Plants Secrete Substances That Mimic Bacterial N-Acyl Homoserine Lactone Signal Activities and Affect Population Density-Dependent Behaviors in Associated Bacteria

Max Teplitski,1 Jayne B. Robinson,2 and Wolfgang D. Bauer1

1Horticulture and Crop Science Department, Ohio State University, Columbus 43210, U.S.A.; 2Biology Department, University of Dayton, Dayton, OH 45469, U.S.A.

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In gram-negative bacteria, many important changes in gene expression and behavior are regulated in a population density-dependent fashion by N-acyl homoserine lactone (AHL) signal molecules. Exudates from pea (Pisum sativum) seedlings were found to contain several separable activities that mimicked AHL signals in well-characterized bacterial reporter strains, stimulating AHL-regulated behaviors in some strains while inhibiting such behaviors in others. The chemical nature of the active mimic compounds is currently unknown, but all extracted differently into organic solvents than common bacterial AHLs. Various species of higher plants in addition to pea were found to secrete AHL mimic activities. The AHL signal-mimic compounds could prove to be important in determining the outcome of interactions between higher plants and a diversity of pathogenic, symbiotic, and saprophytic bacteria.

Additional keywords: chitinase, quorum sensing, swarming, violacein.

In recent years, it has become clear that many important behaviors in bacteria are regulated in a population density-dependent manner, including behaviors such as motility and the synthesis of exoenzymes, exopolysaccharides, surfactants and antibiotics that are crucial to attacking or colonizing plant, animal, or microbial hosts (Eberl et al. 1996; Glessner et al. 1999; for reviews see Fuqua et al. 1996; Pierson et al. 1999; Swift et al. 1999). A recent study has revealed that transcription of over 250 genes is affected by the N-acyl homoserine lactone (AHL) signaling system in Pseudomonas aeruginosa (Whiteley et al. 1999). The synthesis of AHL signals is common among plant-associated bacteria (Cha et al. 1998) and AHL signaling has been well studied in several of these bacteria (for review see Pierson et al. 1999).

Bacterial cell-cell signaling and population density-dependent regulation of behavior (“quorum sensing”) involves the synthesis and perception of small, diffusible signal molecules that move in and out through the cell membrane and between cells. In gram-negative bacteria, AHLs are the best studied of the diffusible signal compounds that govern population density-dependent behaviors, but there is recent evidence for other types of intercellular signal compounds, including cyclic dipeptides and quinolones in P. aeruginosa (Holden et al. 1999; Pesci et al. 1999), unknown low molecular weight substances in Xanthomonas campestris (Poplawsky et al. 1998), and a volatile fatty acid methyl ester in Ralstonia solanacearum (Flavier et al. 1997). In gram-positive bacteria, many behaviors are also regulated in a population density-dependent manner (cf. reviews by Dunny and Leonard 1997; Horinouchi 1999; Lazazzera et al. 1999). The quorum-sensing signal substances in gram-positive species have been identified as γ-butyrolactones, structurally quite similar to AHLs, and post-translationally modified peptides.

A given bacterial species will commonly synthesize several different AHL signals that differ from one another in the length of the N-acyl side chain (four to 14 carbons), the presence or absence of double bonds, or side chain substituents (keto, hydroxyl) (Swift et al. 1999). AHLs usually have their effects by binding semi-specifically to receptor proteins homologous to LuxR, the prototypical AHL receptor from Vibrio fischeri (Choi and Greenberg 1992; Fuqua et al. 1996; McClean et al. 1997). At sufficiently high population densities, the AHLs reach a “threshold” concentration so that AHL binding to the cognate LuxR-like receptor activates the transcription of specific genes or sets of genes (Stevens and Greenberg 1999). A given bacterial species may have several different LuxR-like receptors for AHLs, each controlling different behaviors. In some cases, AHL binding to a receptor induces increased synthesis of the AHL, leading to rapidly self-amplifying behavioral changes in a set of cells (Fuqua et al. 1996). In other cases, the binding of one AHL controls the synthesis of one or more other AHLs, leading to complex regulatory hierarchies (Pesci and Iglewski 1999; Rodelas et al. 1999). There is evidence that AHLs secreted by cells of one species can induce population density-dependent responses in cells of other bacterial species in natural rhizosphere environments (Pierson et al. 1998). Such interpopulation AHL signaling may be significant to bacterial communities.

An important recent study has demonstrated that halogenated furanones produced by the marine red algae Delisea pul-
**RESULTS**

*C. violaceum* wild type and CV026 reporter.

The CV026 reporter strain is a mutant of wild-type *C. violaceum* ATCC 31532, which cannot synthesize N-hexanoyl-L-homoserine lactone (= C₆-HSL). It is thus reliant on exogenously added AHLs for activation of its AHL receptor, CviR, to induce synthesis of the purple antibiotic violacein and a number of exoenzymes (McClean et al. 1997; Chemin et al. 1998; Swift et al. 1999). Violacein production in CV026 is induced by AHLs with a short (C₄-C₈) alkanoyl or 3-oxo-alkanoyl side chain or by a number of their chemical analogs (McClean et al. 1997; Cha et al. 1998). Addition of an AHL with a long alkanoyl or 3-oxo-alkanoyl side chain (e.g., 3-oxo-C₁₂-HSL) competitively inhibits violacein synthesis induced by short side chain AHLs such as N-butanoyl-L-homoserine lactone (= C₄-HSL). Thus, CV026 is an excellent reporter for a wide range of bacterial AHLs (or AHL signal-mimic compounds from plants).

