Changes in Gene Expression in Canola Roots Induced by ACC-Deaminase-Containing Plant-Growth-Promoting Bacteria

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The technique of RNA arbitrarily primed-polymerase chain reaction (RAP-PCR) was used to study changes in gene expression over time in canola roots treated with the 1-aminocyclopropane-1-carboxylate (ACC) deaminase-containing plant-growth-promoting bacterium Enterobacter cloacae UW4 and to compare the changes with those in a mutant of E. cloacae UW4 in which the ACC deaminase structural gene acdS was replaced by homologous recombination with acdS with an intentional knockout containing a tetracycline resistance gene. Genes that were either up- or down-regulated over a three-day period in canola plants treated with wild-type or mutant bacteria were isolated, cloned, and sequenced; all appeared to have high homology with Arabidopsis thaliana genes. The upregulated genes included a cell division cycle protein 48 homolog and a eukaryotic translation initiation factor 3 subunit 7 gene homolog. The downregulated genes included one encoding a glycine-rich RNA binding protein with a function in RNA processing or binding during ethylene-induced stress, which is expressed only in roots, and another gene thought to be involved in a defense signaling pathway. All RAP-PCR results were verified using Northern blotting. These data, indicate that roots isolated from canola seeds treated with the ACC deaminase-producing E. cloacae UW4 up-regulate genes involved in cell division and proliferation but down-regulate stress genes. This data is in agreement with a model in which ACC deaminase-containing plant-growth-promoting bacteria reduce plant stress and induce root elongation and proliferation in plants, largely by lowering ethylene levels.

Plant-growth-promoting bacteria include a diverse group of free-living soil bacteria that can stimulate the growth of plants by one or more of a number of different mechanisms (Glick 1995). These bacteria may stimulate plant growth directly or indirectly. Indirect stimulation of plant proliferation includes preventing phytopathogens from inhibiting plant growth and development (Holguin and Glick 2003; Robison et al. 2001; Wang et al. 2000); direct stimulation includes providing plants with compounds such as fixed nitrogen, phytohormones, or solubilized iron from the soil (Glick 1995; Glick et al. 1999; Jacobson et al. 1994) and that this enzyme can cleave the ethylene precursor ACC to α-ketobutyrate and ammonia and thereby lower the level of ethylene in developing or stressed plants. When ACC deaminase-containing plant-growth-promoting bacteria are bound to the seed coat or root of a developing seedling, they act as a sink for ACC, ensuring that plant ethylene levels do not become elevated to the point at which root growth is impaired (Glick et al. 1998). In addition, by lowering ethylene levels, ACC deaminase-containing plant-growth-promoting bacteria protect plants from the deleterious effects of numerous environmental stresses, including flooding (Grichko and Glick 2001), phytopathogens (Wang et al. 2000), metals (Burd et al. 1998, 2000; Grichko et al. 2000; Nie et al. 2002), drought (Mayak et al. 2004), and salt (Mayak et al. in press). The importance of the enzyme ACC deaminase in the functioning of the bacterium Enterobacter cloacae UW4 in plant growth promotion was recently demonstrated (Li et al. 2000). The ACC deaminase structural gene (acdS) from E. cloacae UW4 was replaced by homologous recombination with a tetracycline resistance gene inserted within the acdS coding region. Upon characterization of this AcdS minus (AcdS-) mutant, it was determined that both ACC deaminase activity and the ability to promote the elongation of canola roots under gnotobiotic conditions were greatly diminished (Li et al. 2000). This result is consistent with the previously postulated model (Glick et al. 1998) that suggests that a major mechanism utilized by plant-growth-promoting bacteria involves the lowering of plant ethylene levels, and hence, ethylene inhibition of plant proliferation by bacterial ACC deaminase. While it is clear that ACC deaminase-containing plant-growth-promoting bacteria can promote the growth of a variety of plants under a range of conditions, the precise nature of the changes in plant gene expression that are caused by the bacterium remains to be elucidated. As a first step toward understanding how ACC deaminase promotes plant growth, in the present study, we have used the RNA arbitrarily primed PCR (RAP-PCR) technique to identify some ACC deaminase-responsive genes in canola (Brassica rapa) cv. Reward roots.

