nidulans trpC promoter and terminator (26, 37, 44). The vector pSB116 was made by replacing the trpc promoter and hph region in pSH75 with a 1.0-kb fragment containing the trpC promoter and the bar open reading frame (ORF) from pBI44MBr (57).

The GFP expression vector pYTGFP-N was used to make a GFP N-terminal fusion vector (see Fig. S1 in the supplemental material). The vector pYTGFP-N contains the GFP (enhanced GFP) ORF fused to the A. nidulans trpC promoter and terminator (19). Because the trpC promoter is constitutively active in A. alternata, we expected it to be useful for expressing the target genes under any culture conditions and in any fungal structures. The AKT1, AKT2, and AKT3-1 cDNAs were amplified from the total RNA of strain 15A with primer pairs AKT1-AKT1-r, AKT2-AKT2-r, and AKT3-AKT3-r (see Table S1 in the supplemental material), respectively, using the RNA PCR kit ver. 2.1 (Takara Bio, Ohtsu, Japan), digested with restriction enzymes, and ligated into pYTGFP-N to make pGFP-AKT1, pGFP-AKT2, and pGFP-AKT3, respectively (see Fig. S1 in the supplemental material). Strain 15A was found to have at least two copies of AKT3 (AKT3-1 and AKT3-2) in its genome (56). The two genes were predicted to encode proteins with 94% amino acid sequence identity and with the same PTS1-like tripeptide, PKL, at their respective C-terminal ends (56). In this study, we used AKT3-1 to make pGFP-AKT3. The AKT1, AKT2, and AKT3-1 cDNAs lacking the nine nucleotides that encode PTS1 were amplified from pGFP-AKT1, pGFP-AKT2, and pGFP-AKT3, respectively, with primer pairs AKT1-AKT1-r2, AKT2-AKT2-r2, and AKT3-AKT3-r2 (see Table S1 in the supplemental material), digested with restriction enzymes, and ligated into pYTGFP-N to make pGFP-AKT1Δ, pGFP-AKT2Δ, and pGFP-AKT3Δ, respectively (see Fig. S1 in the supplemental material).

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Table 1. Plasmids used in this study

| Plasmid | Construction* | Source |
|---------|---------------|--------|
| pSH75   | trpC-hph-trpCt | 26     |
| pII99   | trpC-nptII-trpCt | 44     |
| pSB116  | trpC-bar-trpCt | This study |
| pYTGFPc | trpC-GFP-trpCt | 19     |
| pYTGFP-N| trpC-GFP-trpCt | 19     |
| pGFP-AKT1| trpC-GFP-AKT1-trpCt | This study |
| pGFP-AKT2| trpC-GFP-AKT2-trpCt | This study |
| pGFP-AKT3| trpC-GFP-AKT3-trpCt | This study |
| pGFP-AKT1Δ| trpC-GFP-AKT1Δ-trpCt | This study |
| pGFP-AKT2Δ| trpC-GFP-AKT2Δ-trpCt | This study |
| pGFP-AKT3Δ| trpC-GFP-AKT3Δ-trpCt | This study |
| pAaPEx6 | Coomaid containing AaPEx6 | This study |
| pDNR-PEx6| PEx6-trpC-nptII-trpCt::3') AaPEx6 | This study |
| pGDPEX6n | 5' AaPEx6-trpC-nptII-trpCt::3' AaPEx6 | This study |

* trpC and trpCt, A. nidulans trpC promoter and terminator, respectively (37); AKT1-PTS1, AKT2-PTS1, and AKT3-PTS1, lacking nine nucleotides encoding C-terminal PTS1-like tripeptides.
All of the PCR products cloned into the vectors were sequenced to confirm that no nucleotide substitution had occurred during amplification.

**Fungal transformation.** Protoplast preparation and transformation of A. alternata were performed as previously described (20). Transformants carrying hph or nptII were selected on regeneration medium (26) containing hygromycin B (Wako Pure Chemicals, Osaka, Japan) at 100 μg/ml or Genticin (Gibco BRL, Life Technology, Gaithersburg, MD) at 400 μg/ml (20). Protoplasts transformed with pSB116 carrying the bar cassette were regenerated on minimal agar medium (MAM) (52) supplemented with 1 M sucrose and 50 μg/ml bialaphos (Meiji Seika Kaisha, Ltd, Tokyo, Japan).