AHL-regulated behaviors in *C. violaceum* CV026 were strongly inhibited by substances secreted by pea seedlings. When 5-day-old, aseptically grown pea seedlings were placed on the surface of nutrient agar colonized by a lawn of the CV026 reporter strain, the seedlings induced no detectable violacein synthesis (Fig. 1A), in contrast to the positive C₄-HSL control (Fig. 1C). However, when placed on agar containing 4 μM C₆-HSL, the pea seedlings visibly inhibited C₆-HSL-induced violacein synthesis in the lawn of CV026, leaving colorless areas around the seedlings (Fig. 1B). No inhibition of bacterial growth around the seedlings was observed, and none was detected in subsequent assays of CV026 growth in shake cultures containing pea exudates. The inhibition of violacein synthesis in areas adjacent to the seedlings was similar to that caused by the addition of a long chain AHL to a lawn of C₆-HSL-induced CV026 (see Figure 1D, and McClean et al. 1997).

Substances that inhibited AHL-induced violacein synthesis in *C. violaceum* CV026 were present in crude exudates collected by gentle rinsing of aseptically grown pea seedlings and in methanol extracts of lyophilized exudate. As shown in Figure 2A, C₆-HSL-inducible violacein synthesis was inhibited by a methanol extract of pea exudate in rough proportion to the amount of extract added. Inhibition of violacein synthesis by the extract could be partially prevented or reversed by an...
initial addition of twice as much C₄-HSL to the assay wells (Fig. 2B) or to the area adjacent to live seedlings (not shown). Little restoration of violacein synthesis was seen if additional C₄-HSL was added after 4 h or later. We have observed that CV026 cells remain responsive to C₄-HSL induction of violacein synthesis for a limited time while they are actively growing (data not shown), perhaps accounting for the failure of delayed additions of C₄-HSL to more fully restore exudate-inhibited violacein synthesis. The progressively greater inhibition of AHL-induced violacein synthesis by greater amounts of pea exudate and the partial reversal of such inhibition by additional C₄-HSL both suggest, but do not prove, that pea seedlings secrete one or more substances capable of specifically interfering with the perception of C₄-HSL by CV026.

To more rigorously test the possible disruption of AHL signaling in C. violaceum by substances in pea exudates, we examined the effects of methanol extract preparations on extracellular protease and chitinase activities, both of which are also regulated by the same receptor for AHL signals as violacein synthesis in C. violaceum (Chernin et al. 1998; Swift et al. 1998).

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**Fig. 2.** Effects of pea seedling exudates on violacein synthesis and exoenzyme activities in the CV026 reporter strain. A, Violacein production induced by 0.5 µg (3 µM) of C₄-HSL per ml in the presence or absence of pea exudate was assayed colorimetrically after 20 h of incubation. B, An additional 0.5 µg of C₄-HSL per ml was added either 0 or 4 h after suspension of CV026 cells in fresh medium, with or without one seedling equivalent of the pea exudate extract, and violacein production assayed colorimetrically after 20 h. C, Extracellular protease activity induced after 20 h by 0.5 µg of C₄-HSL per ml in the presence or absence of pea exudate. D, Exochitinase (p-nitrophenyl-β-D-N-acetylglucosaminidase) activity induced after 20 h by 0.5 µg of C₄-HSL per ml in the presence or absence of pea exudate. All assays were performed in duplicate in repeated experiments as described in Materials and Methods.
As shown in Figure 2C and D, the addition of seedling exudate extract substantially inhibited both C₄-HSL-inducible protease and N-acetylglucosaminidase (exochitinase) activities in CV026. When assayed on colloidal chitin plates as described by Chernin et al. (1998), pea seedling exudates also substantially inhibited AHL-dependent endochitinase activity (data not shown). Growth of CV026 was not appreciably altered by pea exudates in these experiments, as determined by plate counts and absorbance measurements (data not shown). In other control experiments, CV026 cultures were first incubated with C₄-HSL to induce extracellular protease and N-acetylglucosaminidase, then mixed with methanol-extracted substances from pea exudate and immediately assayed. Addition of the pea exudate extract after induction by C₄-HSL had no effect on these activities, indicating that the extracts did not contain substances that directly inhibited the enzymes. When C₄-HSL was incubated overnight at room temperature with high-pressure liquid chromatography (HPLC)-purified pea extracts containing the violacein inhibitory activity, it was found that the C₄-HSL could be fully recovered from the mixture by ethyl acetate extraction (data not shown), suggesting that the active materials in seedling exudates did not degrade or immobilize the exogenously added C₄-HSL. Based on these results, it seems unlikely that the coordinate inhibition of AHL-induced violacein, protease, and chitinases by pea exudates seen in Figure 2 is an artifact of some nonspecific interference with growth, violacein biosynthesis, exoenzyme synthesis or activity, or C₄-HSL availability. Reasonable mechanisms of action to consider include direct interactions of the plant compounds with the receptor, CviR (as expected for a true signal-mimic compound like the D. pulchra furanones) or indirect effects such as reduced bacterial uptake of the exogenously added C₄-HSL.

As shown in Figure 3, AHL-regulated extracellular protease activity was inhibited by pea exudate extracts to about the same extent in wild-type C. violaceum as in the CV026 reporter strain supplied with exogenous C₄-HSL. Pea exudate from five seedlings was about as effective as 4 µM exogenous 3-oxo-C₁₂-HSL, a known competitive inhibitor of violacein synthesis (McClean et al. 1997). The addition of pea exudate extract or 3-oxo-C₁₂-HSL had no appreciable effect on growth of the bacteria (Fig. 3). These results are important because they suggest that the normal synthesis of cognate AHLS (C₄-HSL in this case) by wild-type bacteria may not be able to mask or overwhelm the effects of secreted plant signal-mimic compounds in natural encounters. Further studies are clearly needed to explore the question of just how effectively the plant compounds can compete with endogenously synthesized AHLS to affect behavior in wild-type bacteria.

