Identification and Transcriptional Analysis of Priming Genes in *Arabidopsis thaliana* Induced by Root Colonization with *Pseudomonas chlororaphis* O6

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(Received on May 11, 2011; Revised on July 21, 2011; Accepted on July 21, 2011)

Root colonization of *Arabidopsis thaliana* with *Pseudomonas chlororaphis* O6 induces systemic tolerance against diverse pathogens, as well as drought and salt stresses. In this study, we demonstrated that 11 genes in the leaves were up-regulated, and 5 genes were down-regulated as the result of three- to five-days root colonization by *P. chlororaphis* O6. The identified priming genes were involved in cell signaling, transcription, protein synthesis, and degradation. In addition, expression of selected priming genes were induced in *P. chlororaphis* O6-colonized plants subjected to water withholding. Genes encoding defense proteins in signaling pathways regulated by jasmonic acid and ethylene, such as VSP1 and PDF1.2, were additional genes with enhanced expression in the *P. chlororaphis* O6-colonized plants. This study indicated that the expression of priming genes, as well as genes involved in jasmonic acid- and ethylene-regulated genes may play an important role in the systemic induction of both abiotic and biotic stress due to root colonization by *P. chlororaphis* O6.

**Keywords**: disease defense, drought stress, induced systemic resistance, priming

Induced resistance or tolerance against abiotic and biotic stresses is stimulated in plants by two systems. In systemic acquired resistance (SAR), plants develop a broad spectrum of systemic resistance against plant diseases upon pathogen attack. SAR involves elevated salicylic acid (SA) levels and both local and systemic accumulation of pathogenesis-related (PR) proteins (Van Loon et al., 2006). Expression of PR genes, PR-1 in particular, is used as a marker for SAR induction (Durrant and Dong, 2004) and a key regulatory protein is NPR1 (Pieterse and Van Loon, 2004). Acinbenzolar-S-methyl is a chemical that activates the SAR pathway (Bostock, 2005).

Colonization of roots by microbes also elicits systemic resistance against abiotic stresses and a broad spectrum of diseases caused by bacterial, fungal, and viral pathogens (Kim et al., 2011). Induced systemic resistance (ISR) causes the plant to induce different array of genes than SAR. One rhizobacterium, *Pseudomonas chlororaphis* O6, induces ISR for many plant diseases and enhanced drought tolerance (Han et al., 2006; Kim et al., 2004a; Ryu et al., 2007; Spencer et al., 2003). Three days of root colonization by *P. chlororaphis* O6 reduces opening of leaf stomata (Cho et al., 2008), thereby demonstrating a rapidly induced systemic effect.

Enhanced plant gene activation was observed in oilseed rape treated with the ISR-inducers *P. thivervalensis* MLG45 and *Bacillus amyloliquefaciens* (Carieaux et al., 2003; Sarosh et al., 2009), in response to abiotic stress and after challenge by insects (Conrath et al., 2002). Differential expressed cucumber genes were induced in *P. chlororaphis* O6-colonized cucumber after challenge with the fungal pathogen *Corynespora cassicola* (Kim et al., 2004a). One induced plant gene encoding galactinol synthase (CsGolS1), when over expressed in tobacco, increases galactinol levels, enhances disease resistance against *Botrytis cinerea* and *Pectobacterium carotovorum*, and provides drought and salinity tolerance (Kim et al., 2008). Another study of plants inoculated with the ISR-inducer *P. fluorescens* WCS 417r, showed 81 genes to increase in expression after challenge inoculation with the bacterial speck pathogen *P. syringae pv. tomato* (Verhagen et al., 2004). Few changes in gene expression were noted in plants colonized by *P. fluorescens* WCS 417r without pathogen challenge (Van Wees et al., 1999; Verhagen et al., 2004). This enhanced gene activation in the colonized plants by ISR-inducing rhizobacteria is referred to as priming. However, the plant
genes that prime for induced defense are not known so far.

The main objective of this study was to identify and characterize genes involved in priming for ISR induced by root colonization with *P. chlororaphis* O6. The priming genes in *Arabidopsis thaliana* were identified using a differential gene expression technique in plants at three and five days after root colonization by *P. chlororaphis*. In addition, the effects of water withholding on abundance of the identified priming genes were examined. For comparison the effect of root colonization on induction of known defense signal pathway marker genes were studied. This study indicated that root colonization of *Arabidopsis* by *P. chlororaphis* O6 stimulated an array of priming genes, and confirmed increased expression of defense genes regulated by the jasmonic acid- and ethylene-signaling pathways.

