Systemic Induction of the Small Antibacterial Compound in the Leaf Exudate During Benzothiadiazole-elicited Systemic Acquired Resistance in Pepper

Boyoung Lee1,2*, Yong-Soon Park1,2, Hwe-Su Yi2 and Choong-Min Ryu1,2*
1Biosystems and Bioengineering Program, University of Science and Technology, Daejeon 305-806, Korea
2Molecular Phytopathology Laboratory, System and Synthetic Biology Research Center, Korea Research Institute of Bioscience and Biotechnology (KIRIBB), Daejeon 305-806, Korea
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Plants protect themselves from diverse potential pathogens by induction of the immune systems such as systemic acquired resistance (SAR). Most bacterial plant pathogens thrive in the intercellular space (apoplast) of plant tissues and cause symptoms. The apoplastic leaf exudate (LE) is believed to contain nutrients to provide food resource for phytopathogenic bacteria to survive and to bring harmful phytochemicals to protect plants against bacterial pathogens. In this study, we employed the pepper-Xanthomonas axonopodis system to assess whether apoplastic fluid from LE in pepper affects the fitness of X. axonopodis during the induction of SAR. The LE was extracted from pepper leaves 7 days after soil drench-application of a chemical trigger, benzothiadiazole (BTH). Elicitation of plant immunity was confirmed by significant up-regulation of four genes, CaPR1, CaPR4, CaPR9, and CaCHI2, by BTH treatment. Bacterial fitness was evaluated by measuring growth rate during cultivation with LE from BTH- or water-treated leaves. LE from BTH-treatment significantly inhibited bacterial growth when compared to that from the water-treated control. The antibacterial activity of LE from BTH-treated samples was not affected by heating at 100°C for 30 min. Although the antibacterial molecules were not precisely identified, the data suggest that small (less than 5 kDa), heat-stable compound(s) that are present in BTH-induced LE directly attenuate bacterial growth during the elicitation of plant immunity.

Keywords: benzothiadiazole, leaf exudate, pepper, systemic acquired resistance, Xanthomonas axonopodis

Plants protect themselves against virulent and avirulent pathogens and microbes while diverse microbes are exposed in nature (Schneider et al., 1996). The well-characterized and most active plant defense system is the hypersensitive response (HR), which induces rapid programmed cell death at a localized infection site in response to perception of necrotizing non-host pathogens (Mysore and Ryu, 2004). Following HR, systemic acquired resistance (SAR) occurs throughout the whole plant body. In agriculture, SAR has been recognized as a strategy to control plant pathogens because of its evolutionary stability and long-lasting effectiveness against a broad range of pathogens (Deverall, 1995; Hammerschmidt and Kue, 1995).

A number of chemicals including salicylic acid (SA), methyl jasmonate, β-aminobutyric acid (BABA), and benzo-(1,2,3)-thiadiazole-7-carbothioic acid S-methyl ester (BTH) have been shown to induce SAR in plants (Chen et al., 1993; Davis and Ausubel, 1989; Epple et al., 1997; Vallad and Robert, 2004). Among those, BTH (Friedrich et al., 1996; Lawton et al., 1996) and Probenazole (Yoshioka et al., 2001) are commercial chemical inducers for SAR in plants. The two chemicals have been commercialized under the trade names, such as Actigard™ and BION for BTH (Kunz et al., 1997) and ORYZEMATE for Probenazole (Meijiseika Co., Tokyo, Japan). Although BTH activates the production of high levels of the defense-related proteins, chitinase, peroxidase, and glucanase, there is a concurrent reduction in the growth of Arabidopsis thaliana (Dietrich et al., 2005). For example, BTH treated wheat plants under field conditions resulted in a reduction of crop yields in the absence of pathogens (Heil et al., 2000), suggesting that a fitness cost to produce defense-related metabolites should be utilized to make a balance between physiological growth and defense machinery (Heil et al., 2000; Heil and Baldwin, 2002).

Upon infection of bacterial and viral pathogens, SAR might be effective to control these pathogens. The mechanism by which BTH treatment reduces bacterial and viral diseases on plants has remained unanswered because antimicrobial activity mediated by BTH has been uncharacterized. Studies with npr1 mutant plants, which are unable to accumulate SA, show that BTH functions similarly to SA in the induction of defense-related gene expression (Friedrich et al., 1996). However, the factors that confer a
quantitative RT-PCR was conducted using iQ™ (GenBank accession no. AY572427) transcript. The relative comparison between the target transcript and the compatible and incompatible pathovars of genes explained by transcriptional expression of defense-related genes.