RESULTS

Effect of ACC deaminase-containing, plant-growth-promoting bacteria on canola root elongation.

Following five days in gnotobiotic growth pouches, canola roots produced from seeds treated with E. cloacae UW4 showed a significant increase in root length when compared with either the negative control or the AcdS- knockout mutant.
The average root length of the *E. cloacae* UW4 treated plants was 98.0 ± 11.3 mm, compared with those treated with *E. cloacae* UW4/AcdS−, which were 61.0 ± 10.3 mm (Fig. 1) and the roots of plants not treated with bacteria, which were 55.5 ± 8.0 mm long. Thus, the roots of the *E. cloacae* UW4–treated canola were, on average, nearly twice as long as either untreated plants or plants treated with the mutant strain, confirming the relationship between ACC deaminase activity and plant root length (Li et al. 2000).

**Products of RAP-PCR.**

The concentration of total RNA extracted from canola roots treated with either wild-type or mutant *E. cloacae* UW4 was measured by spectrophotometry and then visualized on a 1% formaldehyde-aldehyde gel. To identify differentially expressed canola genes associated with the effects of ACC deaminase, RNA from roots treated with either *E. cloacae* UW4 or *E. cloacae* UW4/AcdS− and obtained five days after seed imbibition was used for RAP-PCR (Fig. 2). A total of six differentially expressed bands were successfully isolated and reamplified; four of these are shown in Figure 2. Even though there were other bands in the gel that appeared to be down-regulated, it was not possible to clone them. In order to identify the proteins that are encoded by these cDNA fragments, the bands were isolated and sequenced, and nucleotide sequences of the differentially expressed cDNAs were compared with sequences residing in the GenBank nucleotide and dbEST databases at the National Center for Biotechnology Information (NCBI). Four of the isolated cDNA fragments were similar to known sequences found in these databases, while RAP5 did not correspond to any significant extent to any known gene or protein, and RAP6 corresponded weakly with a probable retroelement polymerase polyprotein (Table 1). Based on amino acid sequence similarity, RAP1 was identified as a glycine-rich RNA-binding stress protein. At the amino acid level, RAP1 shares 68% identity to the stress protein from *Brassica napus* (Bergeron et al. 1993). An alignment of the amino acid sequences of the four known sequences with their nearest counterparts is shown in Figure 3. RAP2 was identified as a cell division cycle (cdc) 48 protein that showed 90% amino acid identity to the protein from *Arabidopsis thaliana*. RAP3 was identified as eukaryotic translation initiation factor 3/subunit 7. It showed a 69% amino acid identity with the comparable protein from *A. thaliana*. RAP4 was identified as a defense signaling pathway gene, which is homologous (70% identity) to one found in *A. thaliana*.

**Table 1. Differentially expressed genes that were identified using RNA arbitrarily primed (RAP)-polymerase chain reaction**

<table>
<thead>
<tr>
<th>Fragment</th>
<th>Accession no.</th>
<th>Estimated size (bp)</th>
<th>Putative function</th>
</tr>
</thead>
<tbody>
<tr>
<td>RAP1 (−)</td>
<td>AY406110</td>
<td>268</td>
<td>Glycine-rich root-specific stress protein</td>
</tr>
<tr>
<td>RAP2 (+)</td>
<td>AY406108</td>
<td>647</td>
<td>Cell division cycle 48 gene</td>
</tr>
<tr>
<td>RAP3 (+)</td>
<td>AY406109</td>
<td>365</td>
<td>Eukaryotic translation initiation factor 3/subunit 7 gene</td>
</tr>
<tr>
<td>RAP4 (−)</td>
<td>AY406111</td>
<td>213</td>
<td>Defense signaling pathway gene</td>
</tr>
<tr>
<td>RAP5 (?)</td>
<td>CK927465</td>
<td>377</td>
<td>Unknown function</td>
</tr>
<tr>
<td>RAP6 (?)</td>
<td>CK937466</td>
<td>512</td>
<td>Probable retroelement polymerase polyprotein</td>
</tr>
</tbody>
</table>

* (+) and (−) indicate genes up- or down-regulated, respectively, in canola treated with 1-aminocyclopropane-1-carboxylate deaminase–containing plant-growth-promoting bacteria; ? indicates not determinable.