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**Materials and Methods**

**Plant materials and growth conditions.** The parental *A. thaliana* ecotype Columbia (Col-0) was obtained from the Ohio State University Stock Center, Ohio State University, Columbus, OH, USA. The plants were grown in a soil-less mixture. All seeds were surface-sterilized by 2 minutes of soaking in 70% ethanol, followed by 20 minutes of soaking in 1% sodium hypochlorite. The washed and sterile seeds were then planted in a sterile soil-less medium (peat moss: vermiculite: perlite, 7:3:3 v/v) using 500 cm³ of the medium in a 10.5 × 10.5 × 9 cm pot, with 15–30 seeds per pot. These seedlings were watered with 20 ml of sterile water per pot every two days. The seedlings were grown with a 16-h-light/8-h-dark cycle under 40-W fluorescent lights maintained at 22 ± 1°C with a relative humidity of 50–60%.

**Bacterial strains and culture conditions.** The root colonizing bacterium, *P. chlororaphis* O6 (Spencer et al., 2003), was stored at −80°C in Luria-Bertani (LB) broth containing 20% glycerol. To prepare the inoculum, cells were grown overnight in Kings medium B broth, pelleted by centrifugation at 10,000 × g for 10 min, washed once with sterile water, and suspended in sterile water to a concentration of 1 × 10⁸ colony forming units (cfu)/ml adjusted OD600 nm = 0.1.

**Assessment of timing of onset and longevity of ISR.** The roots of two week-old pot-grown *Arabidopsis* seedlings were inoculated by the addition of 35 ml of suspensions of *P. chlororaphis* O6 at 1 × 10⁸ cfu/ml. Sterilized water was applied to the roots of the control plants. At defined times after root treatment, the *Arabidopsis* plants were challenged with *Pectobacterium carotovorum* subsp. *carotovorum* strain SCC 1 or water as a negative control onto a leaf as described previously (Han et al., 2006).

**Assessment of differential gene expression.** Plant genes that were differentially expressed in the leaves after rhizosphere colonization by *P. chlororaphis* O6 were detected by a PCR-based method using annealing controlled primers (ACPs). Total RNAs were prepared from the entire leaves of three pot-grown seedlings harvested three and five days after either the inoculation of the roots with *P. chlororaphis* O6, or simple treatment with water. The leaves were immediately frozen in liquid nitrogen, and total RNAs were extracted with an RNeasy plant mini kit (cat. no. 74904; QIAGEN). First-strand cDNA was generated by reverse transcription. First-strand cDNAs were diluted by the addition of ultra-purified water, then stored at −20°C until being used in a GeneFishing™ (Seegene, Seoul, South Korea) ACP-PCR system (Kim et al., 2004b). ACP-PCR was conducted at 50°C for one cycle in a final reaction volume of 20 µl containing 3–5 µl (about 50 ng) of diluted first-strand cDNA, 1 µl dT-ACP2 (10 µM), 1 µl 10 µM arbitrary ACP, and 10 µl #2 Master Mix (Seegene, Seoul, South Korea). The amplified PCR products were separated in 2% agarose gel stained with ethidium bromide. The differentially expressed bands in plant samples after root-colonization of *P. chlororaphis* O6 were re-amplified and extracted from the gel using a GENCLEAN II Kit (Q-BIO gene, Carlsbad, CA), and directly sequenced with an ABI PRISM 3100-Avant Genetic Analyzer (Applied Biosystems, Foster City, CA).

**Assessment of transcript level using quantitative reverse transcription polymerase chain reaction (qRT-PCR).** For quantitative differential expression of the selected genes (DEG), RT-PCR analysis was performed using with a QuantiTect SYBR green reverse transcription kit (cat. no.204243; QIAGEN Inc., Valencia, CA) was employed and total RNAs were extracted from leaves of control and ISR-induced plants as described above section. Briefly, the roots of two week-old pot-grown *Arabidopsis* seedlings were inoculated by the addition of 35 ml of suspensions of *P. chlororaphis* O6 at 1 × 10⁸ cfu/ml. Sterilized water was applied to the roots of the control plants. Seven days after root treatment, plants grown under pot conditions were exposed to drought stress by withholding water for defined times. Primers for five of the ACP-PCR-detected genes were designed to yield PCR products of lengths between 150–200 bp. The primer pairs employed in the RT-PCR studies are provided in Supplemental Table 1. The controls consisted of the omission of one of the primers and analysis of the RNA without reverse transcription. Because the transcript levels of the *Arabidopsis* β-actin gene were not
altered under the conditions employed herein, these transcript levels were used to normalize the results for each gene. The comparative CT method \(2^{-\Delta\Delta CT}\) method was used to determine the expression level of analyzed genes (Livak and Schmittgen, 2001). Fold units of change in expression were calculated by dividing the normalized expression for genes in plants colonized with \textit{P. chlororaphis} O6 by the normalized expression values for the genes in the untreated controls. The results are expressed as the means and standard deviations of three replicates.

