The P450 Monooxygenase BcABA1 Is Essential for Abscisic Acid Biosynthesis in Botrytis cinerea
Verena Siewers, Jörn Smedsgaard, and Paul Tudzynski

Institut für Botanik, Westfälische Wilhelms-Universität, D-48149 Münster, Germany, and Bio-Centrum DTU, DK-2800 Lyngby, Denmark

Received 7 January 2004/Accepted 22 March 2004

The phytopathogenic ascomycete Botrytis cinerea is known to produce abscisic acid (ABA), which is thought to be involved in host-pathogen interaction. Biochemical analyses had previously shown that, in contrast to higher plants, the fungal ABA biosynthesis probably does not proceed via carotenoids but involves direct cyclization of farnesyl diphosphate and subsequent oxidation steps. We present here evidence that this “direct” pathway is indeed the only one used by an ABA-overproducing strain of B. cinerea. Targeted inactivation of the gene bcab1 encoding a cytochrome P450 oxidoreductase reduced the ABA production significantly, proving the involvement of P450 monoxygenases in the pathway. Expression analysis of 28 different putative P450 monooxygenase genes revealed two that were induced under ABA biosynthesis conditions. Targeted inactivation showed that one of these, bcab1, is essential for ABA biosynthesis: Δbcab1 mutants contained no residual ABA. Thus, bcab1 represents the first identified fungal ABA biosynthetic gene.

Fungi, especially phytopathogenic species, have been shown to produce all major classes of phytohormones (for a review, see reference 35). The best investigated fungal phytohormone system (and the only one in which the biosynthetic genes involved have been identified) is that of Gibberella fujikuroi. It has been shown that the genes involved in gibberellic acid (GA3) biosynthesis are arranged in a cluster (34) and that the biosynthesis in major respects is different from the higher plant pathway (12). In this case, therefore, the original hypothesis that fungal biosynthesis of phytohormones represents a good example for a horizontal gene transfer (6) is negated. This view appears to negatively modulate the salicylic acid-dependent defense pathway in tomato plants during infection with B. cinerea. Thus, ABA could represent a virulence factor for the fungus. The unequivocal proof for a role of fungal ABA in the host-pathogen interaction would require defined mutants, which are absolutely unable to produce ABA in planta. Strains not producing detectable amounts of ABA in axenic culture have been described (21), but it is not clear whether they still have the capability to produce ABA in planta. Therefore, for several reasons the identification of genes involved in the ABA pathway in B. cinerea would be interesting: it would advance evolutionary research by allowing comparison of the pathways in higher plants and fungi, lead to advanced biotechnological methods for the generation of overproducing strains, and provide new insight into phytopathological investigations as indicated above. The objective of the present study was to find conclusive evidence for the postulated alternative ABA biosynthetic pathway in B. cinerea. We show here that P450 monoxygenases are involved in ABA biosynthesis, by targeted inactivation of a cytochrome P450 oxidoreductase gene, and we describe the identification of the first fungal gene involved in ABA biosynthesis, encoding a P450 monooxygenase.

MATERIALS AND METHODS

Fungal strains. Strain ATCC 58025 of B. cinerea Pers.:Fr. (Botryotinia fuckeliana [de Bary] Whetzel) is a nonsporulating overproducer of ABA (18), and B05.10, a haploid strain were obtained after benomyl treatment of strain SAS5 (25).

Bacterial strains. Escherichia coli strain TOP10F* (Invitrogen, Groningen, The Netherlands) was used for propagation of plasmids. Propagation of lambda clones was performed in strains LE392 (Stratagene, La Jolla, Calif.) and XL1-Blue MRF* (Stratagene), respectively.

* Corresponding author. Mailing address: Institut für Botanik, Westfälische Wilhelms-Universität Münster, Schlosergarten 3, 48149 Münster, Germany. Phone: 49-251-83 21601. Fax: 49-251-83 24998. E-mail: tudzyns@uni-muenster.de.
higher plants

FIG. 1. Postulated biosynthetic pathways of ABA in higher plants and in C. pini-densiflora (19, 22), respectively. Intermediates identified in C. pini-densiflora are α-onylidenecacetonic acid (structure 1), ω-SH-α-onylidenethanol (structure 2), 1'-OH-α-onylidenethanol (structure 3), (1'R)-4'R-OH-α-onylidenecetonic acid (structure 4), (1'R)-4'-5-OH-α-onylidenecetonic acid (structure 5), and 1'-deoxy-ABA (structure 6). The site of action of DPA is indicated.

