The Analysis of Fruit Protection Mechanisms Provided by Reduced-Pathogenicity Mutants of *Colletotrichum gloeosporioides* Obtained by Restriction Enzyme Mediated Integration

N. Yakoby, D. Beno-Moualem, I. Kobiler, and D. Prusky

Department of Postharvest Science of Fresh Produce, Agricultural Research Organization, The Volcani Center, Bet Dagan 50250, Israel.

Current address of N. Yakoby: The BioTech Center, Rutgers University, Cook College, New Brunswick, NJ 08901.

Accepted for publication 17 June 2002.

ABSTRACT


*Colletotrichum gloeosporioides* is an important postharvest pathogen that attacks ripe avocado fruit. Two reduced-pathogenicity mutants, Cg-M-142 and Cg-M-1150, previously obtained by restriction enzyme mediated integration, were used for the sequential analysis of the induction of biocontrol in avocado fruit. Plant biochemical indicators, such as H+-ATPase activity and levels of reactive oxygen species, phenylalanine ammonia lyase, epicatechin, and an antifungal diene, were investigated. The main difference between Cg-M-142 and Cg-M-1150 was the lack of appressorium formation by the latter. Preinoculation of avocado fruit with Cg-M-142 enhanced H+-ATPase activity and the production of reactive oxygen species. These early signaling events were followed by higher phenylalanine ammonia lyase activity and higher levels of epicatechin and the antifungal diene, and decay was delayed. Unlike Cg-M-142, Cg-M-1150 did not activate early signaling events related to fruit resistance. We suggest that the initiation of early signaling events affecting fruit resistance is determined by the capability of the pathogen to interact with the fruit during appressorium formation. Furthermore, the intensity of the fruit defense response determines the level of resistance during fruit storage.

Additional keywords: fruit signal transduction, pathogen–fruit interaction, postharvest decay.

The phytopathogenic fungus *Colletotrichum gloeosporioides* (Penz.) Penz. & Sacc. (teleomorph Glomerella cingulata) is an important postharvest pathogen attacking a wide range of tropical and subtropical fruits (8,25). *C. gloeosporioides* initiates its attack by conidium germination and appressorium formation (5), followed by the production of infection hyphae, which penetrate the epidermal cells of the avocado exocarp. The hyphae remain quiescent until the fruit ripens and softens during storage (19). It has been hypothesized (19) that several mechanisms are responsible for the resistance of unripe fruit to fungal attack during quiescence: (i) the lack of nutrient supply for the pathogen in unripe fruit, (ii) the lack of secretion of fungal pathogenicity factors, (iii) the presence of preformed antifungal compounds, and (iv) the presence of inducible antifungal compounds. The last two mechanisms have been suggested as the main factors affecting avocado resistance (19).

The level of 1-acetoxy-2-hydroxy-4-oxo-heneicosa-12,15-diene (antifungal diene) depends on the synthesis and metabolism of the compound (24). The synthesis of diene includes the activation of Δ9 desaturase (L. Madi, W. Xuejun, I. Kobiler, A. Lichter, and D. Prusky, unpublished data), Δ12 desaturase (W. Xuejun et al., unpublished data), and a fatty acid elongase (Xuejun et al., unpublished data). Transcriptional activation of the two genes has been observed in avocado as a result of inoculation with *C. gloeosporioides* (Xuejun et al., unpublished data). The metabolism of the antifungal diene has been attributed to the oxidative activity of lipoxygenase (23). The activity of lipoxygenase is modulated by the level of flavan-3-ol-epicatechin (epicatechin) (12), a natural antioxidant phenol generated in the phenylpropanoid pathway and present in the avocado exocarp. The level of epicatechin declines during fruit ripening, and as a result lipoxygenase can reduce diene levels (24). Transcriptional activation of genes encoding epicatechin-forming proteins in the phenylpropanoid pathway, such as phenylalanine ammonia lyase (PAL), has been shown to increase diene and concomitant fruit resistance (2,4,27). It is still unclear what early events activate the mechanism of resistance in avocado fruit. It was recently reported that the production of reactive oxygen species (ROS) in avocado significantly increased after inoculation with *C. gloeosporioides* conidia (4). It has been suggested that quiescent infection may be a local source of ROS production. This increase in ROS activates the phenylpropanoid pathway, which subsequently increases the level of antifungal diene and enhances fungal inhibition.