### Data analysis
Data were analyzed by ANOVA using IBM SPSS Statistics version 19 (IBM Corporation, Somers, NY, USA). The significance of the effects of \textit{P. chlororaphis} O6 pretreatment on the target genes at different time points was evaluated by Duncan’s multiple range test \((P < 0.05)\).

### Results

#### ISR induction by root colonization with \textit{P. chlororaphis} O6
The induction of ISR for a soft rot bacterial pathogen was observed at three days after root-colonization of \textit{P. chlororaphis} O6 and reached maximal level after seven days of the root colonization with O6 strain. The ISR activity was maintained at this level up to 15 days after \textit{P. chlororaphis} O6 treatment, and then gradually declined over time (Fig. 1).

#### Identification of ISR-specific genes
The use of 120 random ACP primer pairs revealed 16 potential genes with different product intensity from RNA pooled from isolations from plants three and five days after root colonization with \textit{P. chlororaphis} O6 as compared with control plants that lacked this inoculation (Fig. 2A and 2B). The transcript levels of the \textit{Arabidopsis} \(\beta\)-actin gene did not

### Table 1. \textit{Arabidopsis} genes detected by ACP-based PCR as being differentially regulated in plant root colonized with \textit{Pseudomonas chlororaphis} O6

<table>
<thead>
<tr>
<th>Functional category</th>
<th>Representitive gene name</th>
<th>GenBank Accession number</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Up-regulated</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IAA synthesis</td>
<td>Nitrilase 1 (NIT1)(At3g44310)</td>
<td>NM_180680</td>
</tr>
<tr>
<td>Stress response</td>
<td>Cold-responsive protein (cor15a)(At2g42540)</td>
<td>NM_180040</td>
</tr>
<tr>
<td></td>
<td>Developing seed related cDNA clone (M77F05STM)</td>
<td>BE530495</td>
</tr>
<tr>
<td>RNA-binding protein</td>
<td>mRNA-binding protein, putative (At3g63140)</td>
<td>NM_116179</td>
</tr>
<tr>
<td>Transcription</td>
<td>(C3HC4-type RING finger) family protein (At5g63780)</td>
<td>NM_125773.2</td>
</tr>
<tr>
<td>Protein synthesis</td>
<td>40S ribosomal protein S7 (RPS7B) (At3g02560)</td>
<td>NM_111124.2</td>
</tr>
<tr>
<td>Photosynthesis</td>
<td>Chlorophyll A-B binding protein 4 precursor homolog (At3g47470)</td>
<td>AY093080.1</td>
</tr>
<tr>
<td>Protein degradation</td>
<td>Cysteine proteinase (RD21A)/thiol protease (At1g47128)</td>
<td>NM_103612.2</td>
</tr>
<tr>
<td>Signaling</td>
<td>KCBP interacting Ca(^{2+})-binding protein (KIC) mRNA</td>
<td>AY363866.1</td>
</tr>
<tr>
<td>Unknown</td>
<td>Expressed protein (At5g19240)</td>
<td>NM_121929.2</td>
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<td></td>
<td>At5g19240 gene</td>
<td>BT004589.1</td>
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<td>Down-regulated</td>
<td>Developing seed \textit{Arabidopsis thaliana} cDNA clone</td>
<td>BE530495</td>
</tr>
<tr>
<td>Chlorophyll synthesis</td>
<td>Photochlorophyllide reductase precursor (At4g27440)</td>
<td>AY081465.1</td>
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<td>Photosynthesis</td>
<td>Lhcb4.2 protein</td>
<td>AF134127.1</td>
</tr>
<tr>
<td></td>
<td>Protodermal factor 1 (PDF1) (At2g42840)</td>
<td>NM_129845.2</td>
</tr>
<tr>
<td></td>
<td>Acyl carrier family protein/ACP family protein (At4g25050)</td>
<td>NM_118637.2</td>
</tr>
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**Fig. 1.** Sustained ISR in \textit{A. thaliana} upon root colonization with \textit{P. chlororaphis} O6. Seedlings grown in pots were inoculated with \textit{P. chlororaphis} O6 (O6) or water as a control (Control). At defined times after root treatments, leaves were challenged with the soft-rot, pathogen \textit{Pectobacterium carotovorum} SCC1 and disease was assessed 2 days later. Three independent experiments were conducted with 3 plants/treatment. Standard errors in the assessment of the extent of disease are shown.
show any major changes with these conditions (Fig. 2B), thus making it appropriate to use in normalization of expression. Table 1 shows the nature of the genes showing differential expression based on their deduced peptide moieties. Genes with increased and decreased expression were noted and they encoded a variety of functions in the plant cell. The stress responsive genes encoded proteins associated with the activity of plant growth regulators [IAA (NIT1) and ABA (Atcor15a)], the salt stress-associated protease (RD21a), a ring finger protein (At5g63780), and a calcium binding protein (KIC) were selected for further investigated. Down-regulated genes encoded a protochlorophyllide reductase precursor (At4g27440), Lhcb4.2 protein, protodermal factor 1 (PDF1), and an acyl carrier protein/ACP family protein.