Media and culture conditions. B. cinerea strains were grown on 2% malt extract (Oxoid, Ltd., Basingstoke, Hampshire, England) amended with 0.5% glucose, 0.1% casein peptone, 0.1% yeast extract (Difco Laboratories, Sparks, Md.), 0.1% Casamino Acids (Difco), 0.1% yeast extract (Duchefa Biochemie BV, Haarlem, The Netherlands), and 0.02% RNA sodium salt. For ABA production, fungi were cultivated on 50-ml Erlenmeyer flasks with 100 ml of a defined liquid medium (31; modified as described by Kettner [14] containing 20 g of lactose/liter instead of glucose (Sprecher medium). Fungi and culture filtrates were harvested after 3 to 7 days of cultivation on a rotary shaker at 150 rpm and 20°C. For DNA isolation, mycelium was grown for 3 to 4 days at 20°C on complex medium agar (24) with a cellophane overlay.

DNA isolation. Fungal genomic DNA was isolated as described by Cenis (4). Lambda DNA was isolated according to the standard method (28). Plasmid DNA was isolated by using a plasmid DNA preparation kit (Genomed, Bad Oeynhausen, Germany).

Southern blot analysis. Genomic DNA was digested with restriction enzymes, size separated on a 1% agarose gel, and blotted onto Hybond-N²/0.1% sodium dodecyl sulfate, and 50 mM phosphate buffer (pH 6.6) at 65°C for 16 to 20 h in the presence of a random-primed [-32P]dCTP-labeled probe. Membranes were washed in 2X SSPE-0.1% sodium dodecyl sulfate at 65°C before being exposed to an autoradiographic film.

RNA blot analysis. RNA was isolated from mycelial samples by using the RNAgent total RNA isolation system (Promega, Mannheim, Germany). Samples of 10 to 15 µg of RNA were transferred to nylon filters by electroblotting on a 1% agarose gel containing formaldehyde according to the method of Sambrook et al (28). Blot hybridizations were carried out in 0.6 M NaCl, 0.16 M Na2HPO4, 0.06 M EDTA, 1% N-lauroylsarcosine (Sigma), and 10% dextran sulfate (Eppendorf AG, Hamburg, Germany) (pH 6.5) as described for the Southern blots.

Sequencing. DNA sequencing of recombinant plasmid clones was performed with an automated sequencer LI-COR 4200 (MWG Biotech, Munich, Germany) by using the ThermoSequenase fluorescence-labeled primer cycle sequencing kit (Amersham Pharmacia). For sequence analysis and construction of phylogenetic trees, the program DNAStar (Madison, Wis.) was used.

Cloning of becpr1 gene. becpr1 was cloned in a PCR approach by amplifying B05.10 genomic DNA with degenerate primers CPR1 and CPR2 (kindly provided by C. Wasmann, University of Arizona). PCRs contained 25 ng of DNA, 10 pmol of each primer, 200 nM concentrations of deoxyribonucleotide triphosphates, and 1 U of Taq polymerase (Red Taq; Sigma-Aldrich, Deisenhofen, Germany) and were carried out at 94°C for 4 min, followed by 35 cycles of 94°C for 1 min, 48°C for 1 min, 72°C for 1 min, 48°C for 1 min, and 72°C for 1 min. The primers were CPR1 (5'-AAG YTG CAG ACY CGC TAC TAY TCS ATC TC-3') and CPR2 (5'-5'-CCT CAA YTC RTC CTT GTA SAR GAA RTC CTC-3'). The resulting 0.5-kb PCR fragment was used as a probe for screening a genomic EMBL3 library of B. cinerea strain SAS56 (25). Positive and purified phages were subcloned in pUC19 (41) and pBluescriptII SK+ (Stratagene), respectively.