Recently, we generated reduced-pathogenicity mutants of *C. gloeosporioides* on avocado fruit, by insertional mutagenesis using restriction enzyme mediated integration (REMI) technology (36). Isolate Cg-M-142 produced appressoria on the exocarp, like the wild-type isolate Cg-14, but it caused late and minor symptom development in ripening fruit. Isolate Cg-M-1150 did not produce appressoria on the exocarp, nor did it cause any symptoms. Preinoculation of avocado fruit with Cg-M-142 delayed the development of symptoms caused by the wild-type isolate and enhanced diene levels (36). Preinoculation with Cg-M-1150 did not affect the level of diene, nor did it delay the appearance of decay symptoms caused by Cg-14. The present work describes the interaction of the reduced-pathogenicity mutant and avocado, with appressorium formation as a main factor in early events (4) leading to the activation of a mechanism of resistance to *Colletotrichum*. We propose a model to explain the recognition of the fungus by avocado fruit and the importance of appressorium formation as a factor in host-plant interactions.
MATERIALS AND METHODS

Avocado fruit, fungal isolate, and growth media. Experiments were carried out with freshy harvested fruit of the avocado cultivar Fuerte (Persea americana Mill. var. drymifolia (Schltdl. & Cham.) S. F. Blake) from an orchard at Kibbutz Givat Brenner, Israel. A single-conidial isolate of C. gloeosporioides, Cg-14, was obtained from a decayed fruit of the avocado cultivar Fuerte, and conidia were maintained in 10 mM Na-phosphate buffer (pH 7.2), 40% glycerol, at –80°C. Three-week-old conidia were harvested from Mathur’s medium (M,S) plates (3), counted, with the aid of a hemacytometer (Brand, Wertheim, Germany), and used for culture and fruit inoculation. Tissue treatment and inoculation were performed as previously described (4). Experiments were conducted in two consecutive harvesting seasons and repeated at least three times per season. The data presented are usually the average values of three to five replications of a single experiment. In some cases the average of all the replications of repeated experiments are presented with their pooled standard error.

Fruit protection assays and phenotypic characterization of transformed isolates. Twenty freshly harvested avocado fruits (cv. Fuerte) were dipped for 30 s in suspensions of conidia (5 × 10⁶ conidia/ml) of the two C. gloeosporioides mutants (Cg-M-142 and Cg-M-1150) and then stored at 90% relative humidity and 20°C for 24 h. Thereafter, the treated fruit was spot-inoculated with a Cg-14 conidial suspension (1 × 10⁵ conidia/ml) (36). Appressoria were observed under a light microscope (BH-2 Olympus, Tokyo, Japan) in a 0.5-mm-deep slice of inoculated exocarp cut out with a scalpel 24 and 48 h after inoculation. Isolate growth rates were measured in five replicates on exocarp and on glass slides by inoculation with 10 µl of the conidial suspension described above. All experiments were repeated at least three times over two consecutive avocado seasons. Exocarp inoculation was carried out by brushing exocarp strips with a conidial suspension (2 × 10⁶ conidia/ml) (22). As a control, exocarp strips were brushed with distilled water only.

Extraction of epicatechin. Epicatechin was extracted from 3 g of 1-mm-thick avocado exocarp strips, as previously described (21). Epicatechin was quantified in a reverse-phase high-performance liquid chromatography (HPLC) (RP-18) column eluted with 55% methanol and 1% acetic acid in double-distilled water. Calculation of the epicatechin level was based on comparison of the HPLC peak areas quantified at 275 nm with those of the standard, (2R,3S)-2-(3,4-dihydroxyphenyl)-3,4-dihydro-1(2H)-benzopyran-3,5,7-triol (Sigma, Jerusalem, Israel).