**DNA and RNA manipulation.** Isolation of total DNA and RNA from A. alternata and DNA gel blot hybridization were performed as previously described (55). For analysis of nucleotide sequences, DNA was cloned into pBluecript KS+ (Stratagene, La Jolla, CA) or pGEM-T Easy (Promega, Madison, WI). DNA sequences were determined with the BigDye Terminator v3.1 cycle sequencing kit (Applied Biosystems, Warrington, United Kingdom) and an automatic fluorescent DNA sequencer ABI PRISM 3100 genetic analyzer (Applied Biosystems). DNA sequences were analyzed with BLAST (4). Alignment of nucleotide and amino acid sequences was done with the CLUSTAL W program (59).

**Isolation of AaPEX6.** The AaPEX6 fragment was amplified from the total DNA of strain 15A by PCR using the primer pair PEX6-6/PEX6-9 (see Table S1 in the supplemental material) and cloned into the pGEM-T Easy vector (Stratagene, La Jolla, CA) with the In-Fusion Dry-Down PCR cloning kit (Clontech). The 6.5-kb target DNA fragment which includes all of the exons and introns of AaPEX6 was isolated. A 6.8-kb region in pCAPEX6 was sequenced, and a putative ORF of AaPEX6 was identified. The AaPEX6 cDNA was amplified from the total RNA of strain 15A with primer pair PEX6-6/PEX6-9 (see Table S1 in the supplemental material) and cloned into the pGEM-T Easy vector to determine the sequence.

The entire AaPEX6 gene was cloned into pDNA-P-6MV (Clontech, Mountain View, CA) with the In-Fusion Dry-Down PCR cloning kit (Clontech). The 6.5-kb fragment which includes all of the exons and introns of AaPEX6 and the 1.6-kb upstream and 0.5-kb downstream regions was amplified from pCAPEX6 DNA by PCR using primer pair AaPEX6-f/AaPEX6-r (see Table S1 in the supplemental material). The In-Fusion cloning reaction and transformation in Escherichia coli Fusion-Blue competent cells (Clontech) yielded plasmid pDNA-P-PEX6. The AaPEX6-targeting vector pGFP-PENx was prepared by replacing a 3.2-kb BamHI-EcoRI fragment within AaPEX6 in pDNA-P-PEX6 with a 1.9-kb BamHI-EcoRI fragment of the nptII cassette.

**Microscopy.** Transformants with GFP fusion constructs were grown for 3 days at 25°C on glass slides covered with a thin layer of potato agar medium supplemented with glucose or oleic acid. Infection-related morphogenesis of transformants was observed on onion epidermis (10). Some of the cell walls on onion epidermis (about 1 cm²) floating on sterilized water were separately inoculated with drops (10 μl) of each conidial suspension (about 2 × 10⁶ conidia/ml) and incubated at 25°C for 24 h. Conidial germination was observed with a microscope (59).

**Assay for vegetative growth and conidiation.** To test for vegetative growth, strains were grown on PDA at 25°C for 4 days. Agar blocks (3 mm in diameter) carrying mycelia were prepared from the resulting colonies and inoculated onto PDA and MAM supplemented with one carbon source, 1% glucose, 0.5% oleic acid, or 0.5% Tween 80. After incubation at 25°C for 5 days, colony growth was observed. To test for conidiation, strains were grown on oatmeal sucrose agar at 25°C for 10 days and induced for conidiation as previously described (14, 22).

**Assay for infection-related morphogenesis, AK-toxin production, and pathogenicity.** Conidial germination and appressorium formation were observed on PDA (25), and DNA gel blot analysis (see Fig. S2 in the supplemental material). These constructs were introduced into strain 15A by cotransformation with plasmid pSH75, which confers hygromycin B resistance (26). As a control, strain 15A was transformed with plasmid pYGFPc, which carries only GFP (see Fig. S1 in the supplemental material). Transformants were grown on PDA, and their hyphae were observed under a fluorescence microscope. Transformation of strain 15A with pYGFPc, pGFP-AKT1, pGFP-AKT2, and pGFP-AKT3 gave rise to 32, 40, 54, and 36 GFP-expressing transformants, respectively. In all of the pYGFPc transformants, the localization of GFP fluorescence was cytosolic and did not appear to be associated with any specific organelle or component of hyphal cells (Fig. 2). In contrast, GFP fluorescence localized to punctate organelles in hyphal cells of all of the pGFP-AKT1, pGFP-AKT2, and pGFP-AKT3 transformants (Fig. 2), consistent with the hypothesis that these enzymes are targeted to peroxisomes. We verified the presence of the GFP fusion constructs in the transformants by DNA gel blot analysis (see Fig. S2 in the supplemental material).
It is known that supplementation of oleic acid as a carbon source in culture media enhances the proliferation of peroxisomes in hyphal cells (62, 63). Transformants were grown on potato agar supplemented with oleic acid instead of glucose. The number of punctate organelles with GFP fluorescence markedly increased in hyphal cells expressing GFP-Akt fusion proteins grown on oleic acid medium compared with those grown on glucose medium (Fig. 2). The cytosolic localization and intensity of GFP fluorescence in pYTGFPc transformants were similar on both media (Fig. 2). These data are consistent with the hypothesis that the punctate organelles labeled by the GFP fusion proteins are peroxisomes.