Construction of a replacement and a complementation vector for becpr1. For construction of the gene replacement vector pCPR1Rep, the plasmid pOliHP (27) carrying the E. coli bgmR gene (gene hph under control of the Aspergillus niger oliC promoter and tspR terminator was used as a vector. A 0.4-kb PCR fragment was amplified from the 5' region of becpr1 by using the primers CPR5'-F (primer 1; 5'-CAC TGA GAA GAT CTC ATT GC-3') and CPR5'-R (primer 2; 5'-5'TTC TTG TAG CGT CGA CTT TCT GC-3') (artificial XhoI and SalI sites, respectively, are indicated in boldface). From the 3' end of becpr1, a 0.4-kb fragment was amplified by using the primers CPR3'-F (primer 3; 5'-AAG CCG TGT TGA AGC TTT GAG G-3') and CPR3'-R (primer 4; 5'-AAT ATC TCT AGC TTT GAA TTC TGG-3') (artificial HindIII and EcoRI sites, respectively, are indicated in boldface). Both fragments were cloned into pcR2.1-TOPO (Invitrogen), cut with XhoI/SalI and HindIII/EcoRI, respectively, and cloned into the corresponding sites of pOliHP. By cutting with XhoI and EcoRI, the 3.5-kb replacement cassette was isolated from the vector prior to transformation.

For construction of the complementation vector pComCPR1, the 2.7-kb HindIII fragment containing the 5' region of becpr1, including 1.5-kb promoter sequence, was cloned into pnRI (17), a vector that carries the Streptomyces noursei nourseothricin acetyltransferase gene nptII under control of the A. niger oliC promoter and a B. cinerea β-tubulin transcription terminator fragment. Using the internal ClaI site of the HindIII fragment (see Fig. 4), the 2.2-kb ClaI fragment comprising the 3' region of becpr1 was fused to this construct.

Cloning of the babel1 gene. A cDNA fragment derived from an expressed sequence tag (EST) library of strain ATCC 58025 under ABA biosynthesis conditions (V. Sievers, D. Tapadar, P. Schreier, and P. Tuzdinsky, unpublished data) was used to screen the EMBL3 library of SAS56.

Construction of a replacement vector and a complementation vector for babel1. The following primers were used to amplify the 5' and 3' regions of babel1 (see Fig. 7): ABA1-5'-F (primer 10; 5'-ATC ACC ACT CGC GGA TCT GCC CGG-3'), ABA1-5'-R (primer 11; 5'-CAT ATG GAT GTA GTC GTC CCC-3'), ABA1-3'-F (primer 12; 5'-CAT ATG CTC TCA CAA AGC TCT GCC AGC-3'), and ABA1-3'-R (primer 13; 5'-ATA GCA GAT AGC AAA TAA ATG AGG-3'). Primer ABA1-5'-R contains an artificial Sall site, and primer ABA1-3'-F contains an artificial HindIII site (indicated in boldface). Both fragments were cloned into pcR2.1-TOPO, cut with XhoI/SalI and HindIII/EcoRI, respectively, and cloned into the corresponding sites of pOliHP to construct pABA1Rep.

The replacement cassette was isolated from the vector prior to transformation by cutting both Xhol and EcoRI.
In order to construct the complementation vector pComABA1, a 4.1-kb ClaI fragment containing the complete bccaba1 gene (see Fig. 7) was ligated into pNR1.

Transformation of B. cinerea. Protocols for protoplast formation were adapted from an established procedure (32). Mycelium derived from cultures grown for 3 days on complex medium agar with a cellophane overlay was ground in a microfluidizer (Worthington, Freehold, NJ) and then diluted at 20°C and 90 rpm for 1 h at 15 min. Protoplasts were collected by using sterile cheese cloth, pelleted by centrifugation for 10 min at 1,500 × g, and resuspended in KCl buffer. DNA (10 to 15 μg) was added to 10^7 protoplasts in 100 μl of KCl buffer.

Transformed protoplasts were added to 100 ml of liquid medium (1.2%) containing 6.0 M sucrose, 5 mM Tris (pH 6.5), and 1 mM (NH4)2HPO4 (SH agar) prior to pouring on plates. After 24 h, the plates were overlaid with SH agar supplemented with 10 μg of hygromycin B (Calbiochem)/ml or with 70 μg of hygromycin B and 140 μg of nourseothricin (Werner-Biologents, Jena, Germany)/ml. Resistant colonies were transferred to agar plates containing Gamborg’s B5 medium (Duchefa Biochemie BV, Haarlem, The Netherlands) and 2% glucose complemented with 70 μg of hygromycin B/ml or with 70 μg of hygromycin B and 140 μg of nourseothricin/ml.