S. liquefaciens reporters.

In wild-type S. liquefaciens MG1, the initiation of swarming requires surface-stimulated induction of the flhDC operon, leading to formation of hyperflagellated cells (Givskov et al. 1997). Surface swarming also depends on C₄- and/or C₆-HSL-induced synthesis of serrawettin, a lipopeptide surfactant (Eberl et al. 1999). The S. liquefaciens strain MG44 is a swrI::Tn5 mutant of MG1 that cannot synthesize its own C₆-HSL and C₆-HSL and therefore requires exogenous AHLS for induction of serrawettin synthesis and swarming motility (Lindum et al. 1998). Kjelleberg et al. (1997) used MG44 as a reporter to show that the halogenated furanones of D. pulchra inhibited AHL-induced surface swarming and other AHL-regulated behaviors in S. liquefaciens (as well as violacein synthesis in C. violaceum). They subsequently showed that serrawettin synthesis required expression of the swrA gene and isolated PL10, an swrA::Tn5luxAB mutant derivative of MG44. Thus, PL10 serves as a specific transcriptional bioluminescence reporter for the presence of exogenous AHLS capable of binding to the AHL receptor (SwrR) that activates the swrA gene and serrawettin synthesis (Lindum et al. 1998).

As shown in Figure 4, S. liquefaciens swarming was strongly affected by substances secreted by pea seedlings. The reporter strain MG44, unable to make its own AHLS, grew rapidly on the agar surface, but did not swim (Fig. 4A). When aseptically grown pea seedlings were inoculated with MG44, placed on nutrient agar, and incubated overnight, the seedlings induced the bacteria to swarm outward over the agar surface (Fig. 4B). As controls, toothpicks were inoculated with MG44 in the same manner. The inoculated toothpicks developed dense bacterial growth along their length, but did not induce surface swarming (Fig. 4C) unless previously dipped in a solution of C₆-HSL (Fig. 4D). A methanol extract of pea seedling exudate, equivalent to the amount secreted by one 7-day-old pea seedling, strongly stimulated swarming of MG44 when present on a filter disk with the bacteria (Fig. 4E).

The S. liquefaciens strain PL10 was used to test whether the plant-induced swarming of MG44 (Fig. 4B and E) was caused by secretion of a surfactant by the seedlings. Since PL10 is defective in both AHL and serrawettin synthesis, it does not swarm even in the presence of exogenous AHLS. However, PL10 can be stimulated to swarm by exogenous serrawettin or other surfactants (Lindum et al. 1998). PL10 thus provides a direct bioassay for exogenous surfactants. As shown in Figure 4F, pea seedlings did not induce surface swarming of PL10,
indicating that pea does not secrete a surfactant capable of stimulating swarming.

While pea seedlings and methanol extracts of seedling exudate had no effect on swarming of PL10, they did stimulate AHL-dependent luminescence in this reporter, as described below.

Other AHL reporters.

Another set of AHL reporters was recently developed for sensitive, quantitative, real-time detection of exogenous AHLS (Swift et al. 1997; Winson et al. 1998). These AHL reporters normally function as multicopy plasmids in E. coli (a species that does not synthesize AHLS). The plasmids carry a gene for an AHL receptor protein (the LuxR receptor from V. fisheri, AhyR from A. hydrophila, or LasR from P. aeruginosa). Upon addition of an appropriate AHL, the receptor is activated so that it binds to its cognate luxI promoter and initiates transcription of a luxCDABE cassette, generating luminescence that is proportional to the concentration of added AHL (Winson et al. 1998). These three reporters differ in their sensitivity to a series of alkanoyl and 3-oxo-alkanoyl homoserine lactones (Winson et al. 1998). For example, the LuxRI′ reporter (pSB401) responds strongly to nanomolar concentrations of 3-oxo-C₆-HSL and to micromolar concentrations of C₆-HSL and 3-oxo-C₁₂-HSL, whereas the LasRI′ reporter (pSB1075) responds strongly to nanomolar concentrations of 3-oxo-C₁₂-HSL and C₁₂-HSL (Winson et al. 1998). The AhyRI′ reporter (pSB536) responds most strongly to C₇-HSL, its cognate AHL signal (Swift et al. 1997). Thus, collectively, these three reporters can detect a fairly broad diversity of AHLS (or AHL signal-mimic compounds).

Figure 5 shows the luminescence of these three E. coli plasmid reporters and that of S. liquefaciens PL10 after inoculation onto live, 4-day-old pea seedlings. The digital images of the seedlings were overlaid with false color luminescence images taken with a very sensitive Hamamatsu charged couple device (CCD) camera 8 to 9 h after inoculation. The luminescence induced in the reporter strains by the pea seedlings was generally two- to 20-fold above levels of the no-AHL controls. The pea seedlings induced luminescence in each reporter that was an appreciable fraction (10 to 75%) of the maximal levels obtainable with high concentrations of an appropriate AHL. This suggests that the concentration of putative AHL mimic compounds at the plant surface may be high enough to affect AHL-regulated gene expression in wild-type bacteria in natural encounters.

Plant-to-plant variation in luminescence was most frequently within about twofold for each of the reporters. The intensity of luminescence was normally greatest on the lower 2/3 of the pea root for each of the four AHL luminescence reporters. Luminescence was usually low on the upper portion of the root, the epicotyl, and the cotyledons. It will be of inter-

Fig. 4. Effects of pea seedlings and seedling exudates on N-acyl homoserine lactone (AHL)-dependent swarming of the Serratia liquefaciens MG44 reporter strain. Aseptically grown pea seedlings and toothpick controls were inoculated with a suspension of S. liquefaciens MG44 cultured as described in Materials and Methods. Inoculated seedlings or toothpicks were then placed on AB glucose-Casamino Acid agar and incubated for 18 to 20 h. A, Filter disk inoculated with MG44. B, Pea seedling inoculated with MG44. C, Sterile toothpick inoculated with MG44. D, Sterile toothpick dipped in 0.5 µg of C₆-HSL per ml, then inoculated with MG44. E, Filter disk containing 1 seedling equivalent of an evaporated 50% methanol-water extract of pea seedling exudate and inoculated with MG44. F, Pea seedling inoculated with PL10.
est to determine whether this pattern of luminescence reflects differences in the relative amounts of putative AHL mimic compounds secreted by the plant in these different parts of the seedlings or reflects differences in the number or metabolic vigor of associated bacterial reporter cells.