To assess whether the C-terminal tripeptides in Akt1, Akt2, and Akt3-1 act as peroxisomal targeting signals, we constructed transformation vectors pGFP-AKT1Δ, pGFP-AKT1Δ, and pGFP-AKT3Δ, in which AKTI, AKT2, and AKT3-I lacking nine nucleotides that encode C-terminal PTS1-like tripeptides were fused to the 3′ end of GFP (see Fig. S1 in the supplemental material). These constructs were introduced into strain 15A by cotransformation with plasmid pSH75, and 20, 17, and 21 GFP-expressing transformants were obtained with pGFP-AKT1Δ, pGFP-AKT2Δ, and pGFP-AKT3Δ, respectively. We verified the presence of the GFP fusion constructs in the transformants by DNA gel blot analysis (see Fig. S3 in the supplemental material). When these transformants were grown on PDA, none of them showed GFP fluorescence in a punctate distribution in hyphal cells; all GFP fluorescence was cytosolic (Fig. 3). These results confirmed the peroxisome-targeting function of the C-terminal tripeptides in Akt1, Akt2, and Akt3-1 and suggested that peroxisomes are involved in AK-toxin production.

Most host-selective toxins, including AK-toxin, are produced during growth in medium and also during conidial germination (17, 28, 58, 68). Toxins released during conidial germination are required by the producing fungi to penetrate host cells and colonize tissue (17, 28, 58, 68). Conidial suspensions of transformants were dropped onto onion epidermal strips, and germinated conidia forming appressoria and penetration hyphae were observed under a fluorescence microscope at 24 h after inoculation. In pYTGFPc transformant GFP-1, the localization of GFP fluorescence was cytosolic in germ tubes, appre-
soria, and penetration hyphae (Fig. 4). In GA1-1 expressing the GFP-Akt1 fusion protein, abundant punctate organelles with GFP fluorescence were observed in germ tubes, appressoria, and penetration hyphae (Fig. 4). Similar patterns of GFP fluorescence distribution in germinated conidia were observed in GFP-Akt2- and GFP-Akt3-1-expressing transformants. Abundant peroxisomes in germinated conidia suggested that peroxisome function is important for the infection-related morphogenesis of *A. alternata*.

**Isolation and identification of Aa*PEX6**. *PEX6* encodes a peroxin protein which belongs to the AAA (ATPase associated with various cellular activities) protein family and is essential for peroxisome biogenesis in eukaryotic cells (7, 48, 60). We isolated *PEX6* from *A. alternata* (Aa*PEX6*) and analyzed the function of peroxisomes in AK-toxin production and pathogenicity in the Japanese pear pathotype by using strains in which Aa*PEX6* is disrupted.

To isolate the Aa*PEX6* gene, part of Aa*PEX6* was amplified from the DNA of strain 15A by PCR. The resulting PCR product was used as a probe to screen a cosmid genomic library of strain 15A, and a positive clone, pcAa*PEX6*, was isolated. Sequencing of the 6.8-kb region in pcAa*PEX6* detected Aa*PEX6*, which has three exons (175, 3,469, and 688 bp) divided by two introns (57 and 89 bp) and potentially encodes a 1,444-amino acid protein (Fig. 5A). The presence of two introns was confirmed by comparison of the genomic sequence with the cDNA sequence.

A database search using the BLAST algorithm revealed that the deduced amino acid sequence shows 53, 52, 50, and 30% identity with those of the Pex6 proteins from *M. oryzae* (49), *C. orbiculare* (25), *P. chrysogenum* (24), and *S. cerevisiae* (64), respectively (see Fig. S4 in the supplemental material). AaPex6 contains Walker A and B motifs and a signature AAA protein family motif (7) (see Fig. S4 in the supplemental material).

