Differential Interference with *Pythium ultimum* Sporangial Activation and Germination by *Enterobacter cloacae* in the Corn and Cucumber Spermospheres

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Differential protection of plants by *Enterobacter cloacae* was studied by investigating early sensing and response behavior of *Pythium ultimum* sporangia toward seeds in the presence or absence of *E. cloacae*. Ten percent of *P. ultimum* sporangia were activated within the first 30 min of exposure to cucumber seeds. In contrast, 44% of the sporangia were activated as early as 15 min after exposure to corn seeds with over 80% sporangial activation by 30 min. Germ tubes emerged from sporangia after 2.5 and 1.0 h in the cucumber and corn spermospheres, respectively. Seed application of the wild-type strain of *E. cloacae* (EcCT-501R3) reduced sporangial activation by 45% in the cucumber spermosphere, whereas no reduction was observed in the corn spermosphere. Fatty acid transport and degradation mutants of *E. cloacae* (strains EcL1 and Ec31, respectively) did not reduce sporangial activation in either of the spermospheres. Although wild-type or mutant strains of *E. cloacae* failed to reduce seed colonization incidence, pathogen biomass on cucumber seeds was reduced in the presence of *E. cloacae* strains EcCT-501R3 and Ec31 by 4 and 8 h after sowing, respectively. By 12 h, levels of *P. ultimum* on cucumber seeds treated with *E. cloacae* EcCT-501R3 did not differ from levels on noninoculated seeds. On corn seeds, *P. ultimum* biomass was not affected by the presence of any *E. cloacae* strain. When introduced after sporangial activation had occurred, *E. cloacae* failed to reduce *P. ultimum* biomass on cucumber seeds compared with that on nontreated seeds. Also, increasing numbers of sporangia used to inoculate seeds yielded increased pathogen biomass at each sampling time. This indicates a direct link between the level of seed-colonizing biomass of *P. ultimum* and the number of activated and germinated sporangia in the spermosphere, suggesting that *E. cloacae* suppresses *P. ultimum* seed infections by reducing sporangial activation and germination within the first 30 to 90 min after sowing.

*Pythium ultimum* is a widespread and destructive oomycete pathogen of plants, notorious for its rapid pathogenic development in response to germinating seeds. *P. ultimum* infects seeds and seedlings, leading to a disease known as damping-off (10). Disease development is generally initiated from sporangia, which together with oospores serve as quiescent structures in the life cycle, each requiring seed exudates to break dormancy and initiate plant infections (9). Sporangia can transition from quiescence to active growth in as little as 1.5 h after exposure to a seed (7, 19). Seed exudates from many plant species can induce these rapid responses (4, 21). Although the specific exudate elicitors from most plants are unknown, the elicitors from cottonseed exudates have been identified as long-chain unsaturated fatty acids (LCUFA), comprised largely of oleic and linoleic acids (9, 10, 18). Other exudate molecules such as saturated fatty acids, sugars, amino acids, and other organic acids do not elicit sporangial germination (13, 14, 18). Once sporangia have germinated, high levels of seed colonization and subsequent infection are observed within 24 to 48 h, leading to rapid seed and seedling death (1, 2, 8, 9, 24). However, if exudate stimulants are removed by physical or biological means, sporangia do not germinate and little or no seed colonization and disease development occur (3, 7, 12, 15).

*Enterobacter cloacae* is a common seed-associated bacterium that suppresses seed infections, protecting a number of plant species from *P. ultimum*-induced damping-off (3, 4, 8, 12). Plant protection by *E. cloacae* is achieved largely by degradation of LCUFA in seed exudates (20), which eliminates the stimulation of *P. ultimum* sporangia (4, 21). *E. cloacae* suppresses *P. ultimum* infections when applied as a coating onto seeds of plants such as carrot, cotton, cucumber, lettuce, radish, sunflower, tomato, and wheat. However, no disease suppression is observed when *E. cloacae* is applied as a coating onto seeds of corn or pea (4, 12).

Currently, it is not known how quickly sporangial germination and subsequent disease development are initiated on these different host species. Furthermore, it is not known whether reduced disease development that accompanies the removal of exudate stimulants is due solely to reductions in sporangial germination or whether *E. cloacae* may also suppress later stages of pathogenesis. Because most of the current knowledge of seed-induced stimulation of sporangial germination comes from work with in vitro-collected seed exudates (4, 18, 20, 21), little is known of the temporal dynamics of sporangial responses in the spermosphere and how these responses relate to the temporal exudation of stimulatory molecules.