The ability of pea seedlings to stimulate the *E. coli* plasmid reporters provides prima facie evidence that at least some of the active plant compounds interact directly and specifically with the AHL receptor protein and affect its ability to initiate transcription of the *luxCDABE* cassette. In the *E. coli* plasmid reporters, there is essentially nothing else for exogenous substances to interact with to induce luminescence. The seedling-induced luminescence of PL10 is also consistent with direct interaction of the plant substances with the SwrR receptor, with resultant transcriptional activation of *swrA::Tn5luxAB*. In *S. liquefaciens*, however, there may be other, unknown mechanisms for activation of the *swrA* gene.

**AHL signal-mimic activity in other plant species.**

Aseptically grown, uncontaminated seedlings of several other plant species were assayed for possible secretion of AHL mimic activity with the above reporter strains in the same manner as shown in Figures 1, 4, and 5 for pea. None of the plant species tested were able to stimulate detectable violacein synthesis in *C. violaceum* CV026 or phenazine synthesis in the *P. aureofaciens* 30-84I reporter (data not shown). Only pea and crown vetch secreted compounds that inhibited AHL-induced synthesis of violacein in the *C. violaceum* CV026 reporter strain, doing so without any apparent inhibition of growth of the reporter strain (Table 1). Rice, soybean, tomato, crown vetch, and *Medicago truncatula* all activated AHL-dependent swarming in *S. liquefaciens* MG44 (data not shown). As indicated in Table 1, these plant species also stimulated luminescence in *S. liquefaciens* PL10 and in at least one of the three *E. coli* plasmid reporters. Neither lettuce nor *Arabidopsis thaliana* stimulated significant activity in any of the reporter strains under our assay conditions. The results from this small survey suggest that secretion of AHL signal-mimic compounds may be widespread among higher plants and at least quantitatively variable among species. This is what one might expect if AHL mimic synthesis by plants were an important and co-evolved mechanism for dealing with associated bacteria.

**Initial purification and characterization of the AHL signal-mimic substances.**

The chemical nature of the substances from pea that affected AHL-inducible behaviors in the *C. violaceum, S. liquefaciens*, and *E. coli* plasmid reporters is presently unknown. The addition of homoserine at concentrations equivalent to or 10-fold higher than those found in pea root exudates (van Egeraat 1975) had no effect in the *S. liquefaciens* MG44 swarming assay or *C. violaceum* CV026 violacein assays. When 10-mg samples of lyophilized pea seedling exudate were successively extracted with 20-ml portions of chloroform, ethyl acetate, 100% methanol, and 1:1 methanol-water, no activity was detectable in the chloroform or ethyl acetate extracts by either the direct or indirect *C. violaceum* assays or

![Fig. 5](image-url) Pea seedling-induced luminescence of N-acyl homoserine lactone (AHL) reporter strains. Luminescence of reporter strains on seedling surfaces was measured and imaged as described in Materials and Methods. Intensity of luminescence is indicated by color (red>yellow>green>blue) and by numbers (photon counts per second). Relative luminescence of reporters on seedlings is also given as a percentage of maximum luminescence inducible by exogenous AHL, normalized to equal numbers of bacteria. Normally, about 1–2 × 10^6 cells of the *Escherichia coli* plasmid reporter strains and about 5 × 10^7 cells of the PL10 reporter were recovered per seedling. Results are representative of several independent experiments.
by the *S. liquefaciens* swarming or luminescence assays. Chloroform and ethyl acetate can extract the known AHLs made by bacteria, and did extract exogenously added C_4-HSL from crude pea exudate preparations (data not shown). Thus, it appears that the plant substances with AHL signal activity are chemically different from bacterial AHLs.

Almost all of the activity in crude pea exudate preparations and detectable by the *S. liquefaciens* swarming and bioluminescence assays was present in 100% methanol extracts. Microtiter plate dilution endpoint assays could detect about 1/4 of one seedling’s activity in the methanol extracts with the PL10 luminescence reporter and X-ray film as described in Materials and Methods (data not shown). Swarm plate assays with MG44 reached a dilution endpoint at extract concentrations roughly fivefold higher than in the luminescence assays, and were considerably more variable. The activity detectable by the *C. violaceum* CV026 indirect assay was usually present in the 50% methanol:water extract, although it could be fully, or almost fully, extracted into 100% methanol if freshly lyophilized material was used. About 1/8 of one seedling’s activity could be detected by dilution endpoint assays of violacein production with the CV026 reporter, as described in Materials and Methods (data not shown). Thus, essentially all of the AHL mimic substances secreted by pea and detectable with these reporters are soluble in 100% methanol. From these semiquantitative estimates of recovered activity, it seems that the mimic substances may be secreted in sufficient quantities to affect AHL-mediated behaviors of bacteria associated with the plant surface in a natural environment. This is consistent with the effects of intact, live seedlings on the behavior of the reporter strains (Figs. 1, 4, and 5).