To examine the function of Aa*PEX6* in peroxisome biogenesis and AK-toxin production, homologous recombination was employed to replace Aa*PEX6* with plasmid pGDPEx6n containing an allele in which a 3.2-kb region within Aa*PEX6* had been replaced with the *nptII* cassette (Fig. 5B). Strain GA1-1 expressing the GFP-Akt1 fusion protein was transformed with pGDPEx6n, and 10 Geneticin-resistant transformants were isolated. It has been reported in yeast and other fungal species that most peroxisome-defective mutants are unable to grow on media containing fatty acids as sole carbon sources because of their defect in fatty acid β oxidation (62, 63). We observed the growth ability of GA1-1 and the GA1-1 pGDPEx6n transformants on MAM supplemented with oleic acid or Tween 80. GA1-1 and seven transformants could grow on fatty acid medium. However, the remaining three transformants (GDP6-1 to GDP6-3) exhibited very poor growth on the medium (Fig. 5C), suggesting that these transformants were deficient in fatty acid utilization due to the disruption of Aa*PEX6*.

A replacement event at the Aa*PEX6* locus in three transformants showing poor growth on fatty acid medium was confirmed by PCR analysis. The Aa*PEX6* locus was amplified from the total DNA of GA1-1 and transformants by PCR with the primer pair AaPEX6f/AaPEX6r (Fig. 5A). A PCR product of the expected size, ~6.5 kb DNA, was amplified from GA1-1, and an ~5.2-kb product, corresponding to the mutated Aa*PEX6* locus, was amplified from GDP6 transformants (Fig. 5B and D).

To further confirm the disruption of Aa*PEX6* in these

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**FIG. 4.** Intracellular localization of GFP-tagged Akt1 in infection-related structures. A sample of a conidial suspension was dropped onto onion epidermis and incubated for 24 h. GFP-1, pYTGFp: transformant; GA1-1, pGFP-AKT1 transformant; GFP, GFP fluorescence images; DIC + GFP, DIC images merged with GFP fluorescence images. c, conidium; gt, germ tube; ap, appressorium; ph, penetration hypha. Bars = 50 μm.
transformants, genetic complementation of ΔAaPEX6 mutant strain GDP6-1 with wild-type AaPEX6 was performed. Plasmid pDNR-PEX6, containing the entire AaPEX6 gene (including the exons, introns and upstream and downstream regulatory sequences) (Fig. 5A), was introduced into GDP6-1 by cotransformation with plasmid pSB116, conferring resistance to bialaphos. When bialaphos-resistant transformants were tested for growth on oleic acid or Tween 80 medium, two transformants (GDP6c-1 and GDP6c-2) could grow on fatty acid medium (Fig. 5C). PCR analysis of these transformants with the primer pair AaPEX6-f/AaPEX6-r verified the presence of the introduced 6.5-kb fragment containing the wild-type AaPEX6 gene in addition to the 5.2-kb fragment corresponding to the mutated AaPEX6 locus (Fig. 5D).

We investigated the intracellular localization of GFP-Akt1 in ΔAaPEX6 mutant strains (GDP6-1 to GDP6-3) and AaPEX6-complemented strains (GDP6c-1 and GDP6c-2). Mycelia grown on PDA were observed under a fluorescence microscope. All of the ΔAaPEX6 mutant strains showed diffuse GFP fluorescence with no punctate pattern in hyphal cells (Fig. 6). This localization was similar to that seen for GFP or GFP-AKT1 expression in the wild-type strain (Fig. 2 and 3). In contrast, abundant punctate fluorescence was observed in AaPEX6-complemented strains, similar to that seen in the parent strain, GA1-1 (Fig. 6). These results support the hypothesis that AaPEX6 is essential for peroxisome biogenesis in A. alternata.

We also investigated colony growth and conidiation of ΔAaPEX6 mutant strains on nutrient-rich media. The growth rates of ΔAaPEX6 mutant strains on PDA were lower than those of the GA1-1 and AaPEX6-complemented strains (Fig. 7A). The melanization of colonies of the mutants was similar to that of GA1-1 (Fig. 7B). Conidiation on oatmeal sucrose medium was dramatically reduced in ΔAaPEX6 mutant strains relative to that of strain GA1-1; the ΔAaPEX6 mutant strains produced 13.2 to 14.4 times fewer conidia than did GA1-1 (Fig. 7C).
AK-toxin production and pathogenicity in ΔAaPEX6 mutant strains. Strains were grown in PDB, and AK-toxin production was evaluated on the basis of the toxicity of culture filtrates to leaves of susceptible pear cultivar Nijisseki. Although culture filtrates of parent strain GA1-1 showed marked toxicity to pear leaves, those of ΔAaPEX6 mutant strains showed no toxicity (Fig. 7D). Culture filtrates of AaPEX6-complemented strains had toxicity similar to that of GA1-1 (Fig. 7D).