![Fig. 1. Gnotobiotic root-length assay. Canola roots after five days of growth with either *Enterobacter cloacae* UW4 or with the *E. cloacae* 1-amino-cyclopropane-1-carboxylate deaminase knockout mutant UW4/AcdS−. Error bars indicate standard error.](image1)

![Fig. 2. Polyacrylamide gel showing RNA arbitrarily primed-polymerase chain reaction amplified cDNA fragments from canola roots. Genes differentially expressed and identified are boxed and labeled. Lanes 1 and 2, Canola roots treated with *Enterobacter cloacae* UW4. Lanes 3 and 4, canola roots treated with *E. cloacae* UW4/AcdS−.](image2)
Fig. 3. Alignment of the amino acid sequences of RAP1, RAP2, RAP3, RAP4, RAP5, and RAP6 with known amino acid sequences. RAP1 was aligned with *Brassica napus*, whereas all others were aligned with *Arabidopsis thaliana*. An asterisk above a sequence indicates differences, and gaps are filled with bars. Score indicates percent identity. Alignment was done using ClustalW version 1.23.
After sequencing and identifying the cDNAs, changes in the level of expression of the corresponding mRNAs due to the presence or absence of ACC deaminase were evaluated by Northern blot analyses. Total RNA from canola roots treated with either *E. cloacae* UW4 or *E. cloacae* UW4/AcdS— and harvested three or five days after seed imbibition was probed with the different cDNA products (i.e., RAP1, RAP2, RAP3, RAP4, RAP5, and RAP6). The expression of both RAP1 and RAP4 was down-regulated (Fig. 4A and B) in canola roots treated with ACC deaminase-containing bacteria, whereas expression of RAP2 and RAP3 was up-regulated (Fig. 4C and D).

**Fig. 4.** Northern blots of genes putatively turned on or off by 1-aminocyclopropane-1-carboxylate deaminase. **a,** Northern blot of the RAP1 gene identified as encoding a glycine-rich root-specific stress protein. **b,** Northern blot of the RAP4 gene identified as encoding a SH2 binding defense signaling pathway gene. **A,** Canola roots were treated with either *E. cloacae* UW4 or UW4/AcdS— and were harvested 3 or 5 days posttreatment. **B,** A stained blot is shown immediately below the Northern blot. RNA (15 µg in each lane) is from canola roots treated with *E. cloacae* UW4/AcdS— harvested at day 3, canola roots treated with *E. cloacae* UW4/AcdS— harvested at day 5, canola roots treated with *E. cloacae* UW4/AcdS— harvested at day 3, and canola roots treated with *E. cloacae* UW4 harvested at day 5. **C,** Northern blot of the RAP2 gene identified as encoding a cell division cycle 48 gene. **D,** Northern blot of the RAP3 gene identified as encoding an eukaryotic translation initiation factor 3 subunit 7 gene. **E,** Northern blot of the RAP5 gene identified as an expressed protein. **F,** Northern blot of the RAP6 gene identified as a probable retroelement polymerase polyprotein.
In addition, the expression of both RAP1 and RAP4 was time dependent. In *E. cloacae UW4/AcdS* treated canola roots, RAP1 mRNA was significantly more abundant in roots that were harvested after three days than those harvested after five days. Likewise, there was a faint signal for RAP1 in ACC deaminase-treated canola roots harvested after three days that was no longer evident in roots harvested after five days. While there are higher levels of RAP2 expressed in *E. cloacae UW4*-treated canola than in plants treated with *E. cloacae UW4/AcdS*, there is no significant difference in the expression of this mRNA between day 3 and day 5 canola roots treated with either *E. cloacae UW4* or *E. cloacae UW4/AcdS*.

Similarly, there is a higher level of expression of RAP3 in *E. cloacae UW4* treated canola than in that treated with *E. cloacae UW4/AcdS*. There is no observable significant difference between day 3 and day 5 roots treated with *E. cloacae UW4*. However, there is a higher level of expression of RAP3 at day 5 in canola treated with *E. cloacae UW4/AcdS* than at day 3. RAP5 showed strong expression in both *E. cloacae UW4* and *E. cloacae UW4/AcdS*, whereas RAP6 expression was very low in both *E. cloacae UW4* and *E. cloacae UW4/AcdS*.