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**Table 1.** Responses of *N*-acyl homoserine lactone (AHL) reporter strains to substances secreted by seedlings of various plant species

<table>
<thead>
<tr>
<th>Plant</th>
<th>Relative response of luminescence reporters (%) a</th>
<th>Relative inhibition of AHL-induced violacein synthesis.</th>
<th>b</th>
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<tr>
<td></td>
<td>CV026</td>
<td>PL10</td>
<td>AhyRI′</td>
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<tr>
<td>Pea</td>
<td>***</td>
<td>13</td>
<td>35</td>
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<tr>
<td>Crown vetch</td>
<td>++</td>
<td>68</td>
<td>100</td>
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<tr>
<td><em>Medicago truncatula</em></td>
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<tr>
<td>Rice</td>
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<td>Lettuce</td>
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<td><em>Arabidopsis thaliana</em></td>
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*a* Photon counts per second from two individual 4-day-old seedlings inoculated with a luminescence-based reporter strain were measured with the charged couple device (CCD) camera as described in Materials and Methods. Relative luminescence of each seedling is reported individually, and was calculated by subtracting background luminescence of the no-AHL controls from the luminescence of each seedling and expressing the remainder as a percentage of the luminescence of the +AHL controls, normalized to equal numbers of reporter bacteria. Similar results were obtained in repeated experiments.

*b* Relative inhibition of AHL-induced violacein synthesis. *** = very strong; ** = strong.

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**Fig. 6.** High-pressure liquid chromatography (HPLC) fractionation of *N*-acyl homoserine lactone (AHL) mimic activities present in methanol extracts of pea seedling exudate. Fractions from a C_{18} reverse phase HPLC column were collected and samples equivalent to exudate from about two seedlings were assayed for activity as described in Materials and Methods. Luminescence was measured with the charged couple device camera. Fractions that induced *Serratia liquefaciens* PL10 luminescence are indicated by light green bars. These same fractions induced swarming of MG44 (not shown). Fractions that induced luminescence in the LuxRI′ reporter (pSB401) are indicated in black, in the AhyRI′ reporter (pSB536) in red, and in the LasRI′ reporter (pSB1075) in dark green. Relative inhibition of C_4-HSL-induced violacein synthesis in *Chromobacterium violaceum* CV026 is shown in purple, and was equal to 100% in fractions 8 and 9. Thick gray line indicates gradient of increasing acetonitrile concentration (0–100%) used for elution; thin black line indicates A_{254} (0–2). Similar results were obtained in repeated fractionations.
As shown in Figure 6, some important separation of different substances with AHL mimic activity was obtained by chromatographic fractionation of pea exudate extract on a C_{18} reverse phase column. The absorbance of fractions at 254 nm is shown as (arbitrary) indication of where the bulk of the material in the exudate extract eluted. None of the fractions from the C_{18} column contained substances that appreciably stimulated violacein synthesis in CV026 (data not shown). This is consistent with the failure of live pea seedlings to induce detectable violacein synthesis (Fig. 1A). When samples of each fraction equivalent to exudate from approximately two seedlings were assayed with CV026 and the luminescence reporters, all of the activity detectable by the CV026 indirect assay and some of the activity detected with the PL10, LasRI', and AhyRI' reporters and the S. liquefaciens MG44 reporter eluted in or near the void volume (fractions 8 to 15), suggesting that the compounds responsible for these activities could not bind to the C_{18} column because they are dominantly hydrophilic or ionic. Several activities detectable with the PL10, MG44, LasRI', AhyRI', and LuxRI' reporters eluted in fractions near the middle of the acetonitrile gradient (fractions 22 to 45). In addition, activities detectable with the AhyRI' and LasRI' reporters appeared in fractions near the end of the gradient (fractions 50 to 57).

When larger samples of the HPLC fractions were assayed (equivalent to exudate from six seedlings), one major new peak of PL10 activity was detectable in fractions 50 to 53 (data not shown). On a per seedling basis, the relative luminescence induced by material in this peak was about 70% of the +AHL control, likely accounting for most of the PL10 lucinescence seen with inoculated seedlings (Fig. 5). It will be of interest to determine whether fractions that stimulated responses in two or three different reporter strains contain one compound acting on two or three reporters or several different compounds, each acting on a separate reporter. Further purification and analysis to identify the active substances are in progress.

**DISCUSSION**

The halogenated furanones from *D. pulchra* provided the first clear example of AHL mimic compounds produced by a eukaryote (Givskov et al. 1996; Kjelleberg et al. 1997; Manefield et al. 1999). While quite limited, our initial examination of other plant species indicates that the synthesis of AHL signal-mimic substances may prove to be fairly general among higher plants. The apparent synthesis and secretion of AHL mimic compounds by diverse plant species substantially broadens the range of plant-microbe interactions where signal-mimic compounds might play important roles. Future investigations to detect AHL mimic compounds in additional plants could be valuable in exploring this possibility and in the search for new compounds to protect plants as well as humans and other animals from pathogenic bacteria.

The AHL signal-mimic activities detected in pea exudate and with live seedlings do not appear to be artifactual responses of the bacterial reporter strains. Pea seedlings and exudates inhibited three different behaviors under AHL control in *C. violaceum*. In *S. liquefaciens* MG44, the initiation of swarming depends on surface-stimulated induction of the *flhDC* operon, leading to formation of hyperflagellated cells, as well as on AHL-regulated serrawettin synthesis. Thus, swarming is not as specifically dependent on AHL binding as PL10 bioluminescence (Givskov et al. 1997; Lindum et al. 1998). In control experiments, the addition of live pea seedlings or exudate extract preparations did not appear to stimulate the *flhDC* operon, as measured by phospholipase assays (data not shown). Phospholipase is under control of the *flhDC* operon (Eberl et al. 1996). A further indication that swarming of *S. liquefaciens* MG44 was specifically induced by plant compounds acting as AHL mimics is the tightly parallel induction of MG44 swarming and PL10 bioluminescence by live seedlings of pea (Fig. 2), by other plants tested in Table 1, and by at least three separate fractions of pea exudate (Fig. 6). Based on such parallel induction of both swarming and swrA gene expression, and on evidence for the absence of serrawettin-like surfactants in the exudate, or compounds that activated the *flhDC* operon, or compounds that affected growth of the reporter strains, it seems likely that pea seedlings secrete compounds that specifically activate the SwrR receptor governing *swrA* expression and serrawettin synthesis.