To better understand the differential protection of plants by
E. cloacae, this study was focused on the dynamics of P. ultimum during early stages of pathogenesis in the presence and absence of E. cloacae on two differentially protected hosts. The main goals of this work are to identify how quickly E. cloacae reduces P. ultimum sporangial activation and germ tube emergence in the spermosphere and to establish whether this relates directly to reductions in subsequent P. ultimum seed colonization. A further goal is to better understand the potential role of E. cloacae fatty acid degradation during early stages of disease development.

MATERIALS AND METHODS

Plant material. To reduce experimental variability, corn (Zea mays cv. Northern X-tra Sweet) and cucumber (Cucumis sativus cv. Marketmore) seeds were sorted by discarding cracked, deformed, or discolored seeds. Seeds were then surface sterilized for 3 min in 0.05% sodium hypochlorite with 1 to 2 drops of Tween 20, rinsed with sterile deionized water (sdw), and blotted dry. Seeds weighing between 160 and 200 mg and between 24 and 30 mg were used for all corn and cucumber assays, respectively.

Production and germination of Pythium ultimum sporangia. Pythium ultimum P4 was routinely grown on a mineral salts medium containing 0.1% α-phosphatidyldcholine (SM+L). Sporangia produced on this medium mimic the behavior of sporangia produced on live plant tissue (13). Agar discs (4-mm diameter) were cut from 5-day-old cultures kept at 27°C. Discs were leached twice in darkness for 10 minutes each followed by a 3-h leaching by adding leaching buffer [0.01 M Ca(NO3)2·4H2O, 0.04 M MgSO4·7H2O, 0.05 M KNO3, pH 5.8] and replacing it with fresh buffer at the end of each leaching period. Discs were rinsed twice with sdw and kept at 24°C for 2 days in darkness. In preparation of defined levels of sporangia to be used as inocula, entire SM+L plates (9-cm diameter) were leached as described above. Sporangia were harvested by adding 500 µl sdw per plate and scraping sporangia from the surface of the culture. Sporangia were filtered though sterile cheesecloth, centrifuged, and washed twice with sdw. Concentrations of sporangia were adjusted with a hemacytometer. Five microliters of sporangial suspension was then added to the surface of water agar discs (4-mm diameter) and allowed to be absorbed. Sporangial germination rates were determined by staining sporangia with 0.03% acid fuchsin in 85% lactic acid and viewing them microscopically at ×100 to ×250.

Production of E. cloacae cells. E. cloacae strains used in this study were the wild type, EcCT-501R3, and two fatty acid metabolic mutants, EcL1 (fadBA type, EcCT-501R3, and two fatty acid metabolic mutants, EcL1 (3 h and at hourly intervals up to 6 h after exposure to cucumber seeds or at 4 h after exposure to corn seeds. After retrieval from the glass bead matrix, sporangial discs were further incubated so that the total incubation times (activation times in the spermosphere plus subsequent incubation for germ tube emergence) were 6 and 4 h for sporangia exposed to cucumber and corn, respectively. Sporangial discs in wells containing substrate and water but no seeds served as negative controls. Five replicates were prepared for each treatment, and each experiment was repeated three times. At each sampling time, seeds were removed and blotted dry and the seed weight was recorded.

Activation and germ tube emergence assays were also performed with E. cloacae EcCT-501R3 cells added at 105, 106, or 107 CFU/cm2 substrate. A second set of experiments evaluated E. cloacae strain EcCT-501R3, EcL1, or EcL3 at 106 CFU/cm2 substrate. Sporangial discs in wells containing no seeds but with E. cloacae EcCT-501R3 added at a rate of 106 CFU/cm2 substrate served as negative controls. Positive controls consisted of sporangial discs exposed to seeds, with E. cloacae EcCT-501R3 cells added after disc harvest to match the approximate number of cells from the maximum concentration used in the study. This permitted a check on the level of sporangial activation but also demonstrated that E. cloacae does not significantly reduce germ tube emergence during the inoculation period. Three replicates were prepared for each treatment, and each experiment was repeated three times. Seed colonization incidence. Seeds were placed in Corning 12-well tissue culture plates as described above. E. cloacae strains EcCT-501R3, EcL1, and EcL3 were added at 105 CFU/cm2 substrate. Seeds in wells containing no sporangia but with E. cloacae EcCT-501R3 added at a rate of 106 CFU/cm2 substrate served as negative controls. Seeds inoculated with P. ultimum sporangia served as positive controls. Seeds were removed at 0, 1, 2, 4, 6, and 8 h after sowing; rinsed; and then incubated on a medium containing 2% water agar for 3 days in darkness. Sporangia in wells containing substrate and water but no seeds served as negative controls. Five replicates were prepared for each treatment, and each experiment was repeated three times.