Extraction of PAL and analysis of PAL activity. Samples for analysis of PAL activity were extracted by blending 5 g of avocado exocarp in 50 ml of cold acetone (-20°C) and collecting the homogenate on Whatman no. 1 filter paper. This process was carried out four times. The exocarp powder was dried at room temperature overnight and stored at –20°C until used. PAL was extracted from 100 mg of exocarp powder stirred for 2 h at 4°C with 5 ml of 0.1 M sodium borate buffer, pH 8.8. The extract was centrifuged at 10,000 × g for 20 min at 4°C, and the supernatant was used as an enzyme source. The reaction mixture for PAL consisted of 10 mM L-phenylalanine (Aldrich, Jerusalem, Israel), 0.1 M borate buffer (pH 8.8), and 0.75 ml of enzyme extract in a final volume of 2.5 ml. The reaction was incubated at 37°C for 60 min and terminated by the addition of 0.1 ml of 5 N HCl. The product, cinnamic acid, was extracted in 3.5 ml of diethyl ether and quantified by comparing the amount of trans-cinnamic acid formed at 269 nm (15).

Preparation of microsomes and isolation of plasma membranes. Plasma membranes were isolated from 40 g of exocarp strips after treatment with 0.2 mM CaSO₄ for 24 h. Plasma membranes were isolated at several periods after inoculation with C. gloeosporioides. Tissue was homogenized in a Waring blender at 4°C, and microsomes were prepared as previously described (10). The microsomes were suspended in a suspension medium (330 mM sucrose, 5 mM KCl, and 5 mM potassium phosphate buffer, pH 7.8), and the plasma membranes were separated by two-phase partitioning (13). The final composition of the phase system was 6.4% (wt/wt) dextran T500 and 6.4% (wt/wt) polyethylene glycol (PEG 3350) in the suspension medium. The final upper phase was diluted 1:10 with a resuspension medium (330 mM sucrose, 1 mM dithiothreitol, 5 mM NaF, and 5 mM MES-bis-tris propane, pH 7.8), and the purified plasma membranes were pelleted by centrifugation at 55,000 × g. The pellet was resuspended in 330 mM sucrose, 5 mM MES-bis-tris propane, and 1 mM dithiothreitol adjusted to pH 7.0 and brought to a concentration of 1 to 3 mg protein/ml. Protein was assayed according to the Bradford method (6).

Reactive oxygen production measured by cytochrome c reducing activity. The production of ROS was determined by cytochrome c reduction dependent NADPH oxidation, as previously described (18). Reduction of the cytochrome c molecule produces one molecule of reactive oxygen (16). The assay consisted of 20 µM cytochrome c (Sigma type VI), 100 µM NADPH, 25 mM potassium phosphate buffer (pH 7.5), and 10 µg of plasma membrane protein in a total volume of 1 ml. The reaction was initiated with the addition of NADPH, and the rate of cytochrome c reduction was measured on a dual-wavelength spectrophotometer at 550 nm, by the difference in the absence and presence of 100 µg of superoxide dismutase, as described by Owen and Hancock (18). H⁺-ATPase activity and rubidium uptake. Vanadate-sensitive plasma membrane was determined in the presence of 0.01% polyethylene glycol 20 cetyl ether (Brij 58) and in the presence or absence of 0.2 mM Na₂VO₄. H⁺-ATPase activity was assayed as previously described (11), with variations: 4 µg of plasma membrane protein was incubated for 30 min at 35°C in 0.5 ml of a medium composed of 40 mM MES-tris (pH 6.5), 0.25 M sucrose, 100 mM KCl, 3 mM MgSO₄, and 3 mM ATP. H⁺-ATPase activity in the sample plasma membrane was found to be 87% vanadate sensitive.