Similarly, the activation of the LuxRI', AhyRI', and LasRI' plasmid reporters by seedlings of various plant species and by methanol extracts of pea seedling exudates suggests that these plants secrete compounds that specifically activate the AHL receptors, most likely by binding directly to the receptor protein. Our results with the PL10 and *E. coli* plasmid reporters appear to provide the first example of positively acting AHL signal-mimic compounds produced by eukaryotic organisms. It is worth noting, however, that others have shown that compounds that inhibit AHL-stimulated gene expression in cells with single copies of a LuxR-like receptor can show stimulatory activity when the receptor is present in multiple copies (Zhu et al. 1998). Thus, plant mimic compounds that stimulate the multicopy plasmid AHL reporters may act as competitive inhibitors of AHL-induced behaviors in wild-type bacteria.

Substances that can block quorum sensing in bacteria have been recently discussed as potentially valuable anti-infective agents for medicine (Finch et al. 1998). The substantial inhibition of three distinct AHL-inducible, CviR-regulated behaviors in CV026, and the partial reversal of such inhibition by addition of more C_{4}-HSL, makes it reasonable to suggest that pea plants secrete one or more compounds that specifically interfere with perception of AHL signals by the CviR receptor in the CV026 reporter strain. It will be of interest to learn whether exudates from higher plants contain substances that specifically inhibit other AHL-induced activities.

Further studies, preferably with pure mimic compounds, will be needed to rigorously establish the molecular mechanism(s) by which the AHL mimic compounds act. For example, plants might secrete “pseudo” mimic compounds that interfere with AHL synthesis or with AHL transport as well as “real” mimic compounds, like the halogenated furanones, that interact directly with the AHL receptor protein. It is of interest that *Pseudomonas aeruginosa* secretes cyclic dipeptides that act as AHL mimic compounds (Holden et al. 1999). The bacterial dipeptides appear to interact directly with AHL receptors and can either activate or inhibit receptor-mediated transcription, depending on the receptor.

Our initial attempts to purify the AHL mimic activity in pea exudates indicate that pea secretions at least one substance with
inhibitory activity in the CV026 reporter and perhaps half a dozen or more different substances that stimulate one or more of the other AHL reporter strains. Further fractionation may reveal additional activities. The sum of the relative activities observed in the HPLC fractions for each reporter agrees reasonably well on a per seedling basis with the total relative activity seen in Figure 5 with live pea seedlings, indicating that most of the activities present on the seedling surfaces were probably recovered and detected after fractionation. However, given the limited range of concentrations tested and the unknown sensitivity of the reporters toward particular mimic compounds, we suspect that the relative luminescence elicited by substances in these HPLC fractions may not be a very reliable indicator of either the amount or potency of the AHL mimic substances present. We also note that the mimic activities detected in fractions from the C18 column probably represent just those compounds that can interact with the CviR, LuxR, AhyR, LasR, and SwrR receptor proteins. The use of additional reporter strains that have AHL receptors with different binding specificities or affinities might well reveal additional AHL mimic activities. The synthesis and secretion of a diversity of AHL mimic substances, some acting positively in certain bacterial species, some acting negatively in other bacterial species, could be important to the coevolved biology of interaction between plants and their associated bacteria.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions.**

Stock cultures of *S. liquefaciens* MG1 wild-type and the reporter strains MG44 (swrI mutant km7) and PL10 (swrI mutant, swrA::Tn5luxAB km7 tc7), generously provided by M. Giviskov and S. Kjelleberg, were cultured on AB medium with glucose and Casamino Acids as described by Giviskov et al. (1997) with km50 or tc10, respectively. Cultures of *C. violaceum* wild-type ATCC31532 and the reporter strain CV026 (sm7 miniTn5 HgC cvil::Tn5xyIE km7) were generously provided by P. Williams, and cultured in LB (Luria-Bertani) broth. Lux plasmid AHL reporters *E. coli* JM109 pSB401 (luxR::luxCDABE tc7), *E. coli* JM109 pSB536 (ahyR::luxCDABE ap7), and *E. coli* JM109 pSB1075 (luxR::luxCDABE ap7), a generous gift from S. Swift, were grown in LB broth with appropriate antibiotics as described by Winson et al. (1998). *Pseudomonas aureofaciens* 30-84 and its phzI mutant, *P. aureofaciens* 30-84I (Wood et al.1997), a gift of F. Gong, were cultured on LB. All strains were maintained as glycerol stocks at –80°C and small samples were subcultured into fresh medium with antibiotics (when necessary) for the assays.

**Plant material.**

Pea (*Pisum sativum*) cv. Extra Early Alaska (Livingston Seed, Columbus, OH) was surface sterilized for 90 min in 95% ethanol, followed by 60 min in 70% commercial Chlorox bleach. Crown vetch (*Coronilla varia*) cv. Pennfield seeds (Livingston Seed) were scarified for 10 min in concentrated sulfuric acid, then surface sterilized in 95% ethanol for 30 min and in 50% Chlorox for 30 min. Seeds of *Medicago truncatula* genotype A17 (a gift from D. Cook) were surface sterilized for 60 min in 95% ethanol, followed by a 10 min treatment with full-strength commercial Chlorox bleach. Soybean (*Glycine max*) cv. Flint (from S. St. Martin), *Arabidopsis thaliana* ColO (from J.C. Jang), tomato (*Lycopersicon esculentum*) cv. Rutgers Select VFA (Geo. W. Park Seed, Greenwood SC), rice (*Oryza sativa*) cv. Nipponbare (from G. L. Wang), and romaine lettuce (*Lactuca sativa*) cv. Paris Island (Livingston Seed) were surface sterilized for 30 s in 95% ethanol and 15 min in 50% commercial Chlorox bleach.