### DISCUSSION

In this study, we have examined some of the effects of ACC deaminase, present within the cytoplasm of a plant-growth-promoting bacterium bound to the roots or seed coat, on gene expression in canola. RAP-PCR was used to compare the patterns of expressed mRNAs present in canola roots treated with ACC deaminase-containing plant-growth-promoting bacteria to those treated with an AcdS* knockout mutant. Several canola genes showed differential expression; however, only four corresponded to known genes. The differential expression of these four genes was subsequently confirmed by Northern blotting. RAP1 was identified as a glycine-rich root-specific stress gene, RAP2 as a cdc 48 gene, RAP3 as an eukaryotic translation initiation factor (eIF) 3 subunit, and RAP4 as a defense signaling pathway gene. Both RAP1 and RAP4 showed time-dependent expression, whereas RAP2 and RAP3 displayed mostly time-independent expression.

The root-specific stress gene encoded by RAP1 was first identified and characterized in canola (*Brassica napus*) by Bergeron and associates (1993). It contains a glycine-rich domain common to many RNA-binding proteins involved in plant stress responses. In addition, this gene is induced by ethylene-producing stresses, such as cold, wounding, flooding, and treatment with 2-chloroethylphosphonic acid, a compound that generates ethylene. Since ACC deaminase-containing plant-growth-promoting bacteria affect plants by lowering their ethylene levels and alleviating some of the deleterious effects of ethylene-induced gene expression, it is not surprising that deletion of the ACC deaminase gene from *E. cloacae UW4* results in a bacterium that no longer protects against the deleterious effects of ethylene. Thus, plants treated with wild-type *E. cloacae UW4* do not need to express a root-specific stress gene that is expressed in plants treated with an *E. cloacae UW4* mutant that does not have ACC deaminase activity. This result suggests that the ACC deaminase-containing *E. cloacae UW4* helps to protect the seed and root system from the potentially deleterious effects of ethylene. Even though some ethylene is necessary for seed germination, higher levels of ethylene inhibit root growth. The addition of ACC deaminase-containing *E. cloacae UW4* relieves some of the inhibitory effects of ethylene, observed in the significantly larger root lengths from canola (*Brassica rapa* cv. Reward) treated with these bacteria. This result is also in agreement with a model (Glick et al. 1998) that proposes the importance of the bacterial enzyme ACC deaminase in plant stress alleviation and, in addition, provides evidence that ACC deaminase-containing plant-growth-promoting bacteria down-regulate the expression of stress genes linked to the ethylene stress response.

Timmusk and Wagner (1999) previously showed that plants treated with the plant-growth-promoting bacterium *Pseudomonas polymyxa* and then exposed to either biotic or abiotic stress exhibited a strong inoculation-dependent increase in abundance in one particular mRNA. The corresponding gene was identified as *ERD15*, which was previously shown to be responsive to drought stress. Quantification of mRNA levels of several stress responsive genes indicated that *P. polymyxa* itself induced mild biotic stress. These authors suggested that biotic or abiotic stress could induce genes associated with plant defenses. Unlike treatment of a plant with *P. polymyxa*, a bacterium that does not show any significant ACC deaminase activity, treatment of canola with the ACC deaminase-containing plant-growth-promoting bacterium *E. cloacae UW4* does not lead to an increase in other stress or a defense signaling pathway gene. For example, in the present study, we identified a defense signaling pathway gene that is down-regulated in ACC deaminase treated canola. In contrast, canola treated with the *acdS* knockout mutant show significant expression of this defense signaling pathway gene.