The objective of this study was to assess the de novo accumulation of antibacterial compound(s) compatible and incompatible pathovars of Xanthomonas axonopodis from leaf exudate (LE) containing apoplastic fluid (AF) of pepper leaves that had been treated systemically with soil drenching of BTH. The BTH-mediated SAR was validated by up-regulation of defense signaling marker genes such as CaPR1, CaPR4, CaPR9, and CaCHI2. Heat treatment of leaf exudate did not alter its antibacterial activity. Further analysis revealed that the molecular weight of the active ingredient should be less than 5 kDa. Although we could not characterize the compound(s), liquid chromatography-mass spectrometry (LC-MS) provided a clear difference at a certain retention time in LE between BTH- and water-treated leaves. Our data shows that BTH-treated leaves produce antibacterial metabolites against bacterial pathogens, indicating that certain compound(s) directly respond to bacterial pathogens by inhibition of bacterial growth from LE after occurring SAR.

The seeds of pepper plants (Capsicum annuum L. cv. Bukwang) were surface sterilized with 6% sodium hypochlorite, washed four times with sterile distilled water (SDW), and then incubated on MS agar medium (half-strength) at 25°C for 7 days with an 8 h light/16 h dark photoperiod until germination. The germinated seeds were transplanted on soilless media (Punong, Co. Ltd., Gyeongju, South Korea) and grown in a growth chamber at 23°C for 7 days with an 8 h light/16 h dark photoperiod. The roots of 6-week-old pepper plants were treated with either 10 ml of 0.5 mM benzo-(1,2,3)-thiadiazole-7-carboxothioic acid S-methyl ester (benzo[b]thiadiazole = BTH) (Syngenta, NC, USA) or sterile water. Pepper leaves were collected at 7 days after the BTH or water application.

After BTH treatment for a week, plant leaves were ground in liquid nitrogen, and 200 mg of the powder was used for a single process of RNA isolation. The RNeasy mini kit (Qiagen) was used to get pure total RNA according to the manufacturer’s instruction. The first strand cDNA from the purified total RNA was synthesized by using M-MLV reverse transcriptase (Enzymomics, Daejeon, Korea) at 42°C for 1 h. To analyze changes of transcripts, quantitative RT-PCR was conducted using iQ™CYBR Green Supermix (Bio-Rad, CA, USA) and Chromo 4 (Bio-Rad, CA, USA) with an annealing temperature of 55°C. The relative transcript abundance was calculated by comparison between the target transcript and the CaACT1 (GenBank accession no. AY572427) transcript. The relative mRNA expression of CaPR1, CaPR4, CaPR9, and CaCHI2 was examined as reported in previous papers (Park et al., 2001, 2002; Yang et al., 2009). The genes were analyzed using the following primer sets: for CaPR1, 5'-ACTTGGC AATTATGATCCACC-3' and 5'-ACCCGATTTCACTGTTGGA GCCATT-3'; for CaPR4, 5'-AACGGATTGACATGCCTAACG-3' and 5'-AATTCTGATAGCCGTGACGGT-3'; for CaPR9, 5'-GACTAGTTCAAGAGCATACTAC-3' and 5'-AATTGCTAGGCTGAGTC-3'; for CaCHI2, 5'-ATT GGACGATGGAGCCATACACCAG-3' and 5'-ATAATCC GAAATGCTAAAAGTGGAAC-3'; and for CaACT1, 5'-TTGG GACTCTGGGATGGTGTG-3' and 5'-AACATGGITGA GCCACCACTG-3'.

To collect LE from pepper leaves, the leaves including petioles were cut from the plants, collected, and put in distilled water. After incubation at room temperature for 1 day, the distilled water containing leaf exudate was filter sterilized (0.22 µm pore). The sterilized solution was used for assessment of bacterial growth in LE. The effect of pepper LE on the pathogenicity of a pathogenic bacterium X. axonopodis pv. vesicatoria (Xav) and a non-host pathogen X. axonopodis pv. glycinea (Xag) was analyzed. The two pathovars of Xanthomonas were cultivated in LB broth media and washed-out using sterilized distilled water. The bacteria were resolved in the LE solution at a concentration of 200 cfu/ml. After incubation at 30°C for 1 day, the solution containing bacteria was spread out on LB agar media and bacterial colonies were counted the next day.