Homologous integration events and complementation were identified by PCR with the following primers (see Fig. 4 and 7, respectively): pLOF-oilp (primer 6: 5'GGT ACT GCC CCA CTG AGT GGC AGC TCG CG-3'), pAN-T (primer 7: 5'GCC AGA ATG CAC AGG TAC AC-3'), CPR-Bc1 (primer 5: 5'CPR-Bc1 (primer 5: 5'GCC AGA ATG CAC AGG TAC AC-3'), CPR-Bc2 (primer 8: 5'GGT ATT GAT TGG CGG ATC GC-3'). P16c (primer 16; 5'H11032/H11032/H11032-3'GGA GCC AGA CTC TCA GAT TGG CGG ATC GC-3'), and P16e (primer 16; 5'CPR-Bc2 (primer 8; 5'GGT ACT GCC CCA CTG AGT GGC AGC TCG CG-3').

EIA for cis- and trans-zeatin, fusicoccin acid. Enzyme immunoassay (EIA) for detection of ABA in culture filtrates was performed on 96-well microplates (flat bottom; Sarstedt, Inc., Newton, N.C.) in triplicate measurements as described by Weiler (39). A rabbit polyclonal antibody raised against mouse immunoglobulin G (RAMIG), an anti-ABA mouse monoclonal antibody, and an ABA-labeled alkaline phosphatase (tracer) were kindly provided by E. W. Weiler, Ruhr University, Bochum, Germany. p-Nitrophenyl phosphate (Biomol Feinchemikalien GmbH, Hamburg, Germany) was used as phosphatase substrate. ABA standards ranged from 0.05 to 50 pmol/ml. For each sample, ABA was assayed in at least two dilutions. The optical density at 405 nm was measured by using a microplate reader (model 550; Bio-Rad Laboratories GmbH, Munich, Germany).

Liquid chromatography-mass spectrometry (LC-MS) analyses. Fungi were grown on alkaloid-forming agar (26) for 7 days in the dark. Culture extracts were analyzed by EIA (see Materials and Methods). Since the amount of ABA produced by the control and the inhibitor-treated culture did not differ significantly (data not shown), the inhibitor obviously had no effect on ABA synthesis by strain ATCC 58025. This supports the view that also in B. cinerea the carotenoid pathway is not involved in ABA biosynthesis.

Cytochrome P450 monooxygenases are involved in ABA biosynthesis in B. cinerea. The postulated pathway for ABA biosynthesis in B. cinerea (Fig. 1) involves several hydroxylation and oxidation steps, which in fungi regularly are catalyzed by P450 monooxygenases (reviewed by van den Brink et al. [38]). Since B. cinerea probably possesses more than 50 enzymes of this class (42), we first used a general approach to test whether indeed P450 monooxygenases (and not dioxygenases) are likely to be involved in ABA biosynthesis. In spite of the high number of P450 monooxygenases, most fungi seem to have only one cytochrome P450 oxidoreductase which acts as the unspecific partner for all of the specific monooxygenases facilitating the electron transfer from NADPH via FAD and FMN to the heme group of the P450 enzyme (37).

Cytochrome P450 oxidoreductase (CPR) genes have been isolated and functionally characterized thus far only from a few fungi such as Aspergillus niger (37) and G. fujikuroi (17). Based on these fungal cpr sequences, degenerate primers were used to amplify a PCR fragment from genomic DNA of B. cinerea showing significant homology to cpr genes of fungi (data not shown). The PCR fragment was used as probe to screen a genomic EMBL3 library of B. cinerea. Two positive plaques were purified and then subcloned. Sequence analysis revealed an open reading frame of 2,348 bp interrupted by two introns of 58 and 211 bp, respectively; it encodes a polypeptide of 692 amino acids corresponding to a molecular mass of 77 kDa. The derived gene product has significant overall homology to fungal CPRs, its closest homologues being the cpr genes of G. fujikuroi and A. niger with 75 and 67% identity, respectively (Fig. 2), and thus represents a bona fide cytochrome P450 oxidoreductase. The gene was therefore named bccpr1.