In addition to genes that are down-regulated, cdc 48 and eIF 3 genes are up-regulated in canola treated with an ACC deaminase-containing plant-growth-promoting bacterium, as compared with canola treated with the same bacterium containing an AcdS* knockout mutant. eIF 3 is the largest of the eIFs (Asano et al. 1997). It is a multiprotein complex of about 600 kDa and is necessary for protein production. The upregulation of this gene in canola treated with an ACC deaminase-containing plant-growth-promoting bacterium can be explained by the fact that, when canola seeds are treated with *E. cloacae UW4*, the observed average root lengths after five days of growth are almost twice as long as when canola is treated with *E. cloacae UW4/AcdS*. To achieve this additional root growth, more protein has to be produced; hence, it is reasonable to expect upregulation of one of the major transcripts required for plant root protein production. Similarly, the cdc 48 gene, which was previously isolated from *Arabidopsis thaliana* in which it was shown to be essential for cell division and growth, is highly expressed in the proliferating cells of the vegetative shoot and root (Feiler et al. 1995). In addition, it was recently demonstrated that certain *Arabidopsis* gene families, including ATPases and cell division cycle proteins, are involved in plant nutrition and root uptake systems and are necessary for plant survival (Maathuis et al. 2003). For *E. cloacae UW4*-treated roots to be twice as long as roots treated with *E. cloacae UW4/AcdS*, the root cells require the upregulation of genes that are involved in root cell growth and proliferation.

By using the technique of RAP-PCR, we have demonstrated that, at the genetic level, the ACC deaminase-containing plant-growth-promoting bacterium *Enterobacter cloacae UW4* down-regulates a gene involved in ethylene-induced plant stress and up-regulates genes involved in plant growth. In addition, contrary to some plant-growth-promoting bacteria that do not contain ACC deaminase, canola treated with *E. cloacae UW4* does not show an upregulation in a defense signaling pathway gene. This data lends further support to the previously proposed model that attempts to explain how ACC deaminase-containing plant-growth-promoting bacteria facilitate the growth of plants (Glick et al. 1998).
MATERIALS AND METHODS

**Bacterial strains.**

*Enterobacter cloacae* UW4 was isolated from the rhizosphere of reeds growing on the campus of the University of Waterloo based on its ability to utilize ACC as a sole source of nitrogen and to promote the growth of canola seedlings under nonotobiotic conditions (Glick et al. 1995). *E. cloacae* UW4 was subsequently changed to *E. cloacae* UW4 based on fatty acid analysis characterization of the bacterium (Shah et al. 1997). *E. cloacae* UW4/AcdS is a genetic construct in which a tetracycline resistance gene was inserted into the coding region of the bacterial ACC deaminase gene (Li et al. 2000).

**Growth pouch assay.**

The protocol for assessing the effect of bacterial strains on the elongation of canola seedling roots followed a modification (Penrose and Glick 2003) of the method described by Lifshitz and associates (1987). Single colonies of bacterial strains were cultured in 5 ml of tryptic soy broth (TSB) medium (supplemented with 15 µg/ml tetracycline for transconjugants) at 30°C for 24 h with shaking (200 rpm). The bacterial cells were then centrifuged at 5,000 × g at room temperature for 10 min. The bacterial pellet was washed twice with 5 ml of 0.85% NaCl before being resuspended in 0.85% NaCl and adjusted to an absorbance of 0.3 to 0.4 at 600 nm, prior to incubation with seeds.

Canola seeds (*Brassica rapa* cv. Reward), provided by G. Brown (Agrium, Inc., Saskatoon, Saskatchewan, Canada) and previously stored at 4°C, were surface-sterilized by soaking them in 1.5% sodium hypochlorite sodium (bleach) for 10 min. The seeds were subsequently rinsed with sterile distilled H2O five times and then incubated in either 5 ml of 0.85% NaCl, which acted as a blank control, or in the appropriate bacterial suspensions in 0.85% NaCl for 1 h at room temperature (22°C ± 1°C). The seeds were then put into growth pouches, as described by Penrose and Glick (2003). After five days, primary root lengths were measured and analyzed using a one-way analysis of variance in Instat version 2.01 for Macintosh (GraphPad Software, San Diego, CA, U.S.A.). Harvested roots and shoots were frozen in liquid nitrogen and stored at −80°C until further analysis. Seeds that failed to germinate in 2 days after sowing were marked, and roots that subsequently developed from these seeds were not measured.

**RNA extraction.**

Frozen roots were ground, using a mortar and pestle, to a fine powder in liquid nitrogen. Total RNA for RAP-PCR and Northern blot hybridization was extracted using the plant RNAeasy system (Qiagen, Mississauga, Ontario, Canada), following the manufacturer’s instructions.