Quantification of P. ultimum seed colonization by qPCR. Quantitative PCR (qPCR) was used to quantitatively assess seed colonization by determining the amount of P. ultimum biomass on seed surfaces. To equate DNA levels with mycelial biomass equivalents, P. ultimum was grown for 6 days at 27°C on potato dextrose agar (Difeo, United States) plates overlaid with cellophane. Mycelium was harvested, weighed, frozen, and lyophilized. After dry weights were determined, lyophilized mycelium was stored at −20°C prior to DNA extraction. DNA was extracted using the Qiagen DNeasy plant minikit (Qiagen Sciences, Maryland) according to the manufacturer’s instructions for optimized yield. DNA was quantified using the Quant-iT PicoGreen double-stranded DNA assay kit (Molecular Probes, Oregon). Since times existing amounts of P. ultimum mycelia were linearly correlated with increasing amounts of DNA {r = 0.920, P < 0.0001, DNA (ng) = [26.72 × mycelium fresh weight (mg)] + 57.865}, DNA levels were subsequently used as a biomass estimator.

To prepare standards for qPCR analysis, P. ultimum was grown as described above. Mycelial mass was harvested, ground in liquid nitrogen, and transferred to 4 ml extraction buffer (0.05 M EDTA, 0.1 M Tris, 0.5 M NaCl, 0.75% β-mercaptoethanol, 0.12% sodium dodecyl sulfate, pH 7.2) and then resuspended in Tris-buffered saline. Cell density was adjusted to 106 CFU/ml at an optical density at 600 nm (0.56) to make a stock cell suspension. The stock was diluted to make suspensions that were linearly correlated with increasing amounts of DNA {r2 = 0.920, P < 0.0001, DNA (ng) = [26.72 × mycelium fresh weight (mg)] + 57.865}, DNA levels were subsequently used as a biomass estimator.

To prepare standards for qPCR analysis, P. ultimum was grown as described above. Mycelial mass was harvested, ground in liquid nitrogen, and transferred to 4 ml extraction buffer (0.05 M EDTA, 0.1 M Tris, 0.5 M NaCl, 0.75% β-mercaptoethanol, 0.12% sodium dodecyl sulfate, pH 7.2) and then resuspended in Tris-buffered saline. Cell density was adjusted to 106 CFU/ml at an optical density at 600 nm (0.56) to make a stock cell suspension. The stock was diluted to make suspensions that were linearly correlated with increasing amounts of DNA {r2 = 0.920, P < 0.0001, DNA (ng) = [26.72 × mycelium fresh weight (mg)] + 57.865}, DNA levels were subsequently used as a biomass estimator.

To determine seed-colonizing biomass of P. ultimum, seeds were prepared in glass beads and Corning 12-well tissue culture plates as described above. Treatment mixtures consisted of inoculated seeds with either E. cloacae strain EcCT-501R3 or strain EcL1 at 105 CFU/cm2 substrate. A second set of experiments evaluated E. cloacae strain EcCT-501R3 added at the time of sowing for the first experiment and at 2 h after sowing for the second experiment. Nontreated, inoculated seeds served as positive controls, and noninoculated seeds were used as negative controls for all experiments. P. ultimum biomass was measured on both corn and cucumber seeds in the first experiment but only on cucumber in the second experiment. In a third series of seed colonization experiments, cucumber seeds were inoculated with 0, 100, 300, or 600 sporangia per seed (sporangia were obtained as described above). Seeds were removed at 4, 8, and 12 h after sowing; rinsed; cut into six to eight pieces, transferred to cryo-vials; and frozen in liquid nitrogen. Seeds were kept at −80°C until extracted. Seed pieces were ground in a mortar and pestle and with glass beads in liquid nitrogen. DNA was then extracted using the Qiagen DNeasy plant minikit according to the manufacturer’s instructions for optimized yield. DNA samples were kept at
The amount of \( P. \text{ultimum} \) DNA on and in seeds was measured in triplicate in 96-well plates (Phenix, North Carolina) using qPCR. qPCR mixtures were prepared by mixing 7 \( \mu \text{L} \) water, 0.5 \( \mu \text{L} \) primer AFP276 (5'-TGATGAGA CGCTGCATT-3') (5), 0.5 \( \mu \text{L} \) 10 \( \mu \text{M} \) primer ITS4 (5'-TCTCGCGCTATTGGA TATGC-3') (22), 10 \( \mu \text{L} \) Platinum Sybr green qPCR SuperMix-uracil DNA glycosylase with 6-carboxyl-X-rhodamine (Invitrogen, California), and 2 \( \mu \text{L} \) template. Samples were amplified using the ABI Prism 7000 sequence detection system (Applied Biosystems, California) with the following conditions: initial denaturation at 95°C for 3 min followed by 40 cycles of denaturation at 95°C for 30 s, annealing at 56°C for 30 s, extension at 72°C for 60 s, and a final extension at 72°C for 10 min. A dissociation protocol starting at 60°C was used to assess the presence of primer-dimer formation and other artifacts. Quantification was achieved by comparing cycle threshold (CT) values in samples to \( C_T \) values of exogenous \( P. \text{ultimum} \) DNA standards ranging from 1.5 fg to 75 ng. Standards were established for every 96-well plate analyzed, and the relationship between \( C_T \) and DNA amount was used to calculate the amount of \( P. \text{ultimum} \) DNA in seed samples. All regression coefficients were \( >0.98 \). Slopes from the standard curves were used to calculate the efficiency (\( E \)) of the PCR using the formula \( E = -10^{\left(-1/\text{slope}\right)} \). The average efficiency was 106\% (\( \pm 2\% \)) for all tested models.