Since Malonect et al. (17) found that the single cpr gene of G. fujikuroi is induced under gibberellin biosynthesis conditions, we performed a Northern analysis to test the expression of bccpr1. As shown in Fig. 3A, the gene is induced under conditions of ABA biosynthesis. The addition of mevalonic acid, the ABA precursor, led to an increased expression of bccpr1 for up to 120 min (Fig. 3B). This expression pattern...
supports the possible role of P450 monooxygenases in ABA biosynthesis. To prove this role unequivocally, we performed a targeted gene inactivation by using a gene replacement approach as outlined in Fig. 4 (for details, see Materials and Methods). The gene replacement fragment was used for transformation of \textit{B. cinerea} protoplasts. Hygromycin-resistant transformants were screened by PCR for the presence of a knockout allele, with primers 5 and 6 and primers 7 and 8, respectively (Fig. 4). Transformants yielding the corresponding diagnostic PCR fragments were genetically purified by protoplast subculturing (the standard method of single-spore isolation was not applicable because the recipient strain ATCC 58025 does not sporulate) and analyzed again by PCR. Transformants showing only the diagnostic PCR fragments, but not the wild-type-specific fragment of 1 kb generated by primers 5 and 9 (data not shown), were analyzed by Southern hybridization: genomic DNA of the wild type and the three different transformants was digested with XbaI, separated in an agarose gel, and transferred to a nylon membrane. As shown in Fig. 5A, transformants \textit{ΔBccpr1-3}, \textit{ΔBccpr1-8}, and \textit{ΔBccpr1-16} lack the wild-type fragment of 1.9 kb hybridizing to the labeled \textit{bccpr1} 3’ flank and show the 1.03-kb fragment expected for the knockout situation (Fig. 4), proving that they are homokaryotic deletion mutants. A Northern analysis confirmed this conclusion: in the mycelia of the mutants grown under ABA biosynthesis conditions, no \textit{bccpr1} transcript could be detected (Fig. 5B). The mutants showed a slightly reduced growth rate in axenic culture but retained the wild-type morphology (data not shown). An EIA analysis showed that mutants \textit{ΔBccpr1-8} and \textit{ΔBccpr1-16} produced significantly less ABA than the wild type (Table 1).

This finding was confirmed by LC-MS analysis (Fig. 6) by using accurate mass and analysis of a standard. ABA elutes at 5.305 min, and the mass spectrum shows a significant ion at \textit{m/z} 247.1335 due to the loss of water from the protonated molecular ion at \textit{m/z} 265.1483 (Fig. 6A). Using narrow ion traces with a width of 30 ppm around the significant ions, shown for \textit{m/z} 247.1335 in Fig. 6B, the presence of ABA was confirmed in

**FIG. 2.** Cladogram of fungal cytochrome P450 oxidoreductases based on amino acid sequences. The accession numbers are as follows: \textit{Aspergillus niger} CprA, CAA81550; \textit{Candida maltosa} NCP1, P50126; \textit{Candida tropicalis} NCP1, P37201; \textit{Coriolus versicolor} CPR, BABB3588; \textit{Cunninghamella echinulata} CPR, AAF89959; \textit{Cunninghamella elegans} CPR, AAF89958; \textit{Gibberella fujikuroi} CPRGf, AJ576025; \textit{Phanerochaete chrysosporium} CPR, AAG31350; \textit{Rhizopus stolonifer} CPR isoenzyme 1, AAG23833; \textit{Rhizopus stolonifer} CPR isoenzyme 2, AAG23834; \textit{Saccharomyces cerevisiae} NCP1, P16603; and \textit{Schizosaccharomyces pombe} CCR1, P36587.

**FIG. 3.** Expression analysis of genes \textit{bccpr1}, \textit{bcaba1}, and \textit{P450-12}. (A) Total RNA was extracted from ABA-producing \textit{B. cinerea} strain ATCC 58025 (lanes +) and nonproducing strain B05.10 grown for 4 days (\textit{bccpr1} and \textit{bcaba1}) or 11 days (\textit{P450-12}) in Sprecher medium. (B) Total RNA was extracted from strain ATCC 58025 grown for 3 days in Sprecher medium. Mycelia were harvested 30, 60, 90, 120, and 180 min after the addition of 3.8 mM mevalonic acid lactone. Fragments of the 3’ region of \textit{bccpr1} (see Fig. 4), the 5’ region of \textit{bcaba1} (see Fig. 7), and an internal 720-bp PCR fragment of \textit{P450-12} were used for probing. Loading of lanes with RNA was checked by probing with ribosomal DNA.
the wild type, but it is not found in the mutants. Both mutants show a drastic reduction of other (nonidentified) metabolites produced, as expected since P450 monooxygenases are involved in many pathways (data not shown).