For rubidium uptake, exocarp tissue disks (about 5 mm in diameter and 1 mm thick, almost free of mesocarp) were prepared as described for microsome preparation. The disks were preincubated for 30 min in 0.2 mM CaSO₄ at 25°C. [¹⁸Rb]K⁺ was added to the vial for the last 20 min of the reaction. The uptake of [¹⁸Rb]K⁺ into 70 mg of avocado disk tissue (from five disks) was assayed in 5 ml of 0.2 mM CaSO₄ and 0.1 mM [¹⁸Rb]K⁺ (700 Bq/µmol). Exchangeable was removed with three washings over a 5-min period in 15 ml of ice-cold 10 mM CaSO₄. The disks were transferred to vials for scintillation counting; 0.2 ml of 1 N NaOH was added for discoloration, and then 4 ml of Opti-Fluor O (Packard, Groningen, The Netherlands) was added. Samples were counted in an MR 300 Lumtron Automatic liquid scintillation system (Packard).

Extraction and quantification of the antifungal diene. A 10-g sample of avocado exocarp (1–2 mm thick) was homogenized in 95% ethanol in an Omni-Mixer (Sorvall, DuPont Company, Newtown, CT) at full speed for 3 min. The ethanol extract was dried in a rotary evaporator at 40°C and then redissolved in 10 ml of distilled water, and the organic phase was extracted by fractionation with dichloromethane. After two extractions, the organic phases were pooled, dried with anhydrous MgSO₄ (Riedel-deHaen, Seelze, Germany), and evaporated to dryness. Samples were redissolved in 1 ml of ethanol AR (Bio Lab, Jerusalem, Israel) and analyzed by HPLC (26). The average value of three separate extractions is presented. The experiment was repeated twice in each of two consecutive avocado seasons.

RESULTS

Phenotypic analysis of REMI mutants. Two C. gloeosporioides REMI mutant isolates, Cg-M-142 and Cg-M-1150, were compared to the wild-type (WT) isolate Cg-14 for conidial germination, germ tube elongation, and appressorium formation on glass
slides (Fig. 1A) and on avocado exocarp (Fig. 1B) 24 and 48 h after inoculation. Cg-M-142 and Cg-M-1150 were similar to the wild type in germination and germ tube elongation on the exocarp, but the two mutants were retarded on glass slides. Appressorium formation was delayed in Cg-M-142 and absent in Cg-M-1150 on both glass slides and exocarp. Appressorium formation by Cg-M-142 was similar to that of the WT within 48 h after inoculation, while Cg-M-1150 did not produce appressoria at all (Fig. 1A and B), even after 96 h (data not presented). Analysis of germ tube elongation showed that Cg-M-142 and Cg-M-1150 formed longer germ tubes on inoculated fruit than on glass slides (Fig. 1A and B).

Effect of preinoculation with REMI mutants on the development of decay. Inoculation of avocado fruit with Cg-M-142 increased the level of the antifungal diene approximately threefold within 4 days (Fig. 2). The diene level remained higher in treated fruit even 11 days after inoculation with Cg-M-142. Decay symptoms were delayed by 2 days in fruit preinoculated with Cg-M-142, compared to the control fruit and fruit preinoculated with Cg-M-1150. Diene levels were similar in fruit inoculated with Cg-M-1150 and the control fruit, during the monitored period. In addition, the development of symptoms in fruit preinoculated with Cg-M-1150 was similar to that in control fruit (Fig. 2).

H⁺-ATPase activity, rubidium uptake, and O₂ generation-dependent NADPH oxidation. Exocarp slices inoculated with the mutant strains were tested for early signaling affects. H⁺-ATPase activity in the plasma membrane extracted from exocarp inoculated with Cg-M-142 increased fourfold within 2 h after inoculation (Fig. 3A) and 2 h later dropped to the same level as the control. Inoculation with Cg-M-1150, however, had no affect on H⁺-ATPase activity 2 h after treatment, with a small increase 4 h after treatment. This observation was further confirmed by monitoring Rb⁺ uptake, which showed a pattern similar to that of H⁺-
ATPase activity, while Cg-M-1150 followed the same pattern as the control (Fig. 3B). A transient sixfold increase in NADPH oxidase activity of exocarp inoculated with Cg-M-142 occurred 1 to 2 h after inoculation (Fig. 4). In exocarp inoculated with Cg-M-1150, cytochrome c reduction was unaffected 2 h after inoculation and a increased slightly 4 h after inoculation (Fig. 4).