**Statistical analysis.** The PROC GLM procedure in SAS v9.1 (SAS Institute, Cary, NC) was used for statistical analyses. All assays were analyzed individually and thereafter pooled and reanalyzed. Data sets for inoculated corn and cucumber seeds were also pooled and analyzed for comparison. Seed imbibition, sporangial activation, germ tube emergence, \( P. \text{ultimum} \) seed colonization incidence, and \( P. \text{ultimum} \) seed colonization biomass data were analyzed using two-way analysis of variance with “water uptake,” “activation percent,” “germination percent,” “colonization percent,” and “amount DNA” as response variables, respectively. “Time” and “treatment” were used as continuous and categorical predictors, respectively. Additionally, simple linear regression (SLR) was performed individually on all response variables with the exception of the “water uptake” variable (pooled data sets) and treatments using “time” as a continuous variable. This was followed by multiple regressions with “treatment” as an additional predictor variable and analysis of covariance (ANCOVA) for comparison of regression slopes. SLR, multiple regression, and ANCOVA for activation and germination data were performed for data within the linear response so that time points with no activation/germination and full activation/germination were omitted. The relationship between DNA level and \( P. \text{ultimum} \) mycelial weight was analyzed using SLR with “amount DNA” as the response variable and “mycelial weight” as the predictor. qPCR standard calibrations of \( P. \text{ultimum} \) DNA versus \( C_T \) values were also analyzed using SLR for estimation of the curve formula. Percentage data were transformed [arcsine \( \sqrt{p} \)] where \( p \) is the proportion of sporangia activated or with germ tubes and seed colonization incidence) to stabilize the normality and variance when necessary. The response variable “amount DNA” in the \( P. \text{ultimum} \) biomass data sets was transformed \( \left[-\log (\text{pg DNA} + 1)\right] \) to achieve normality. Appropriate diagnostic plots were established to ensure that all assumptions of the tests were fulfilled and to check for influential points. Insignificant terms (\( \alpha = 0.05 \)) were dropped in all tested models. Results considered to be significant are comparisons with \( P < 0.05 \).

**RESULTS**

**Seed characterizations.** Seed imbibition and germination percentages were used to gauge seed exudation rates and ensure seed viability and seedling quality. The germination rates of both cucumber and corn seeds were \( >99\% \). Both cucumber and corn seeds rapidly imbibed a significant amount of water within the first 30 min and continued to take up water over the 12-h sampling period (data not shown). When imbibition rates were expressed as mg H\(_2\)O/mg seed, no differences were observed between corn and cucumber seeds during the first 4 h of imbibition. Beyond 4 h, significantly greater amounts of water were taken up by corn seeds than by cucumber seeds. Corn seeds weighed an average of 163 (\( \pm 2.5 \)) mg, which is about seven times greater than the weight of cucumber seeds (25 [\( \pm 0.6 \)] mg). Therefore, expressing imbibition rates as mg H\(_2\)O/seed revealed that the overall amount of water imbibed by corn seeds greatly exceeded that of cucumber as early as 30 min after sowing.

\( P. \text{ultimum} \) sporangial activation and germ tube emergence in the spermosphere. A low but significant percentage of \( P. \text{ultimum} \) sporangia were activated for germination in the cucumber spermosphere within the first 30 min of exposure to the seed (Fig. 1A). Maximum levels of sporangial activation (60 to 70\%) were observed by 1.5 to 2.5 h of exposure to seeds. In contrast, 44\% of the sporangia were activated in the corn spermosphere as early as 15 min after exposure to seeds. Maximum activation levels (80 to 100\%) were observed as early as 30 min after seed exposure.