To be sure that the loss of ABA production is the result of deletion of \textit{bccpr1}, we performed a complementation test: mutant \textit{\textit{H9004}Bccpr1-8} was transformed with vector \textit{pComCPR1} containing a full-length genomic copy of \textit{bccpr1} (see Materials and Methods). One of the complemented mutants obtained, \textit{Bccpr1C5}, was shown to contain obviously more than one correctly sized wild-type copies of the gene (Fig. 5A). Northern analysis confirmed the transcription of the \textit{bccpr1} gene copies (Fig. 5B). EIA analyses proved a complementation of the mutant phenotype, i.e., ABA production in the complemented mutant \textit{Bccpr1C5} reached the level of the wild type again (Table 1), finally proving that the loss of ABA production in the mutants was caused by the deletion of \textit{bccpr1}.

Taken together, these data strongly suggest that P450 monooxygenases are involved in ABA biosynthesis. Therefore, a screening for candidate ABA biosynthesis genes within the P450 monooxygenase gene family of \textit{B. cinerea} was performed. Altogether, 28 different putative P450 monooxygenase genes identified in different EST libraries of \textit{B. cinerea} were probed in a dot blot analysis with labeled cDNAs from ABA-producing and -nonproducing mycelia, respectively. Two of them, \textit{bcP450-12} and \textit{bcaba1}, showed increased expression under ABA biosynthesis conditions (data not shown). This was confirmed by Northern analysis (Fig. 3A): both genes are strongly induced under ABA biosynthesis conditions and show almost no constitutive expression. The addition of the precursor mevalonic acid (Fig. 3B) resulted in a significant and persistent induction of \textit{bcaba1} after 90 min, whereas \textit{P450-12} showed a transient induction at 60 min. Thus, both genes are candidates for ABA biosynthetic genes and, because of the persistent expression under mevalonic acid, we focused on \textit{bcaba1}. The genomic copy of \textit{bcaba1} was isolated from the genomic EMBL3 library of strain SAS56 by using the cDNA clone as a probe. Sequencing of subcloned lambda fragments hybridizing to the probe revealed an open reading frame of 1,769 bp.

### Table 1. ABA production of \textit{B. cinerea} wild-type and mutant strains

<table>
<thead>
<tr>
<th>Strain</th>
<th>ABA (nmol liter(^{-1}))</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>ATCC 58025</td>
<td>2,927</td>
<td>621</td>
</tr>
<tr>
<td>\textit{\textit{H9004}Bccpr1-8}</td>
<td>3.77</td>
<td>3.75</td>
</tr>
<tr>
<td>\textit{\textit{H9004}Bccpr1-16}</td>
<td>3.14</td>
<td>1.84</td>
</tr>
<tr>
<td>\textit{Bccpr1C5}</td>
<td>3.374</td>
<td>582</td>
</tr>
<tr>
<td>\textit{Bcaba1-2}</td>
<td>0.14</td>
<td>0.14</td>
</tr>
<tr>
<td>\textit{Bcaba1-27}</td>
<td>0.38</td>
<td>0.45</td>
</tr>
<tr>
<td>\textit{Bcaba1C5}</td>
<td>3,174</td>
<td>349</td>
</tr>
<tr>
<td>Control</td>
<td>0.39</td>
<td>0.03</td>
</tr>
</tbody>
</table>

\(a\) ABA was determined in 7-day-old liquid cultures by EIA with noninoculated Sprecher medium as a control.
FIG. 6. LC-MS analyses. (A) Mass spectrum of ABA standard showing a predominant peak at $m/z$ 247.1335 ($M^+ - H_2O$). (B) Ion traces for $m/z$ 247.1335 with a width of 30 ppm of metabolites extracted from wild-type ATCC 58025 (dotted line) and replacement mutants ΔBccpr1-16 (dashed line) and ΔBcaba1-27 (full line). (C) Ion traces for $m/z$ 235.172 with a width of 30 ppm of metabolites extracted from wild-type ATCC 58025 (dotted line) and replacement mutant ΔBcaba1-27 (full line). (D) Mass spectrum of the unknown metabolite from replacement mutant ΔBcaba1-27 eluting at 4.854 min.
interrupted by four introns of 48, 49, 83, and 59 bp, respectively, with a coding capacity of 509 amino acids, yielding a calculated molecular mass of 57.2 kDa. As expected from the EST data, the derived amino acid sequence of \( \text{bcaba1} \) showed significant homology to fungal P450 monooxygenases.