**Confirmation of ACP Data by Real-Time RT-PCR.** Examination of transcript levels from five of these potential priming genes confirmed their expression as a consequence of colonization by *P. chlororaphis* O6. Expression of the selected DEG genes showed unique transcript patterns. For the gene encoding NIT1, maximum expression was observed five days after root colonization whereas for the other genes the maximum response occurred later at nine days (Fig. 3A).

**Effect of drought stress on expression of genes primed by root colonization by *P. chlororaphis* O6.** We examined whether the selected genes were regulated by drought and whether root colonization by *P. chlororaphis* O6 exerted a priming effect on their expression upon the abiotic stress of water withholding (Fig. 3B). Water withholding of plants with roots colonized by *P. chlororaphis* O6, resulted significant increases in transcripts from four of the genes over drought-stressed control plant, with exception of the gene encoding KIC. In colonized plants expression was
increased more at 3 and 7 days after drought treatment for the \textit{NIT1}, \textit{Zinc finger}, and \textit{RD21a} genes, and at 5 days for \textit{Atcor15A}. Expression of the \textit{KIC} gene showed a trend for reduced expression in colonized plants subjected to water-withholding (Fig. 3B). These results indicated that specific genes showed a priming effect for \textit{P. chlororaphis} \textit{O6}-induced drought tolerance.

\textbf{Discussion}

Root colonization of \textit{A. thaliana} with \textit{P. chlororaphis} \textit{O6} alters plant performance, improving plant growth (Ryu et al., 2007), resilience to pathogenic attack (Spencer et al., 2003) and drought and salt tolerance (Cho et al., 2008). However, molecular mechanisms involved beneficial phenomenon in plants primed by \textit{P. chlororaphis} \textit{O6} awaits resolution. In this study, we conclusively demonstrated that the root inoculation of \textit{Arabidopsis} with \textit{P. chlororaphis} \textit{O6} without other stresses altered gene expression both positively and negatively early after root inoculation. Both reduction in stomatal apertures, important in drought stress tolerance, and the onset of ISR required three days after root inoculation (Cho et al., 2008). Thus, root inoculation by \textit{P. chlororaphis} \textit{O6} rapidly induced systemic effects at the cellular and gene expression levels.

Five up-regulated priming genes were related to abiotic stresses and plant growth. The functions of \textit{NIT1} and \textit{KIC} may be correlated with the improved growth, observed with \textit{P. chlororaphis} \textit{O6} root colonization of tobacco (Kang et al., 2006; Ryu et al., 2007). \textit{NIT1} (At3g44310) encodes nitrilase 1, which releases IAA from indole-3-acetonitrile (Bartel and Fink, 1994; Bartling et al., 1992). Zhang et al., (2007) previously reported increased expression of genes encoding for nitrilases 1, 2 and 3 as the result of treatment of \textit{Arabidopsis} with volatiles from a plant growth-promoting isolate of \textit{Bacillus subtilis}. These volatiles included butanediol, which also is produced by \textit{P. chlororaphis} \textit{O6} (Han et al., 2006). The nitrilase genes, along with an expansin gene, were up-regulated more markedly in the leaves than in the roots, thus suggesting that they are involved in the enhanced leaf growth observed by \textit{P. chlororaphis} \textit{O6} in normal grown plants significantly increased transcript accumulation of defense marker genes encoding \textit{PDF1.2} and \textit{VSP1} associated with the jasmonic acid and ethylene pathway. Transcripts from the ethylene sensitive \textit{HEL} were slightly increased upon root colonization of \textit{P. chlororaphis} \textit{O6} at later time points (Fig. 4A). However, transcription of the SAR marker gene \textit{PR-1a} was slightly decreased.