No significant germ tube emergence (direct evidence for sporangial germination) occurred in the cucumber spermosphere until 2 to 2.5 h of exposure to the seed (Fig. 1B). By 5 to 6 h after exposure to cucumber seeds, the percentage of sporangia with emerged germ tubes reached a maximum level of 60 to 70\%. In contrast, a significant percentage of sporangia with emerged germ tubes were observed in the corn spermosphere as early as 1 to 1.5 h after exposure to seeds. Maximum germ tube emergence (>90\% germination) was observed after 4 h of exposure to corn seeds.

Sporangial activation and germ tube emergence in response to corn and cucumber seeds increased linearly and significantly (\( P < 0.0001 \)) over the first few hours of seed germination until the response was maximized (Table 1). The rate of increase in the cucumber spermosphere was significantly lower than that in the corn spermosphere.

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**FIG. 1.** Activation (A) and germ tube emergence (B) of \( P. \text{ultimum} \) sporangia in the corn (○) and cucumber (□) spermospheres and in the absence of seeds (○) during the first 4 and 6 h of seed germination, respectively. Activation was assessed as percent sporangia with visible germ tubes after sporangia were exposed to seeds for various periods of time and then incubated in the absence of seeds, allowing germ tubes to emerge. Each point with the error bars represents the mean ± standard deviation of 15 observations from three repeated experiments.
Effects of \textit{E. cloacae} cell density and \textit{fad} mutants on \textit{P. ultimum} sporangial activation in the spermosphere. These experiments focused on sporangial activation and not germ tube emergence with the reasoning that the exudates responsible for inducing initial sporangial activation are those that \textit{E. cloacae} must inactivate to effectively prevent germ tube emergence. Significant reductions in sporangial activation were observed upon introducing \textit{E. cloacae} cells to cucumber seeds but only when cells were applied at the highest rate of $10^8$ cells/cm$^3$ substrate (Fig. 2A and C). Significant reductions in percentages of activated sporangia were observed at all sampling points up to 6 h (Table 2). The wild-type strain of \textit{E. cloacae} (strain EcCT-501R3) reduced the percentage of activated sporangia by approximately 45% in the cucumber spermosphere over that for seeds not treated with \textit{E. cloacae}. However, \textit{fad} mutants of \textit{E. cloacae} (strains EcL1 and Ec31) applied at $10^8$ cells/cm$^3$ substrate were ineffective in reducing sporangial activation in the cucumber spermosphere at any time point tested (Fig. 2C; Table 2). No reductions in sporangial activation at any cell density by any of the \textit{E. cloacae} strains were observed in the corn spermosphere at any sampling time (Fig. 2B and D and Table 2).

\textit{P. ultimum} colonization of seeds treated with \textit{E. cloacae}. The incidence of corn and cucumber seeds colonized by \textit{P. ultimum} increased at a constant rate during seed germination, reaching 100% colonization incidence by 6 h after sowing (Fig. 3A and B). No significant differences in colonization incidence between corn and cucumber seeds were observed at any sampling point. Neither strain EcL1 nor Ec31 reduced the incidence of \textit{P. ultimum} seed colonization in the cucumber spermosphere. \textit{E. cloacae} strain EcCT-501R3 reduced \textit{Pythium} colonization incidence in the cucumber spermosphere only at the 8-h sampling point. The rate of increase in seed colonization incidence in the cucumber spermosphere was significantly lower for seeds treated with \textit{E. cloacae} strain EcCT-501R3 than for nontreated seeds (Table 2). None of the \textit{E. cloacae} strains reduced seed colonization incidence in the corn spermosphere at any sampling time (Fig. 3B and Table 2).

Biomass of \textit{P. ultimum} on corn and cucumber seeds treated with \textit{E. cloacae}. The incidence of seeds colonized by \textit{P. ultimum} provided little information on the absolute amount of \textit{P. ultimum} seed colonization. On the other hand, quantitative estimates of \textit{P. ultimum} biomass provided a better indication of the overall level of seed colonization. \textit{P. ultimum} was detected on both cucumber (43 pg \textit{P. ultimum} DNA/seed) and corn (75 pg \textit{P. ultimum} DNA/seed) seeds as early as 4 h after sowing (Fig. 4A and B). By 8 h after sowing, an approximately 10-fold increase of \textit{P. ultimum} biomass was detected (501 and 721 pg DNA per cucumber and corn seed, respectively). However, these levels were not significantly different from those at 4 h after sowing. Levels of \textit{P. ultimum} biomass on cucumber seeds at 12 h after sowing (677 pg \textit{P. ultimum} DNA/seed at 12 h) did not differ significantly from levels detected at 8 h after sowing. In contrast, levels of \textit{P. ultimum} biomass on corn seeds were significantly greater by 12 h after sowing (3,470 pg \textit{P. ultimum} DNA/seed) than those observed at 8 h after sowing. No \textit{P. ultimum} DNA was detected on nontested seeds at any of the sampling times.