\text{bcaba1} is part of the ABA biosynthetic pathway. To clarify the role of this candidate gene unequivocally, a gene disruption approach was performed, comparable to the one described for \( \text{bccpr1} \). The gene replacement fragment \( \text{pABA1Rep} \) (see Fig. 7 and Materials and Methods) was used to generate transformants of strain ATCC 58025. As described above, these transformants were checked by PCR and Southern analysis. Transforms of strain ATCC 58025. As described above, these transformants were checked by PCR and Southern analysis. Transforms \( \Delta \text{Bcaba1-2}, \Delta \text{Bcaba1-27}, \) and \( \Delta \text{Bcaba1-36} \) showed the diagnostic PCR fragments of 0.8 and 0.7 kb generated by primer pairs composed of primers 14 and 6 and of primers 7 and 15, respectively, and lacked the wild-type fragment of 0.8 kb generated by primers 16 and 15 (data not shown). Hybridization of HindIII-digested genomic DNA with the labeled \( \text{bcaba1} \) 5’ flank showed lack of the 1.5-kb wild-type fragment and the presence of the expected 3.5-kb replacement fragment (Fig. 8A). Transformant T35, on the other hand, is an ectopic transformant: it contains the wild-type fragment and an additional hybridizing fragment of 3.0 kb. Northern analyses confirmed this interpretation. In the mycelia of mutants \( \Delta \text{Bcaba1-2} \) and \( \Delta \text{Bcaba1-27} \) grown under ABA biosynthesis conditions, no \( \text{bcaba1} \) transcript could be detected (Fig. 8B). All mutants showed a normal growth rate and morphology in axenic culture. Mutants \( \Delta \text{Bcaba1-2} \) and \( \Delta \text{Bcaba1-27} \) were subjected to a detailed biochemical analysis. EIA tests showed that both contain no detectable ABA, even less than the \( \text{bccpr1} \) mutants (Table 1). LC-MS analyses confirmed these findings: both mutants lack the specific ABA peak, proving unequivocally that BcABA1 is involved in ABA biosynthesis (as shown in Fig. 6C for mutant \( \Delta \text{Bcaba1-27} \)).

As a final proof for the role of \( \text{bcaba1} \) in ABA biosynthesis, mutant \( \Delta \text{Bcaba1-2} \) was transformed with vector \( \text{pComABA1} \) containing a full-length copy of \( \text{bcaba1} \), by using a nourseothricin-resistance selection cassette (see Materials and Methods). Altogether, 12 complemented mutants were screened for the presence of the complete gene by PCR (data not shown), and the bona fide complemented mutant Bcaba1C5 was further characterized. Since it contains probably multiple copies of the gene (see Fig. 8A), expression of \( \text{bcaba1} \) in Bcaba1C5 is not only recovered but exceeds the wild-type level (see Fig. 8B). ABA production of the complemented mutant is comparable to the wild type (Table 1).

The LC-MS analyses showed that a peak at retention time 4.854 min with a major ion peak at \( m/z \) 235.172 was significantly increased in the mutants (Fig. 6C and D). The molecular mass corresponds to the postulated ABA intermediate \( \alpha-\gamma \)-ionylidene acetic acid (see Fig. 1, component 1), although unequivocal identification of this metabolite would require further detailed analyses. Taken together, these data confirm that \( \text{bcaba1} \) represents the first identified fungal ABA biosynthesis gene.
DISCUSSION