**Induction of PAL and epicatechin level.** Inoculation of exocarp slices with Cg-M-142 increased PAL activity more than twofold within 10 h (Fig. 5A), whereas PAL activity was reduced in exocarp inoculated with Cg-M-1150, compared to the control. Epicatechin also increased twofold in exocarp inoculated with Cg-M-142, whereas no change in epicatechin levels was found after inoculation with Cg-M-1150 (Fig. 5B). Epicatechin levels increased when whole avocado fruit was preinoculated with Cg-M-142 or Cg-M-1150. However, only the epicatechin increase in fruit inoculated with Cg-M-142 was significant (a twofold increase within 120 h after inoculation) (Fig. 6), while no significant increase was found in fruit inoculated with Cg-M-1150.

**DISCUSSION**

Development of disease caused by pathogens in the genus *Colletotrichum* requires adhesion of conidia to the surface of host tissue, followed by conidial germination and the formation of appressoria to penetrate the host tissue (3,19). Two reduced-pathogenicity REMI mutants, Cg-M-142 and Cg-M-1150 (36), did not differ from the wild-type *C. gloeosporioides* in germination rate or germ tube elongation on fruit exocarp. The most striking differences between the two mutants and the wild type were (i) the lack of appressorium formation by Cg-M-1150 and the delayed appressorium formation by Cg-M-142 and (ii) the capability of Cg-M-142 to induce resistance to the wild-type *C. gloeosporioides* isolate. The importance of appressoria in fungal attack has been extensively studied (30,37); however, it has not been examined for its importance in triggering signals leading to increased resistance.

Avocado fruit exocarp, like other plant organs (7,9), recognizes penetrating hyphae of wild-type *C. gloeosporioides* appressoria at the level of the host plasma membrane (4). This recognition occurs by the release of soluble cell wall sugars, probably sensed by a heterotrimeric G-protein receptor that activates a G-protein (32, 33) that transduces the signal by activating one or more membrane-bound phosphatases, resulting in the dephosphorylation of the host plasma membrane H^+-ATPase and activation of NADPH oxidase (4) (Fig. 7). We cannot exclude the presence of other receptors in the plasma membrane that may sense the penetrating fungus as well. However, activation of H^+-ATPase and H_2O_2 production was stimulated by the appressorium-forming isolate Cg-M-142 and not by the non-appressorium-forming isolate Cg-M-1150. The latter caused a marginal delayed increase in both signals, had no effect on rubidium uptake, and failed to alter fruit resistance. The lack of induction of resistance indicates the importance of appressorium formation in initiating defense signals, such as the proton electrochemical gradient across the plasma membrane (28). In avocado fruit, H^+-ATPase was induced by the penetrating appressoria, along with Rb^+ uptake, confirming the opening of K^+ channels as an early reaction to fungal invasion. ROS production has been shown to be an early response in plant pathogen interactions in general (1,17) and in avocado fruit (4). Inoculation of avocado exocarp tissue with the wild-type Cg-14 increased ROS production threefold within 2 h, while inoculation with Cg-M-142 increased ROS production sixfold, which can explain its increased resistance effect. Cg-M-1150 did not significantly affect ROS levels. In addition, the application of Ca^{2+} ionophore (A-23187), which opens Ca^{2+} channels, stimulated the production of H_2O_2 (4) (Fig. 7). Thus the level of H_2O_2 in the fruit may predict the level of fruit resistance against fungal attack. In addition to these responses, Cg-M-142 activated the resistance-related phenylpropanoid pathway, where PAL activity was higher and epicatechin accumulation was 30% higher than that induced by the wild-type Cg-14 (4). Moreover, in vivo defense experiments on whole fruit showed a threefold increase in epicatechin production in fruit inoculated with Cg-M-142, compared to the control. Activation of the phenylpropanoid pathway increases the production of epicatechin, an antioxidant, which protects the reduction of the antifungal diene by inhibiting lipoxygenase (2). Epicatechin has also been found to inhibit *C. gloeosporioides* polygalacturonase and pectate lyase (34,35) (Fig. 7), and the latter was recently found to be an important virulence factor in fruit-fungus interactions (35). The induction of PAL activity by H_2O_2 in avocado fruit is in agreement with the results of studies of *Arabidopsis* and tobacco cell cultures, in which the addition of exogenous H_2O_2 triggered PAL expression (14,17). This suggests that PAL activity is stimulated after direct penetration by appressoria, as was observed in barley leaves after inoculation with *Erysiphe graminis*.