Introducing \textit{E. cloacae} strain EcCT-501R3 to cucumber seeds significantly reduced \textit{P. ultimum} biomass development compared to that observed on nontreated seeds at all time points tested (Fig. 4A). The rate of biomass increase over time was significantly lower on seeds treated with strain EcCT-501R3 than on nontreated seeds (Table 2). \textit{P. ultimum} biomass on corn was not significantly reduced by either of the two \textit{E. cloacae} strains (Fig. 4B and Table 2).

Reduction of \textit{P. ultimum} biomass development on cucumber seeds by \textit{E. cloacae} strain Ec31 was variable (Fig. 4A). By 4 and

### Table 1. Regression analysis of \textit{P. ultimum} sporangial activation and germ tube emergence in response to cucumber and corn seeds during the first 4 h of seed germination

<table>
<thead>
<tr>
<th>Assay and seed type</th>
<th>$r^ab$</th>
<th>LS mean$^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Activation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Corn</td>
<td>0.803**</td>
<td>97.84 a</td>
</tr>
<tr>
<td>Cucumber</td>
<td>0.822**</td>
<td>23.65 b</td>
</tr>
<tr>
<td>Germ tube emergence</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Corn</td>
<td>0.910**</td>
<td>72.16 a</td>
</tr>
<tr>
<td>Cucumber</td>
<td>0.922**</td>
<td>15.59 b</td>
</tr>
</tbody>
</table>

$^*$ Regression coefficient for response (activation and germ tube emergence) versus time.

$^b$ Significant ($P < 0.001$) by F test for the association between the response variable and time.

$^c$ Least-squares (LS) mean estimates from ANCOVA comparing the slopes for the activation and germination assays. Estimates followed by different letters are significantly different.
8 h after sowing, significantly lower levels of *P. ultimum* biomass developed on seeds treated with strain Ec31. However, by 12 h after sowing, levels of *P. ultimum* biomass developing on treated and nontreated seeds did not differ. Furthermore, the linear rate of biomass increase was not significantly lower in the presence of strain Ec31 than in its absence (Table 2).

**Relationship between *P. ultimum* sporangial activation and cucumber seed colonization.** Adding wild-type cells of *E. cloacae* to seeds after sporangia were fully activated (2 h after sowing) resulted in no significant decreases in *Pythium* biomass on seed surfaces compared to that on nontreated seeds at any sampling time after inoculation (Table 3). The rate at which *Pythium* biomass increased on seeds over time was significantly higher (*P* < 0.001) when *E. cloacae* cells were introduced 2 h after sowing than when cells were added at the time of sowing.

![FIG. 3. Frequency of cucumber (A) and corn (B) seeds colonized by *Pythium ultimum* in the presence of *E. cloacae* strain EcL1 (○), Ec31 (■), or EcCT-501R3 (△). Controls consisted of noninoculated seeds (□) as well as inoculated seeds not treated with bacteria (■). Each point represents the mean of three observations (six replicates each) from three repeated experiments.](image)

![FIG. 4. *Pythium ultimum* biomass development on cucumber (A) and corn (B) seeds in the presence of *E. cloacae* strain Ec31 (○) or EcCT-501R3 (△). Seeds were exposed to *Pythium* for 4, 8, and 12 h, after which amounts of *P. ultimum* DNA were determined using qPCR. Seeds that were inoculated but not treated with *E. cloacae* (■) and noninoculated seeds (□) served as positive and negative controls, respectively. Each point with the error bars represents the mean ± standard deviation of six seeds from two repeated experiments. Note the different y axes.](image)

### TABLE 2. Regression analysis of sporangial activation, seed colonization incidence, and seed colonization biomass development by *P. ultimum* on cucumber and corn seeds treated with *E. cloacae* strain EcCT-501R3, EcL1, or Ec31

<table>
<thead>
<tr>
<th>Seed type and treatment</th>
<th>Activation</th>
<th>Incidence</th>
<th>Biomass</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td><em>r</em>²a</td>
<td>LS meanb</td>
<td><em>r</em>²a</td>
</tr>
<tr>
<td>Cucumber</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Noninoculation</td>
<td>0.047 (NS)</td>
<td>0.106 c</td>
<td>0.000 (NS)</td>
</tr>
<tr>
<td>Inoculation with strain</td>
<td>0.920**</td>
<td>34.05 a</td>
<td>0.829**</td>
</tr>
<tr>
<td>EcCT-501R3</td>
<td>0.538**</td>
<td>7.14 b</td>
<td>0.532**</td>
</tr>
<tr>
<td>EcL1</td>
<td>0.853**</td>
<td>28.21 a</td>
<td>0.654**</td>
</tr>
<tr>
<td>Ec31</td>
<td>0.873**</td>
<td>28.21 a</td>
<td>0.709**</td>
</tr>
<tr>
<td>Corn</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Noninoculation</td>
<td>0.035 (NS)</td>
<td>0.079 b</td>
<td>0.004 (NS)</td>
</tr>
<tr>
<td>Inoculation with strain</td>
<td>0.783**</td>
<td>53.54 a</td>
<td>0.814**</td>
</tr>
<tr>
<td>EcCT-501R3</td>
<td>0.860**</td>
<td>48.66 a</td>
<td>0.797**</td>
</tr>
<tr>
<td>EcL1</td>
<td>0.906**</td>
<td>47.35 a</td>
<td>0.788**</td>
</tr>
<tr>
<td>Ec31</td>
<td>0.954**</td>
<td>47.96 a</td>
<td>0.555**</td>
</tr>
</tbody>
</table>