As this was the first trial for identification of inhibitory molecules in pepper LE, we separated exudate into two sizes using membrane filters (Amicon®, Millipore, USA) with molecular cut-offs of 10 kDa and 5 kDa and checked their activity to inhibit bacterial growth. The exudate was initially purified with a syringe filter of 0.22 µm pore size to remove impurities, and then the purified exudate was separated into two samples using two different sizes of centrifugal filters (10 kDa and 5 kDa). The serial filtering of separated exudate was conducted from 10 kDa to 5 kDa for further analysis. All of the centrifugation steps were performed at 4,000 x g (Union 32R, Hanil science medical, Korea) at 4°C.

In this study, analysis of variance for experimental data sets was performed using JMP software v.5.0 (SAS Institute, Cary, NC, USA). Significant effects of treatment were determined by the magnitude of the F value (P = 0.05). When a significant F test was obtained, separation of means was accomplished by Fisher’s protected least significant difference (LSD) at P = 0.05.

It is known that SAR can be induced by avirulent pathogens (Reglinski et al., 2007) and chemicals (Reignault and Walters, 2007). Specifically, Lawton et al. (1996) demon-
strated that BTH could induce SAR to virus and bacteria in *Arabidopsis*. Treatment of tobacco with BTH activates SAR against Tobacco mosaic virus (TMV), *Cercospora nicotianae*, *Pectobacterium carotovorum*, *Pseudomonas syringae* pv. *tabaci*, and *Phytophthora parasitica* (Friedrich et al., 1996). Several reports presented that BTH can play a role in inducing strong SAR in response to diverse pathogens in crop plants (Brisset et al., 2000; Görlich et al., 1996; Maxson-Stein et al., 2002; Smith-Becker et al., 2003). Recently, it has been reported that BTH applied to pepper plants is likely to enhance plant defenses to even insect pests including whitefly and aphid infestation as well as against microbial pathogens (Lee et al., 2012; Yang et al., 2011).

In our system, the growth parameters of pepper plants treated with 0.5 mM BTH were similar to water-treated control plants, indicating that 0.5 mM BTH does not have a negative effect on plant fitness in pepper (data now shown). Given that the concentration of BTH used was proper for plants, we tested whether SAR was induced in pepper leaf by soil drench-application of BTH. In general, *CaPR1* and *CaPR9* have been implicated in the SA-dependent signaling pathway, *CaPR4* has a role in the jasmonic acid (JA)/ethylene (ET) pathway, and *CaCHI2* is a marker gene of the ET-related pathway. To confirm SAR induction, transcriptional expression of *CaPR1*, *CaPR4*, *CaPR9*, and *CaCHI2* in pepper seedling leaves was examined by qRT-PCR analysis at 7 days after BTH treatment on roots. After normalization of the expression levels of each gene to constitutively expressed *CaACT1*, the expression of *CaPR1*, *CaPR4*, *CaPR9*, and *CaCHI2* was up-regulated by a 4.0-, 12.0-, 22-, and 7.5-fold, respectively, at 7 days after drench-application of 0.5 mM BTH to roots when compared with water-treated control (Fig. 1). Meanwhile, lower bacterial population in leaves was confirmed in BTH soil-drenched plants (data not shown). The data indicate that root-applied BTH can activate induced resistance in leaves, in which the genes involved in SA and JA/ET signaling may be strong candidates for the induction of SAR at the transcriptional level. In contract to our data, the two *PR1* genes in wheat were not induced by BTH, suggesting that BTH-modulated plant defense may be differentially regulated in monocots as compared to dicots.

Although several pathogenesis-related (PR) genes were induced in pepper leaves after application of BTH on the roots (Fig. 1), additional evidence was necessary for the molecular mechanism of BTH-induced SAR against pathogens. To obtain answers for how SAR-elicited plants are able to actually suppress growth of bacteria, experiments were performed to identify antibacterial substance(s) from systemic tissue during elicitation of SAR by BTH treatment on roots. The trial experiments were conducted to screen appropriate solvents for maximum extraction of leaf exudate (LE) from pepper leaves (data not shown); these experiments showed that water was an effective solvent for leaf exudate within 24 h at room temperature.