The surface-sterilized seeds were extensively rinsed with sterile tap water, imbibed for 4 to 6 h in sterile water, and germinated in the dark on moist Whatman #1 paper. After the seedlings were harvested, the germination paper was touched to the surface of an LB agar plate to check for microbial contamination. Only uncontaminated seedlings were used for the bioassays.

**Preparation of pea exudates and extracts.**

Batches of approximately 100 aseptically grown, 7-day-old pea seedlings (and the washed, autoclaved Whatman filter paper used to maintain moisture for the seedlings) were flooded with 100 ml of sterile water and gently swirled for several minutes to collect the crude seedling rinsate. One-hundred-microliter samples of the rinsates were plated on LB agar to check for microbial contamination, and contaminated batches were discarded.

Contaminant-free rinsates were spun down, filtered, frozen, and lyophilized. On average, 5 to 10 mg of dry crude exudate was collected per 100 seedlings. Lyophilized exudate preparations were mixed with 50% methanol:water (1 ml per 10 mg of lyophilized material) and centrifuged after 15 min to remove insoluble materials. The 50% methanol extract was used for routine assays of violacein, swarming, and exoenzyme activity, as shown in Figures 2 and 3. For further purification of activities, an additional volume of 100% methanol was added to the 50% methanol extract to bring the extracts to 95% methanol. Isopropanol (1 volume of isopropanol to 4 volumes of 95% methanol:water) was then added to precipitate inactive contaminating materials. After 15 min of incubation, the insoluble material (approximately 25% by mass) was removed by centrifugation. All extraction steps were carried out at room temperature. The organic solvents were removed by rotary evaporation under vacuum at 30°C. The resulting yellow viscous exudate extract was dissolved in 1.0 ml of water per 10 mg of dry residue for the assays in Figures 3 and 4.

**HPLC of pea extract.**

For reverse phase HPLC, 1 ml of a 30% methanol:water solution of the isopropanol-precipitated extract, prepared as described above, containing exudate from about 650 seedlings, was injected onto a water-equilibrated, semi-preparative C18 column (Whatman Partisil 10 ODS-3). The column was eluted for 15 min in water, followed for 45 min with a linear water: acetonitrile gradient at a flow rate of 3 ml/min. Absorbance was monitored at 254 nm. Fractions were collected every minute and bioassayed with the reporters as described below.

**Bioassays for AHL mimic compounds from plants: Violacein and phenazine production.**

*C. violaceum* CV026, *P. aureofaciens* 30-84, and a C6-HSL negative mutant, 30-84I, were grown in LB broth overnight on a shaker at 27°C. Five hundred microliters of the cultures,
containing approximately 2–3 \times 10^6 bacteria, was spun down and resuspended in 5 ml of warm LB containing 0.5% agar. For indirect CV026 bioassays to measure inhibition of violaeclin synthesis by seedlings or exudate, 3.5 µg of C_4-HSL dissolved in 7 µl of ethyl acetate was added to the soft agar suspension of bacteria. For assays with live seedlings (Fig. 1, Table 1), the bioassay agar suspensions were poured on a surface of regular LB agar, and aseptically grown seedlings were placed on the thin layer of soft agar containing the bacteria. As controls, glass fiber disks containing appropriate AHLs were placed on the surface of the bioassay agar. To semi-quantitatively estimate the ability of seedling exudates or exudate extract preparations to inhibit AHL-induced pigment synthesis, 20- to 30-µl samples from a two- or threefold serial dilution of the exudate or extract (with organic solvents removed by evaporation) were added to the wells of a 96-well Falcon microtiter plate and mixed with 50 µl of the warm bioassay agar containing CV026 and C_4-HSL (or P. aureofaciens 30-84 with no AHL) as described above. The plates were incubated for 15 to 25 h at 27°C. Violaeclin was extracted from the wells with 700 µl of 95% ethanol, and absorbance measured at 575 nm. The reversibility of pea exudate-induced inhibition of violaeclin synthesis was tested by adding an additional 0.5 µg of C_4-HSL per ml either 0 or 4 h after mixing of the CV026 bioassay agar with one seedling equivalent of the pea exudate extract and incubation for 20 h.

To test whether compounds in pea root exudates can degrade or immobilize C_4-HSL, HPLC fractions containing violaeclin synthesis inhibiting activity from 25 pea seedlings were re-dissolved in 250 µl of sterile distilled water and mixed with a 50-µl solution containing 25 µg of C_4-HSL in water and incubated overnight. As a control, 300 µl of sterile, distilled water containing 25 µg of C_4-HSL was incubated similarly. C_4-HSL was extracted with ethyl acetate (2 \times 2 ml) and the solvent removed by rotary evaporation. The C_4-HSL residue was re-dissolved in sterile distilled water and samples from a twofold dilution series assayed by the semiquantitative microtiter plate method described above.