We have presented here conclusive evidence that the biosynthesis of ABA in *B. cinerea* follows—as postulated earlier on the basis of biochemical data—a direct pathway, including direct cyclization of farnesy1 diphosphate and sequential oxidation. (i) DPA, which has been shown to be a potent inhibitor of phytoene dehydrogenase, i.e., of carotenogenesis, in several fungi, such as *Phycomyces blakesleeanus* (7) and *Mucor circinelloides* (11), did not affect ABA biosynthesis in *B. cinerea* as previously described for *C. pinii-densiforme* (21). (ii) Deletion of the *bcpcr1* gene probably encoding a cytochrome P450 oxidoreductase drastically reduced ABA biosynthesis, strongly suggesting the involvement of P450 monooxygenases in this fungal biosynthetic pathway (in contrast to the plant pathway). Most fungi investigated thus far seem to contain only one CPR encoding gene. However, ∆*bcpcr1* deletion mutants still seem to exhibit some residual ABA producing activity as determined by EIA. Furthermore, mutants are viable and even able to infect bean plants (data not shown), although P450 monooxygenases are also involved in primary metabolism, e.g., the formation of ergosterol (37). This leads to the assumption that at least one more electron-donating system has to be present in *B. cinerea*. In the zygomycete fungus *Rhizopus nigricans*, two putative CPR encoding genes have been identified sharing 66% identity (16). Therefore, the presence of a second CPR in *Botrytis* cannot be excluded completely. However, it is also possible that another electron donor, as described for cytochrome b5, in *Saccharomyces cerevisiae* (33), can (at least partially) overcome the defect and thus be responsible for residual ABA production. (iii) Deletion of a P450 monooxidase gene, *bcaba1*, completely abolished ABA biosynthesis, identifying the product of this gene as essential for ABA biosynthesis. These findings also rule out a minor role of the carotenoid pathway in this strain. This clearly demonstrates that the fungal ABA pathway, at least in *B. cinerea*, is different from that in higher plants (such as the pathways of GA3 and ethylene). *bcaba1* thus represents the first fungal ABA biosynthesis gene to be identified. The availability of this gene opens fascinating new opportunities for further investigations.

(i) The role of fungal ABA production in host-pathogen interactions can now be studied in detail. Expression of the gene in planta can be easily monitored and can give a clue about the timing of ABA synthesis; more importantly, a functional analysis by targeted inactivation in a pathogenic field isolate can give unequivocal evidence for the importance of fungal ABA in the pathogenic process (deletion of *bcaba1* in the standard strain B05.10 is under way). However, in this context it has to be considered that also fungal ABA precursors such as α-(γ)-ionylideneacetac acid can exhibit biological activity (reviewed by Oritani and Kiyota [23]). Therefore, overexpression experiments would also be helpful in this context. In addition, although the *bcaba1* mutants do not show any defects in vegetative properties, an influence of ABA on other parts of the (nonpathogenic) life cycle, e.g., sexual propagation, cannot be ruled out thus far and can now be analyzed in detail.

(ii) Since genes of several secondary metabolic pathways in fungi are arranged in gene clusters, such as penicillin (9), sterigmatocystins (3), gibberellins (34), and ergot alkaloids (36), there is a high probability of identifying other genes encoding enzymes of the ABA pathway by a chromosome-walking approach. Indeed, preliminary analyses identified at least two genes downstream of *bcaba1* that are candidates for further ABA biosynthetic genes (V. Siewers and P. Tudzynski, unpublished data).

(iii) The increasing number of fungal genomes available allows comparative evolutionary studies, since ABA biosynthesis seems to have been evolved in quite diverse groups of fungi, including ascomycetes such as *Ceratocystis* spp., zygomycetes such as *Mucor* spp., and basidiomycetes such as *Agrocybe praecox* (8, 10).

(iv) Detailed expression studies will make it possible to define more accurately the conditions for ABA production in axenic culture, an important prerequisite for biotechnological evaluation. In the same way that transformation of a *G. fujikuroi* wild-type strain with additional copies of the gibberellin gene cluster led to an up to threefold-enhanced gibberellin production (P. Linnemannstons and B. Tudzynski, unpublished data), overexpression of *bcaba1* (and other genes of the pathway) could be a helpful tool for the design of better ABA production strains.

We also demonstrated here the options provided by a “genomics” approach. Although only a part of the *Botrytis* genome sequence is publicly available, a concerted approach with EST data, expression studies, and gene disruption was successful in identifying the first ABA biosynthesis gene. Conventional approaches, e.g., by differential cDNA screening, had been unsuccessful. Thus, there is also a good chance for identification of genes involved in other phytohormone pathways in *B. cinerea* in the near future, especially for ethylene, which probably has significant impact on pathogenicity and might also affect other aspects of the fungal life cycle.

ACKNOWLEDGMENTS

This study was supported by the Deutsche Forschungsgemeinschaft (Tu 50/9) and the European Community-Access to Research Infrastructure action of the Improving Human Potential Programme.

The technical support of Bayer Crop Science in establishing an EST library for strain ATCC 58025 is gratefully acknowledged. We thank Bettina Tudzynski and Peter Schreier (Bayer Crop Science) for helpful manuscript, Anke Boettig for excellent technical assistance, and E. Weiler (Bochum) for providing ABA-specific antisera and for help in establishment of the ABA-EIA.

REFERENCES