* a Regression coefficients for response (activation, colonization incidence, and colonization biomass) versus time, followed by the significance level of the F test for the association between the response variable and time. NS, nonsignificant; *, *P* < 0.01; **, *P* < 0.001.

* b Least-squares (LS) mean estimates from ANCOVA comparing treatment slopes. Estimates followed by different letters are significantly different.

* c ND, not determined.
TABLE 3. Regression analysis of P. ultimum biomass associated with cucumber seeds inoculated 2 h after sowing with E. cloacae strain EcCT-501R3 or Ec31

<table>
<thead>
<tr>
<th>Treatment</th>
<th>( r^{2b} )</th>
<th>LS mean(^a )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Noninoculation</td>
<td>0 (NS)</td>
<td>-0.000 b</td>
</tr>
<tr>
<td>Inoculation with strain</td>
<td>0.575**</td>
<td>813.96 a</td>
</tr>
<tr>
<td>EcCT-501R3</td>
<td>0.600**</td>
<td>485.34 a</td>
</tr>
<tr>
<td>Ec31</td>
<td>0.605**</td>
<td>610.34 a</td>
</tr>
</tbody>
</table>

\(^a\) Regression coefficient of the regression line for biomass versus time, followed by the significance level of the F test testing the association between the response variable and time. NS, nonsignificant; **, \( P < 0.001 \).

\(^b\) Least-squares (LS) mean estimates from ANCOVA comparing the slopes of treatments. Estimates followed by different letters are significantly different.

TABLE 4. Regression analysis of P. ultimum biomass in and on cucumber seeds inoculated with different numbers of sporangia per seed

<table>
<thead>
<tr>
<th>Time (h)</th>
<th>( r^{2b} )</th>
<th>LS mean(^a )</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>0.297*</td>
<td>18.23 a</td>
</tr>
<tr>
<td>8</td>
<td>0.637**</td>
<td>13.55 a</td>
</tr>
<tr>
<td>12</td>
<td>0.438**</td>
<td>8.46 a</td>
</tr>
</tbody>
</table>

\(^a\) Regression coefficient of the regression line for biomass versus number of sporangia, followed by the significance level of the F test testing the association between the response variable and time. *, \( P < 0.01 \); **, \( P < 0.001 \).

\(^b\) Least-squares (LS) mean estimates from ANCOVA comparing the slopes of treatments. Estimates followed by the same letter are significantly similar.

Strain Ec31 of E. cloacae behaved similarly to the wild-type strain.

Adjusting the potential number of activated sporangia in the spermosphere by exposing cucumber seeds to increasing dosages of sporangia ranging from 0 to 600 sporangia per seed resulted in increasing levels of P. ultimum biomass on cucumber seeds for each sampling time point (Table 4). Biomass levels at each of the sampling times were approximately 10% of those from experiments where sporangial inoculum was not manipulated (i.e., in sporangial disks) (data not shown). Within a given dosage, P. ultimum biomass levels on seeds did not increase significantly over time between 4 and 12 h.

**DISCUSSION**

The results of this study revealed that P. ultimum sporangia respond to exudates released into the spermospheres of corn and cucumber within 30 min after exposure to seeds, nearly 45 to 75 min before germ tubes emerge. Such an early response indicates that sufficient concentrations of LCFA must be present in the spermosphere within 30 min after sowing since LCFA are the only known elicitors of P. ultimum sporangial germination (10). Recent studies of fatty acid exudation from seeds have confirmed that LCFA, especially oleic and linoleic acids, are released from both corn and cucumber seeds as early as 15 min after sowing (23). Although it is difficult to know the precise concentration and distribution of LCFA in the spermosphere at any point in time, the calculated concentrations of oleic and linoleic acids are well within the concentrations previously reported to induce high levels of sporangial germination (18, 20). Assuming a spermosphere radius of 5 mm (10) surrounding a 5-mm-diameter seed, the volume of the spermosphere would be approximately 0.5 cm\(^3\). In this scenario the combined levels of oleic and linoleic acids released over the first 6 h of germination would yield around 226 \( \mu \)g/cm\(^3\), approximately the level known to induce maximum sporangial germination (18, 20).