**RAP-PCR.**

RAP-PCR was carried out according to McClelland and Welsh (1994). A RAP-PCR primer set was obtained from Sigma Genosys (Oakville, Ontario, Canada). Each reverse transcription reaction contained 3 µg of total RNA in a volume of 13 µl of water. Two primers (1 µl of each), primer A1 (5′-AATCTAGAGCTCTCC-3′) and primer A2 (5′-AATCTA GACCTCTTCCTGG-3′), each diluted to 25 µM, were added to the sample. The mixture was heated to 70°C for 10 min and subsequently chilled on ice. For reverse transcription, a master mix was made containing 4.5 µl of 5× M-Muloney murine leukemia virus (MuLV) buffer (500 mM Tris-HCl, pH 8.3, 0.1 M NaCl, 1 mM EDTA, 5 mM dithiothreitol, 0.1% Triton X-100, 50% glycerol), 1 µl of each of the four dNTPs (25 µM), 1 µl RNase inhibitor, and 1 µl M-MuLV V reverse transcriptase (5 units/µl). All reagents were obtained from Fermentas, Inc. (Oakville, Ontario, Canada). The reverse transcription reaction was carried out at 37°C for 1 h, followed by inactivation of the transcriptase by heating at 95°C for 5 min. The cDNA obtained was diluted 1:10 with water, and to 5 µl of the diluted cDNA, 1 µl of primer A1 (25 µM) and 1 µl of primer A2 (25 µM), 1 µl of dNTPs (25 µM), 1 µl of Taq polymerase (5 units), 5 µl of Taq polymerase buffer (×10) was added. All reagents were obtained from Fermentas, Inc. RAP-PCR was carried out using the following conditions: denaturation at 94°C for 1 min, annealing at 36°C for 5 min, extension at 72°C for 5 min in an MJ Instruments PTC-100 thermocycler (Waltham, MA, U.S.A.). This was followed by 39 cycles of denaturation at 94°C for 1 min, annealing at 35°C for 2 min, and extension at 72°C for 2 min. Subsequently, the cDNA products were labeled with Cy-5 primer (5′-AATCTAGAGCTC-3′) by adding 1 unit of exo− Klenow fragment (Fermentas) for 2 h at 37°C. The labeled products were subsequently run in a vertical DNA 1.4% polyacrylamide sequencing gel for 4 h at 2,000 V. The gel was visualized using a Storm 860 optical scanner (Molecular Dynamics, Sunnyvale, CA, U.S.A.); differentially expressed bands were excised from the gel, and the cDNA products were gel-isolated using the Qiaxx II kit from Qiagen (Oakville, Ontario, Canada).

**Cloning, sequencing and analyses.**

cDNA products were cloned into the pGEM-T easy vector system from Promega (Mississauga, Ontario, Canada). The cloned partial cDNA sequences were sequenced using the dye-terminator cycle sequencing method on an ABI Prism model 377 system (Applied Biosystems, Foster City, CA, U.S.A.). The nucleotide sequences obtained were submitted to the NCBI server for BLASTN and BLASTX searches against nucleotide and protein sequences available in various databases.

**RNA blot hybridization.**

Total RNA (20 µg) was size-separated on a formaldehyde denaturing 1.2% agarose gel according to Sambrook and associates (1989). RNA was capillar transferred to a positively charged nylon membrane (Roche, Quebec, Canada), using 20x SSC (1x SSC is 0.15 M NaCl plus 0.015 M sodium citrate) as the transfer medium. RNA was fixed to the membrane by UV-crosslinking for 3 min (UV Stratalinker 2400; Stratagene, La Jolla, CA, U.S.A.). Membranes were probed using cDNA probes labeled with α-32P-dCTP (Amersham, Toronto). Membranes were probed overnight and afterwards washed two times at room temperature for 15 min and then one time at 65°C for 30 min with 5x SSC and 0.1% sodium dodecyl sulfate (SDS). The washed membrane was visualized using a Storm 860 optical scanner (Molecular Dynamics).

For subsequent probing, the membrane was stripped of bound probe by boiling three times in 0.1% SDS, for 15 min and then reused.

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**LITERATURE CITED**


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