Eventually, the water-based extraction method was selected to extract LE from pepper leaves in sterile plastic boxes at 7 days after root drenching with BTH (Fig. 2A). The same number of two pathogenic bacteria (10^7 cfu/ml), *Xav* (compatible) and *Xag* (incompatible), was used at 0 h time point per each treatment. The bacterial number was counted on the LB media by dilution plating method at 24 h after cultivation in LE. The number of bacteria of *Xav* and *Xag* in LE of BTH-treated plants showed an 8.4- and 10.2-fold reduction, respectively, when compared with water-treated plants (Figs. 2B and 2C). The data showed that the level of bacteria population of *Xav* and *Xag* was similar to initial number of bacteria but not completely be diminished level, suggesting that unidentified compounds could have a bacteriostatic effect. Next, to evaluate the heat stability of the antibacterial compounds in LE treated with BTH, the collected exudate from water- or BTH-treated plants was boiled at 100°C for 30 min. Boiling did not change the antibacterial activity against the two bacterial pathogens. The data showed that a significant reduction of bacterial number was observed in *Xav*-BTH (16.4-fold) and *Xag*-BTH (20.1-fold) (Figs. 2B and 2C).

Although the bacterial growth of *Xav* and *Xag* was not affected by heat stress in BTH treated exudate, bacterial fitness of *Xav* in the boiled water but not *Xag* was affected (Figs. 2B and 2C). It indicates that any complex forms or long-chain bacterial nutrients (food) can be degraded at high temperature to useful forms for *Xav* bacterium. Alternatively, antibacterial molecules can be naked to be less active in water-LE to facilitate of *Xav* rather than *Xag*. Taken together, any compounds in exudate from BTH treatment can modulate the antibacterial activity to the two

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**Fig. 1.** Confirmation of systemic acquired resistance by BTH in pepper. Expression of defense-related marker genes *CaPR1*, *CaPR4*, *CaPR9*, and *CaCHI2* in pepper leaves at 7 days after treatment with BTH on roots (black bars) or water control (white bars) was examined by qRT-PCR.
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bacterial pathogens, even if exposed to heat stress. In addition, any substances in exudate accumulated by BTH treatment can activate induced resistance to both compatible and incompatible pathogens, suggesting that BTH-treated pepper plants may enhance basal defense to general bacterial pathogens.

To characterize the compounds in LE from water and BTH-treated pepper, the exudates were analyzed by using LC-MS. The data showed that the peaks in chromatograms of BTH-treated pepper plants were different from water-treated pepper plants at approximately 7.36, 21.76, and 29.23 min retention time (data not shown). Although candidate peaks were identified, the exact name(s) of molecule(s) were not elucidated using our system up to now. Further analyses (i.e., NMR) are remained to characterize the molecular compound(s) that have an antibacterial activity in LE from BTH treated plants.

There are a number of phytoalexins in plants, among which capsidiol is the main bicyclic sesquiterpene in tobacco (Nicotiana tabacum) and pepper (Capsicum annuum) (Maldonado-Bonilla et al., 2008). Also, pepper genes involved in biosynthesis of capsidiol were up-regulated in BTH treated plants when compared with control plants (data now shown). Unexpectedly, no capsidiol was detected in this study. Several possible scenarios were considered to address this aspect. First, the amount of capsidiol is too low to be detected by LC-MS used in this study. Second, the activity of capsidiol could be inhibited by activation of other substances in LE. Alternatively, capsidiol activity may be eliminated under certain conditions of LE used in this study. Finally, the activity of capsidiol in LE may be compromised during modification with post-transcriptional and/or post-translational processing, or in part, concentration of BTH used here could not activate capsidiol.

Although candidate peaks were identified (data not shown), no more information was obtained from previous find-
As part of an effort to identify the antibacterial ingredient(s) from BTH treated LE containing AF, the exudate was divided into two samples according to size using membrane filters that have molecular cut-offs at 10 kDa and 5 kDa each. The exudates below 10 kDa and below 5 kDa were further examined for their inhibition of bacterial growth. After incubation of Xav and Xag in the fractionized exudates for 24 h, the surviving bacteria were counted on agar media. The results showed that fewer bacterial colonies were observed in the BTH treated LE than in the water treated control regardless of the size fractionation and bacterial strains (Figs. 3A and 3B). These results suggest that the size of the inhibitory molecule is small, approximately below 5 kDa. Converting the 5 kDa mass into amino acids, which have an average molecular weight of 110 Da, the inhibitory molecule is likely to be a peptide consisting of 45 amino acids, with a gene of approximately 135 bp in length. Based on the data, the antibacterial molecule(s) may possess the bacteriostatic activity rather than the bactericidal effect.

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