---

**Figure 5.** Phenylalanine ammonia lyase activity A, and epicatechin B, in slices of avocado fruit exocarp inoculated with *Colletotrichum gloeosporioides* mutant isolates Cg-M-142 (■) and Cg-M-1150 (○) or with water (▲). The bars represent standard error in one representative experiment run four times with three replications. The experiment was repeated in two consecutive avocado seasons.

**Figure 6.** Epicatechin in avocado fruit exocarp inoculated with *Colletotrichum gloeosporioides* mutant isolates Cg-M-142 (■) and Cg-M-1150 (○) or with water (▲). The bars represent standard error in one representative experiment run four times with three replications. The experiment was repeated in two consecutive avocado seasons.
or *E. pisi* (29), via the production of H$_2$O$_2$. Although Cg-M-1150 showed a marginal increase in early signals as well as in PAL activity, it did not have any effect on fruit resistance. It is possible that compounds secreted by the fungus during its growth on the exocarp generate this effect. Since PAL is the first enzyme in the phenylpropanoid pathway that branches into various routes, the increase in PAL may affect other downstream products unrelated to resistance.

The genetic differences between the wild-type Cg-14 and its derived isolate Cg-M-142 in their ability to induce resistance has not yet been elucidated. Several genes have been reported to affect appressorium formation (37). A nonpathogenic *C. lagenarium* mutant in mitogen-activated protein (MAP) kinase, CMK1 gene, showed reduced conidiation and failed to form appressoria (30). The disruption of *Botrytis cinerea* BMP1, a gene encoding MAP-kinase, prevented appressorium formation and pathogenicity (38). Yang and Dickman (37) showed that strains of *C. trifolii* deficient in protein-kinase A were unable to infect intact alfalfa plants, likely because of a failure to penetrate. Cg-M-142 has been shown to harbor a single REMI integration site, which suggests a single gene disruption, whereas two independent integration sites have been found in Cg-M-1150, possibly disrupting one or two genes to cause its unique phenotype (36). The genes disrupted by REMI technology in both mutants are currently under investigation. Although the flanking regions have been cloned, we did not find any homology in the known published genetic data.

The two REMI mutant isolates allowed us to demonstrate the importance of appressorium formation dynamics in triggering fruit resistance. In general appearance, the fruit inoculated with Cg-M-142 was less decayed than fruit inoculated with Cg-M-1150 or with the control (data not shown). This observation indicates that early signaling by Cg-M-142 had succeeded in elevating resistance in fruit naturally infected with *C. gloeosporioides* as well. We suggest that the induction of fruit resistance is a result of greater epicatechin accumulation, derived from early signals generated by delayed appressorium formation and invading Cg-M-142 hyphae, in contrast to the fast penetration by the wild-type isolate (35,36). This fast penetration by the wild-type isolate has generated a fast and transient activation of resistance mechanism, but this effect did not delay symptom development in ripening fruit (20). Lack of appressorium formation by the Cg-M-1150 mutant is assumed to be the reason for the lack of induction of resistance and failure to colonize the fruit.

**ACKNOWLEDGMENTS**

We thank R. Ardi for drawing the initial general scheme model. This research was supported by Research Grants IS-2686-96 to D. Prusky from BARD, the U.S. Binational Agricultural Research and Development Fund, the German–Israel Agricultural Research Agreement (GIARA) for the Benefit of the Third World, and the U.S.–Israel Cooperative Development Research (CDR) Program. The first author, N. Yakoby, was granted personal scholarships from the Israel Fruit Marketing Board, Ministry of Agriculture and Rural Development, and from the Association of Trustees and Friends of the Agricultural Research Organization.

**LITERATURE CITED**