**Luminescence bioassays with the CCD camera.**

*S. liquefaciens* PL10 was grown overnight at 27°C in AB with glucose and Casamino Acids (Eberl et al. 1996) with tc10. *E. coli* JM109 pSB401 (LuxRI'), *E. coli* JM109 pSB536 (AhyRI'), and *E. coli* JM109 pSB1075 (LasRI') were grown at 37°C in LB with tc10 or amp100. The bacterial suspensions were then centrifuged, and the pellets resuspended in 10 volumes of sterile water to an A_600 of approximately 0.10. Aseptically grown, etiolated seedlings were completely submerged in the suspension of bacteria, then gently blotted on a piece of sterile, absorbent tissue and incubated on the surface of AB agar at 27°C (for PL10) or LB agar at 37°C (for the plasmid reporters) for 5 to 9 h. As controls, suspensions of the reporter strains were inoculated onto small slabs (3 × 20 × 70 mm) of nutrient agar, incubated, and then observed at the same time as the seedlings. For the positive controls, the agar slabs contained a high, maximally activating concentration of C_4-HSL for the LuxRI' and AhyRI' (2 µM) and PL10 (6 µM) reporters or 3-oxo-C_12-HSL (5 µM) for the LasRI' reporter. No AHL was added to the agar slabs of the negative controls. Images of the seedlings and luminescing bacteria were taken with a Hamamatsu C2400 intensified CCD camera and analyzed with an Argus-20 image processor. The false color images were superimposed on the digital images of the seedlings in Adobe Photoshop version 4.0.1. The total number of photons emitted and counted per second from the entire surface area of the seedling (or corresponding streak colonies for the controls) was determined. The relative luminescence of the reporter strain growing on an inoculated seedling was then calculated by subtracting the background luminescence of the uninoculated reporter strain (growing on agar containing no AHL), and then expressing this net luminescence as a percentage of the luminescence of the positive controls. Relative luminescence values were normalized to equal numbers of reporter bacteria based on recovery of the reporter bacteria from seedling and agar surfaces and plating on agar to determine the number of CFU.

For quantitative bioluminescence bioassays, 20 to 30 µl of seedling exudate extract or HPLC fractions was placed onto 11-mm-diameter Whatman GFC glass fiber disks. After evaporation of the samples to dryness in a laminar flow hood, the disks were inoculated with 10 µl of 10-fold-diluted cultures of the luminescence reporters and then incubated for 5 to 6 h in the shallow wells of a 96-well microtiter plate lid with 25 µl of AB glucose/Casamino Acids medium (PL10) or LB (plasmid reporters) prior to CCD camera measurement of luminescence. Luminescence bioassays with PL10 required exposure to n-decanal vapor (30 µl in a large lidded box for 8 to 20 min) prior to luminescence measurements.

**Luminescence assays with X-ray film.**

For routine, semiquantitative detection of lux-inducing activity induced by pea exudates, extracts, or HPLC fractions, samples (or samples from a twofold dilution series) were inoculated on filter disks in the presence of the lux reporter strains as described above. Luminescence was then detected by placing the microtiter plate lid on a piece of Kodak Bio-Max Light-1 X-ray film for 4 to 8 min. The incubation times required to induce maximal luminescence and maximal net luminescence above background varied for each of the lux reporters, requiring about 4 to 6 h for the LuxRI', AhyRI', and LasRI' reporters and about 9 to 11 h for the PL10 reporter. The luminescence of reporter strains inoculated onto seedlings could be readily detected in the same manner.

**Swarming assays.**

*S. liquefaciens* MG44 and PL10 were grown overnight in AB with Casamino Acids and glucose with amp50 or tc10 at 27°C to a late log phase. The cells were then pelleted, and re-suspended in 10 volumes of sterile water. Aseptically grown seedlings or sterile toothpicks were dipped in the reporter suspensions, rinsed in sterile water, blotted on a sterile, absorbent tissue, and incubated for 10 to 15 h on a surface of AB agar with glucose and Casamino Acids as in Givskov et al. 1997. Swarm colonies were photographed with a Polaroid camera. Swarm assays were also conducted in 24-well microtiter plates by inoculating the surface of agar containing seedling exudate or extract with the MG44 reporter strain followed by overnight incubation.

**Protease assays.**

Late log phase suspensions of *C. violaceum* CV026 (10 µl) were added to 1 ml of fresh 1/10 strength LB medium con-
taining 0.5 µg of C_4-HSL per ml and 0, 1, 5, or 15 seedling equivalents of pea exudate extract, with the organic solvents previously removed by evaporation. After 20 h of shake culture growth at 27°C, the bacteria were removed by centrifugation and the culture supernatants assayed for protease activity with azocasein (Sigma Chemical, St. Louis, MO) as described (Gerhardt et al. 1994). Similarly, C. violaceum wild type was inoculated into 1/10 LB medium with or without pea exudate extract and with or without 3-oxo-C_12-HSL (2.3 µg/ml) and assayed after 20 h incubation for protease activity on azocasein.

**Chitinase assays.**

For exochitinase (p-nitrophenyl-β-D-N-acetylglucosaminidase) activity, 10 µl of CV026 overnight LB shake culture was inoculated into 1/10 strength LB-based synthetic medium (SM) with colloidal chitin prepared as described by Chernin et al. (1998) containing 0.5 µg of C_4-HSL per ml and 0, 1, 5, or 15 seedling equivalents of pea exudate extract. After 20 h of incubation on a shaker at 27°C, exochitinase activity in culture supernatants was measured with p-nitrophenyl-β-D-N-acetylglucosaminide (Sigma) as described in Chernin et al. (1998). Endochitinase activity was assayed in 1/10 LB-based SM agar with colloidal chitin (as in Chernin et al. 1998) containing 0.5 µg of C_4-HSL per ml and 0, 1, 5, or 15 seedling equivalents of a pea exudate extract per 1 ml of the medium. The plates were stab inoculated with CV026 and chitin degradation estimated by clearing of the medium surrounding the bacterial colonies after 3 to 4 days of incubation.

**Phospholipase assay.**

For S. liquefaciens cultures (MG44 and PL10) incubated with pea seedling extracts, phospholipase activity was measured in egg yolk agarose as described by Givskov and Molin 1993.

In general, growth of bacteria was monitored in shake cultures by both turbidimetric measurements at A_{600} and by dilution platings of gently sonicated cultures on LB agar. All assays were done in duplicate and all experiments were conducted at least twice with essentially the same results. The C_4-HSL used in this study was synthesized as previously described (Chhabra et al. 1993) and verified by HPLC fractionation and MS analysis, while the 3-oxo-C_12-HSL was provided by B. Iglewski.

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**LITERATURE CITED**