The Japanese pear pathotype produces two related molecular species, AK-toxins I and II (Fig. 1), with toxin I being the most abundant and biologically activity species (41, 45). Toxin I in culture filtrates was quantified by reverse-phase HPLC. Each value represents the average of two determinations.

Table 2. AK-toxin I and EDA production and pathogenicity of ΔAaPEX6 mutant strains

| Strain | AK-toxin I (μg/ml) | EDA (μg/ml) | Pathogenicity |
|--------|-------------------|-------------|---------------|
| GA1-1  | 3.3               | 24.8        | +             |
| GFP-AKT1/ΔAaPEX6 mutants | | | |
| GDP6-1 | ND*               | ND          | -             |
| GDP6-2 | ND                | ND          | -             |
| GDP6-3 | ND                | ND          | -             |
| GFP-AKT1/ΔAaPEX6/ΔAaPEX6 mutants | | | |
| GDP6c-1| 2.9               | 25.6        | +             |
| GDP6c-2| 2.5               | 22.0        | +             |

* AK-toxin I and EDA in culture filtrates were analyzed by reverse-phase HPLC. Each value represents the average of two determinations.

The strain Nijisseki was wounded slightly, treated with culture filtrate of each strain, and incubated for 24 h (D). Leaves were spray inoculated with a conidial suspension (about 5 × 10⁵ conidia/ml) of each strain and incubated for 24 h (E).
the GA1-1 and AaPEX6-complemented strains produced toxin I, those of ΔAaPEX6 mutant strains did not produce detectable toxin I (data not shown). Thus, it appeared that the presence of functional AaPEX6 is essential for AK-toxin production. These results also indicated that the peroxisome localization of peroxisomal enzymes involved in AK-toxin production is required for the biosynthesis of EDA and AK-toxin.

AK-toxin-minus mutants completely lose pathogenicity on host pear leaves (47, 55, 56). The ΔAaPEX6 mutant strains were tested for pathogenicity to Nijisseiki leaves by spray inoculation of conidial suspensions. Although the GA1-1 and AaPEX6-complemented strains caused a number of lesions on pear leaves within 24 h after inoculation, all of the ΔAaPEX6 mutant strains caused no lesions on pear leaves (Fig. 7E). This result suggested that these mutants lost pathogenicity due to loss of AK-toxin.

We attempted to restore pathogenicity to the ΔAaPEX6 mutant strain by the addition of AK-toxin I to conidia. Conidia of the Japanese pear pathotype germinate within 6 h after inoculation on host leaves, subsequently form appressoria, and begin to form penetration hyphae as early as 8 h after inoculation (16, 47). Lesions begin to appear as early as 18 h after the inoculation of conidial suspensions and become obvious after 24 h (47). We previously observed that this pathogen produced about 0.1 μg AK-toxin I per ml in the first 12 h when a conidial suspension (about $5 \times 10^5$ conidia/ml) was sprinkled onto paper towels and incubated at 25°C (16). AK-toxin I constantly causes necrosis on Nijisseki leaves at a concentration of more than 0.005 μg/ml within 24 h when a toxin solution is dropped onto wounded sites of the leaves (45). To assess whether the addition of AK-toxin can restore pathogenicity to the ΔAaPEX6 mutant strain, conidia of GA1-1 and GDP6-1 suspended in water or AK-toxin I solution (0.1 μg/ml) were tested for the ability to infect Nijisseki leaves. We could not evaluate whether the addition of AK-toxin I can restore the ability of GDP6-1 to cause lesions because spray inoculation of AK-toxin I solution with and without conidia developed approximately the same numbers of similar-size lesions on Nijisseki leaves within 24 h, due to the toxicity of AK-toxin I. Thus, we observed the elaboration of penetration hyphae at 12 h after inoculation.

Conidia of GDP6-1 germinated and formed appressoria on pear leaves at a frequency similar to that of conidia of GA1-1, and the addition of AK-toxin I to the conidia had no significant effect on conidial germination and appressorium formation (Fig. 8). In GA1-1, about 38% of the appressoria formed penetration hyphae after 12 h when conidia were suspended in water or AK-toxin I solution (Fig. 8). The penetration frequency was dramatically reduced in GDP6-1 relative to GA1-1: only 3.2% of the appressoria formed penetration hyphae in water (Fig. 8). The penetration ability of GDP6-1 was partially, but not completely, restored by the addition of AK-toxin I to conidia: 11.2% of the appressoria formed penetration hyphae in the presence of AK-toxin I (Fig. 8). Restoration of the penetration ability of GDP6-1 was not observed in the presence of 0.01 μg/ml AK-toxin I. These results suggested that peroxisome function is involved in the penetration ability, in addition to AK-toxin production, of the Japanese pear pathotype.