Given that LCFA are present in the corn and cucumber spermospheres at stimulatory concentrations as early as 15 to 30 min after sowing, we would predict that the percentage of activated sporangia should be reduced in the cucumber spermosphere but not in the corn spermosphere in the presence of E. cloacae. Although previous studies demonstrated that E. cloacae can reduce the stimulatory activity of cucumber but not corn seed exudates collected for as little as 2 h after initiating seed imbibition in vitro (4), it was not previously known whether exudate inactivation can occur within 30 min in the spermosphere. Our results confirm that exudate inactivation occurs in the cucumber spermosphere but not the corn spermosphere in the presence of E. cloacae within 30 min of sowing, suggesting that E. cloacae actively degrades fatty acid elicitors present in the spermosphere, which can explain the observed reduction in sporangial activation.

Sporangial activation experiments with cucumber and corn in the presence of fatty acid degradation mutants of E. cloacae were carried out to provide further evidence that the failure of E. cloacae to suppress sporangial activation in the corn spermosphere is due to the failure to degrade fatty acid elicitors in the spermosphere. Previous in vitro studies with cottonseed exudates demonstrated that fatty acid degradation by E. cloacae was required to prevent P. ultimum sporangial germination and eliminate subsequent disease development (20, 21). If the degradation of fatty acid elicitors by E. cloacae can explain the reduced sporangial activation in the cucumber spermosphere, such reduced activation should be eliminated in the cucumber spermosphere in the presence of the E. cloacae mutants. Our results support these predictions since both the fatty acid mutants failed to reduce sporangial activation in the cucumber or corn spermospheres at any time after sowing.

Collectively, our observations of sporangial activation in the presence of E. cloacae indicate that the failure of E. cloacae to protect corn seedlings from Pythium damping-off is due not to a lack of synchronization between E. cloacae fatty acid degradation and seed exudation of fatty acids but to other factors. It is more likely that the inability of E. cloacae to protect corn seeds from infection is due to the failure of E. cloacae to degrade fatty acid elicitors released into the spermosphere soon after sowing. It has been shown in other studies (16, 17) as well as in preliminary studies in our lab (S. Windstam and E. B. Nelson, unpublished data) that corn seeds support a high level of growth and activity of E. cloacae in the spermosphere. This suggests that the growth and development of E. cloacae are not inhibited in the corn spermosphere, making it unlikely that the lack of seed protection is due to the general inhibition of E. cloacae metabolism by corn seed exudates. We hypothesize, therefore, that the inability of E. cloacae to reduce P. ultimum infections in the corn spermosphere is due to the suppression of \( \beta \)-oxidation by other components of corn seed exudates. This hypothesis is explored more fully in a companion paper (23).

Successful disease development by P. ultimum is dependent not only on the activation and germination of sporangia in the
spermosphere but also on the rapid colonization of seeds and seedling surfaces, ultimately resulting in embryo or seedling infection (11). Our results have shown that P. ultimum colonization levels on cucumber seeds are reduced when E. cloacae is active. Yet, either such reductions in colonization could be due to reduced sporangial activation and germ tube emergence or E. cloacae may continue to interact with P. ultimum mycelia after germ tube emergence to reduce seed colonization and subsequent disease development.

Two approaches were used to determine whether the reduction in P. ultimum biomass by E. cloacae on cucumber seeds was due solely to reductions of sporangial activation. In the first approach, E. cloacae cells were added to cucumber spermospheres only after sporangia were fully activated (2 h). If disruption of sporangial activation is the principal cause of reduced P. ultimum biomass, adding E. cloacae cells after full activation should result in seed colonization levels greater than or equal to those observed on nontreated seeds or seeds coated with E. cloacae at the time of sowing. In the second approach, seeds were exposed to increasing levels of activated sporangia in the spermosphere, there should be a direct correlation between sporangial dosage and biomass development. Such dose-dependent responses have been frequently described for Pythium species (6). Using both approaches, our results confirmed the experimental predictions—P. ultimum seed colonization was not reduced if E. cloacae cells were added after full sporangial activation. Furthermore, increasing the numbers of sporangia to which seeds were exposed resulted in increasing amounts of seed colonization biomass. These findings are significant because they provide further evidence that the behavior of sporangia alone can establish the ultimate levels of seed colonization and subsequent disease incidence and severity. Therefore, reductions in sporangial activation should translate directly into reductions in disease development.

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REFERENCES