Significant increase in \textit{VSP1} expression was observed in water limited plants pretreated with \textit{P. chlororaphis} \textit{O6} over drought-stressed control plant (Fig. 4B). Similarly transcripts levels for \textit{PDF1.2} and \textit{HEL} were slightly increased to be maximum 9 days after water withholding. However, no significant increasing of \textit{PR-1a} transcript was observed (Fig. 4B). These results indicated that ethylene and jasmonic acid signal pathway played an important role in the priming effect of \textit{P. chlororaphis} \textit{O6}-induced drought tolerance.
in plants with roots colonized by both the *Bacillus* and the *Pseudomonas* strains.

Transcript from gene (At2g46600), encoded KIC, a Ca\(^{2+}\)-binding protein, connected with cell division, (Bowser and Reddy, 1997; Reddy et al., 2004; Smirnova et al., 1998; Vos et al., 2000). The control of Ca\(^{2+}\) flux by such binding proteins may be associated with regulation of stomatal apertures (Suhita et al., 2004), which were modified by root colonization *P. chlororaphis* O6 (Cho et al., 2008). Previous reports indicated the gene encoding Atcor15a to be drought and cold ABA-regulated (Lin and Thomashow, 1992; Wilhelm and Thomashow, 1993). *RD21a* encodes a cell wall-associated cysteine proteinase that is transcriptionally regulated by high salt (Koizumi et al., 1993; Yamaguchi-Shinozaki et al., 1992). Consequently, we propose that transcriptional induction of *RD21a* and *Atcor15a* upon root colonization with *P. chlororaphis* O6 could be involved in the abiotic stress tolerance induced by *P. chlororaphis* O6-colonization (Cho et al., 2008).

Our studies in tobacco (Spencer et al., 2003) demonstrated that induced resistance was correlated with increased expression of the genes *PR-1g* (Eyal et al., 1992), *HMREG* (Choi et al., 1994) and *Lox* (Heitz et al., 1997), thereby indicating the activation of the defense genes regulated by the ethylene- and jasmonate-regulated pathways. Other studies with *Arabidopsis* defense signaling mutants were verified that these pathways were required for induced resistance to pathogens (Cho et al., 2008). Our data in this paper confirmed the importance of a jasmonate-related pathway through the strong induction in plants colonized with *P. chlororaphis* O6 for the *PDF1.2* and *PDF1.2* genes. Although we have measured increased levels of SA in plants showing stomatal closure (Cho et al., 2008) expression from the SA-responsive gene, *PR-1a*, was not elevated in the colonized plants up to 9 days of inoculation, perhaps as the result of antagonism between the SA and JA-pathways (Thaler et al., 2002).

Our findings illustrate that root colonization induced cross protection for both resistance to plant pathogens and tolerance to abiotic stresses. Previously tolerance to pathogens and high salt, drought, and low temperatures was found upon overexpression of genes for endochitinase and basic PR1 in pepper (Hong et al., 2005). These finding indicated that genes related to pathogen defense may play important roles also in induced tolerance to abiotic stresses, such salt and drought. Rhizobacteria-mediated ISR for abiotic and biotic stresses in plants seems to share common defense signal pathways as demonstrated in this paper. In this study, *P. chlororaphis* O6 colonization primed certain genes for enhanced expression when plants were subjected to the abiotic stress of water withholding. Priming effects were noted for the drought-associated genes, *Atcor15a*, *RD21a*, *Zinc finger*, and *NIT1*. A reverse situation, repression of expression, was noted with *KIC*. However, early priming was detected only with *VSP1* when the defense genes were evaluated. Later priming was observed on days 9 of water withholding for *PDF1.2*. Thus, priming occurred at different times for different genes indicating complexity of regulation.

In summary, root colonization by *P. chlororaphis* O6 primed expression of an array of genes to show increased expression upon drought stress. To our knowledge, this is the first report on identification of plant priming genes for ISR. The priming genes are associated with the cell signaling pathways regulated by ABA, jasmonic acid, and ethylene. Our findings are consistent with the fact that *Arabidopsis* mutants lacking in the jasmonate signaling pathways display neither induced drought resistant nor ISR in plants colonized by *P. chlororaphis* O6 (Cho et al., unpublished). Thus, our work demonstrated that resilience to both abiotic stress and biotic challenge can result from enhanced systemic activation through root colonization by certain microbes.

**Acknowledgements**

This work was supported by a grant from BioGreen 21 program (code# 2007041034032), Rural Development Administration, and supported by the World Class University project of the National Research Foundation of Korea (grant no. R32-2009-000-20047-0).

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