**Infection-related morphogenesis of ΔAaPEX6 mutant strains.** We further observed the infection-related morphogenesis in ΔAaPEX6 mutant strains on glass and onion epidermis. To observe conidial germination and appressorium formation, conidial suspensions in water were dropped onto glass and incubated for 24 h. There were no significant differences in conidial germination, germ tube elongation, or appressorium formation among the parent, AaPEX6 disruption-containing, and AaPEX6-complemented strains (Fig. 9A; see Fig. S5 in the supplemental material). To determine the ability of ΔAaPEX6 mutant strains to penetrate plant epidermal cells, conidial suspensions were dropped onto onion epidermal strips and monitored to detect the elaboration of penetration hyphae at 24 h. The ΔAaPEX6 mutant strains successfully formed penetration hyphae from appressoria at a frequency similar to those of the parent and AaPEX6-complemented strains (Fig. 9B and C), suggesting that peroxisome function is not essential for the penetration of onion epidermal cells by *A. alternata*.

*A. alternata* conidia are known to invade leaf tissue and produce visible lesions on leaves of resistant pear cultivars with defense responses partially compromised by heat shock (46). Leaves of the resistant pear cultivar Chojuro, which are insensitive to AK-toxin and completely resistant to the Japanese pear pathotype, were dipped in water at 50°C for 50 s, spray inoculated with a conidial suspension, and then incubated for 24 h. The parent, AaPEX6 disruption-containing, and AaPEX6-complemented strains caused no lesions on untreated leaves (Fig. 10A). The parent and complemented strains caused many visible lesions on heat-shocked leaves (Fig. 10B). In contrast, the ΔAaPEX6 mutant strains produced fewer and smaller lesions than the parent strain (Fig. 10B). Together with the finding that the addition of AK-toxin I to conidia partially, but not completely, restored the penetration ability of the ΔAaPEX6 mutant strain, this result suggested that loss of peroxisome function results in a
reduction in the penetration into pear leaf epidermal cells and the proliferation in leaf tissue of this *A. alternata* pathotype.

**DISCUSSION**

**Contribution of peroxisomes to AK-toxin biosynthesis and pathogenicity.** In this study of the Japanese pear pathotype of *A. alternata*, we verified that the AK-toxin biosynthetic enzymes Akt1, Akt2, and Akt3-1, which have PTS1 tripeptides at their C-terminal ends, are localized in peroxisomes. To assess the role of peroxisome function in AK-toxin biosynthesis, we isolated *AaPEX6*, which encodes a peroxin protein essential for peroxisome biogenesis, from the Japanese pear pathotype of *A. alternata*. *AaPEX6* mutant strains failed to form functional peroxisomes and completely lost AK-toxin production and pathogenicity. Because the strawberry and tangerine pathotypes have closely related orthologs of *AKT1*, *AKT2*, and *AKT3* which are involved in the biosynthesis of AF-toxin and ACT-toxin (15, 33, 35), Akt1, Akt2, and Akt3 of the Japanese pear pathotype and their orthologs in the strawberry and tangerine pathotypes participate in the biosynthesis of EDA, a common moiety of AK-, AF-, and ACT-toxins. Indeed, the *AaPEX6* mutant strains of the Japanese pear pathotype failed to produce not only AK-toxin but also EDA. EDA is detected in culture filtrates of the Japanese pear, strawberry, and tangerine pathotypes (12, 15, 34). When 3H-labeled EDA was added to a growing liquid culture of the Japanese pear pathotype strain, it was efficiently converted to AK-toxin (12). This result clearly shows that EDA is an intermediate in the AK-toxin pathway. A 13C nuclear magnetic resonance analysis of EDA purified from culture filtrates supplemented with [2-13C]sodium acetate suggested that EDA is biosynthesized by the condensation of six molecules of acetic acid, followed by modifications, including reduction, dehydration, and decarboxylation (42). We identified *AFT9*, which encodes a polyketide synthase (PKS), in the AF-toxin biosynthetic gene cluster of the strawberry pathotype and found the *AFT9* homologues from the Japanese pear and tangerine pathotypes by DNA gel blot analysis (51). We verified that the *AFT9* ortholog of the Japanese pear pathotype, *AKT9*, resides in the AK-toxin biosynthetic gene cluster (S. Takaoka and T. Tsuge, unpublished data). It is likely that the acetyl-CoA-derived backbone of the EDA molecule is produced by the activity of PKS and then modified by other enzymes, including Akt1, Akt2, and Akt3.

In addition to *AFT9* and *AKT9*, we have identified several other genes involved in toxin production by the three pathotypes (1, 15, 20, 34, 35, 36, 51). They include genes common to all three pathotypes, which are probably involved in EDA biosynthesis, and genes specific to each pathotype. Analysis of their predicted amino acid sequences by various sorting algorithms did not identify potential PTS sequences in any of these proteins other than Akt1, Akt2, and Akt3 of the Japanese pear.
pathotypes and their orthologs in the strawberry and tangerine pathotypes. While these data suggest that only part of the EDA biosynthetic pathway takes place in peroxisomes, peroxisomal proteins lacking either of the two known PTS sequences have been found (48, 60), and a definitive subcellular localization of the other proteins involved in toxin production in hyphal cells remains to be determined.

The importance of peroxisomes in secondary metabolism has been reported in penicillin biosynthesis in *P. chrysogenum* (38, 39). Penicillin biosynthesis consists of at least three enzymatic steps (23, 24, 38, 39). The first step is a nonribosomal condensation of activated l-α-aminoacidic acid, l-cysteine, and l-valine to δ-(l-α-aminoacidipyl)-l-cysteinyl-d-valine tripeptide (ACV), catalyzed by a single enzyme, ACV synthetase. In the second step, isopenicillin N synthetase converts ACV to isopenicillin N. The final step is the exchange of the acyl-CoA side chain of isopenicillin N for a hydrophobic side chain catalyzed by acyl-CoA:isopenicillin N acyltransferase. It was suggested that ACV synthetase and isopenicillin N synthetase are membrane-associated and cytosolic enzymes, respectively (39), while acyl-CoA:isopenicillin N acyltransferase has a PTS1 sequence and is located in peroxisomes (38, 39). Thus, the three steps of penicillin biosynthesis take place in different cellular domains.

The possible involvement of peroxisomes has also been suggested for the biosynthesis of two polyketide mycotoxins, aflatoxin produced by *Aspergillus flavus* and *A. parasiticus* and sterigmatocystin produced by *A. nidulans* (32). Sterigmatocystin is the penultimate intermediate in aflatoxin biosynthesis (23). Although peroxisomal localization of any of aflatoxin biosynthetic enzymes has not been observed, a precursor, norsolorinic acid (NOR), was found to accumulate in peroxisomes of all three *Aspergillus* species (32). NOR, a fluorescent compound, is produced by the activity of PKS (StcA) and fatty acid synthase (α subunit StcJ and β subunit StcK), which form a complex (67). These enzymes lack either of the two known PTS sequences, and their peroxisomal localization has not been verified. However, NOR ketoreductase, which lacks either of the two PTS sequences and catalyzes the next step of NOR synthesis in the aflatoxin pathway, has been observed in the cytoplasm of *A. parasiticus* (30). Thus, it is likely that biosynthetic pathways of the penicillins of *P. chrysogenum*, aflatoxin of *Aspergillus* spp., and EDA ether toxins require intracellular transport steps for intermediates and/or products.

Host-selective toxin biosynthetic genes of *A. alternata* pathotypes appeared to be clustered on small chromosomes of <2.0 Mb in most of the strains tested (2, 3, 13, 15, 20, 21, 33, 35). We demonstrated that small chromosomes that contain toxin biosynthetic genes from the strawberry, apple, and tomato pathotypes are conditionally dispensable (CD) chromosomes (2, 15, 21). We are now focusing on the structural and functional analysis of the 1.05-Mb CD chromosome that contains the *AFT* genes of the strawberry pathotype to identify the entire gene cluster and to understand the evolution of toxin biosynthesis and the origin of CD chromosomes. Identification of the entire gene cluster and subsequent analysis of the predicted amino acid sequences of AF-toxin pathway enzymes by sorting algorithms should provide more information concerning the subcellular localization of EDA and EDA ether toxin biosynthesis in *A. alternata*.

**Contribution of peroxisomes to plant infection.** *ΔP*EX6 mutant strains of *C. orbiculare* and *M. oryzae* completely lose pathogenicity (25, 49, 65). These pathogens produce dome-shaped appressoria which are darkly pigmented with dihydroxynaphthalene melanin (61). *ΔP*EX6 mutant strains of these pathogens form smaller appressoria with severely reduced melanization that fail to form infection hyphae (25, 49, 65). The reduced melanization of *ΔP*EX6 mutant strains suggests that melanin biosynthesis in appressoria depends on acetyl-CoA produced by fatty acid β oxidation in peroxisomes. *A. alternata* also produces dihydroxynaphthalene melanin, which accumulates in the cell walls of conidia and hyphae (22, 26, 54). However, its appressoria are smaller than those of *C. orbiculare* and *M. oryzae* and are colorless. We observed that melanin-deficient mutants and the wild-type strain of the Japanese pear pathotype developed approximately the same numbers of similar-size lesions on susceptible pear leaves in laboratory tests (22, 54). Thus, the ability to produce melanin is
probably not relevant to plant infection by the Japanese pear pathotype of *A. alternata*.

We examined the infection-related morphogenesis of the ΔAaPEX6 mutant strains of the Japanese pear pathotype. Conidia of the ΔAaPEX6 mutant strains germinated and formed appressoria as did those of the wild type on glass and host pear leaves. When the ability to penetrate onion epidermal cells was tested, the frequency of successful production of penetration hyphae from appressoria of the ΔAaPEX6 mutant strains was not significantly different from that of the wild type, suggesting that the mutants retain the basic appressorium-mediated penetration ability. However, the ability of the ΔAaPEX6 mutant strains to penetrate host epidermal cells could be partially, but not completely, restored by the addition of AK-toxin I to a conidal suspension prior to inoculation. When conidia of the ΔAaPEX6 mutant strains were used to inoculate leaves of a resistant pear cultivar with defense responses partially compromised by heat shock, they caused fewer and smaller lesions on the leaves than did those of the wild-type strain. We also observed that germ tubes, appressoria, and penetration hyphae of this pathogen contained abundant peroxisomes. These results suggest that peroxisome function is also necessary for initial plant infection and for tissue colonization by this *A. alternata* pathotype.

The ΔPEX6 mutant strains of *M. oryzae* retained the capacity to form polarized infection hyphae on onion epidermal layers, albeit at a reduced frequency (65). The loss of pathogenicity of ΔPEX6 mutant strains of *M. oryzae* was not solely attributable to the lack of appressorium function because wounded seedlings inoculated with the mutants did not develop rice blast symptoms or support fungal growth (65). Virulence could be partially, but not completely, restored to the mutants by the addition of glucose to a conidal suspension (65). We also observed that the AaPEX6 deletion mutation caused a marked reduction of conidiation in *A. alternata*. The AaPEX6 mutant strains produced about 13 times fewer conidia than the wild type. Because the conidiation medium contains sucrose as a carbon source, peroxisome functions other than fatty acid β-oxidation and glyoxylate metabolism are likely to contribute to conidiation. Conidiation in the *M. oryzae* ΔPEX6 mutant strains was also dramatically reduced; ΔPEX6 mutant strains produced about 40 times fewer conidia than wild-type strains (65). These data suggest that the diverse functions of the peroxisomes, including β-oxidation, glyoxylate metabolism, and peroxide detoxification, are required for plant pathogenesis in fungi.

In *M. oryzae* and *C. orbiculare*, the glyoxylate cycle, the enzymes of which are localized in peroxisomes, has been shown to be necessary for full virulence by using ΔCICL1 mutants, which lack isocitrate lyase (6, 66). The importance of the glyoxylate cycle in plant pathogenesis has also been shown in the *Brassica* pathogen *Leptosphaeria maculans* (18) and the wheat pathogen *Stagonospora nodorum* (53). Peroxisome-associated carnitine acetyltransferase is required for the elaboration of penetration hyphae during infection by *M. oryzae* (8, 49). Carnitine acetyltransferase catalyzes the conversion of acetyl-CoA into acetylcarnitine prior to the transport of acetyl units from peroxisomes to the correct intracellular compartment for subsequent utilization (11). The ΔCART1 (ΔPTH2) mutants, which lack carnitine acetyltransferase, had weakened cell walls, probably due to defects in cell wall biosynthesis (49). Recently, it has been reported that host invasion by *C. orbiculare* requires Atg26, a sterol glucosyltransferase that activates pexophagy (5). It is likely that plant pathogenesis in fungi depends on several peroxisome functions, including fatty acid metabolism, acetyl-CoA generation, secondary metabolism, cell wall biogenesis, and peroxisome homeostasis.

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