Resistance to Citrus Canker in Key/Mexican Lime
Induced by β-Aminobutyric Acid and Green Tea

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Abstract: Citrus bacterial canker, caused by Xanthomonas citri subsp. citri (Xcc), is a destructive
disease. So far used chemicals to control this pathogen are either not effective or have harmful effects
on the environment. To improve control of this disease, lime (Citrus aurantifolia) plants inoculated
with Xcc were treated with β-Aminobutyric Acid (BABA), ascorbic acid (vitamin C), thiamin (vitamin
B1), green tea (Camellia sinensis), copper oxychloride and distilled water. Lesion diameters of
inoculated leaves were evaluated twenty days after treatment. The results showed that BABA and
green tea had inhibitory effects on disease development. None of the agents used for plant treatment
had direct antimicrobial activity on Xcc, except copper oxychloride. This indicated that the inhibitory
effects of BABA and green tea resulted from strengthening the defense capacities of the plant. To
support this claim, partial coding sequences of Pathogenesis-Related (PR) genes from lime were cloned
and sequenced. Analysis of PR gene expression showed increased mRNA levels of β-1,3-glucanase and
chitinase, during disease development. Reduction in lesion size and lack of antimicrobial activity indicate
that BABA and green tea might be useful treatments against Xcc infection.

Key words: Induced resistance, pathogenesis-related genes, plant activator, Xanthomonas. citri subsp.
Citri, β-Aminobutyric Acid (BABA), Salicylic Acid (SA), Copper Ammonium Carbonate
(CAC), copper bacteriocides include, citrus canker, green tea

INTRODUCTION

Asiatic citrus canker, caused by the endophytic bacterium Xanthomonas citri subsp. Citri (Schaad et
al., 2006) (Xcc) is a serious disease of commercial citrus varieties and some citrus relatives, distributed in
widespread areas of the world (Goto, 1992). The pathogen causes lesions on leaves, twigs and fruits
which later develop into white or yellow spongy pustules. These pustules then become darker and
thicker and turn into brown corky cankers which are often surrounded by water-soaked margins. Severe
infections can result in defoliation, badly blemished fruits, premature fruit drop, twig dieback and general
tree decline (Rossetti, 1977; Stall and Seymour, 1983; Chand and Pal, 1982; Civerolo, 1981). Non-marketable
quality due to lesions and premature fruit drop are the most economically important damages. Since all citrus
cultivars are susceptible to canker, prevention, quarantines, control and eradication programs are the
only tools available to control the disease, but come at a high annual cost for producers. Differences in
susceptibility are mostly attributed to differences in stomata frequency and size (Goto, 1992). Stomata and
wounds are Xcc’s sites of entry into host plants. Within four days after entry, water soaked lesions appear on
the infected tissue. A survey evaluated the effectiveness of thirteen bactericidal chemicals on three citrus species
to control canker by spray application. In this study, Copper Ammonium Carbonate (CAC) with 8% copper
content was consistently the most effective agent in controlling Xcc (Mur et al., 1996). CAC is a contact
poison and used to achieve adequate control on susceptible citrus hosts such as grapefruit and sweet
orange (Graham et al., 2004; Schubert et al., 2001; Leite Jr et al., 1987). Disadvantages of long-term use of
copper bacteriocides include induced copper resistance in xanthomonad populations (Marco and Stall, 1983;
Rinaldi and Leite Jr, 2000) and accumulation of copper in the soil with potential phytotoxic and adverse environmental effects (Alva et al., 1995). However, other contact bactericidal agents were not as effective as copper because they lack sufficient residual activity to protect leaf and fruit surfaces for extended periods (Graham et al., 2006; Rinaldi and Leite Jr, 2000; McGuire, 1988). Various synthetic and biological compounds are capable of controlling a large variety of plant diseases without displaying a direct antibiotic effect (Jakab et al., 2001). In recent years, the importance of vitamins as nutrients and as disease control agents has been emphasized (Pavet et al., 2005; Beyer et al., 2002). The objective of our study was the control of Xcc by use of new compounds with low adverse environmental effects. Moreover, we tested compounds that have systemic effects inside the plant which would be advantageous over contact bactericidal agents like copper containing compounds.

**MATERIALS AND METHODS**

**Plant materials:** Two year old lime juveniles were raised from seeds and used as plant material. They were maintained at 25-30°C and 60% relative humidity conditions in a greenhouse. The study was performed from 2005-2009.

**Bacterial culture:** Bacterial inoculum was prepared with Xcc strain J1 which we isolated from a lime leaf showing canker symptoms and confirmed by PCR using specific primer, XACF [CGTCGCAATACGATTGGAA] and XACR [CGGAGGCATTGTCGAAGGAA] (Park et al., 2006). The bacterial strain was maintained by serial transfer on Nutrient Agar (NA) (Merck) and periodically re-isolated from inoculations described below to maintain virulence. For greenhouse tests, the strain was grown in nutrient broth and incubated on a shaker at 28°C for 24 h.

**Inoculation and treatment of plants:** To achieve high relative humidity, plants were covered with plastic bags one day prior to inoculation. All chemicals were purchased from Sigma-Aldrich Chemical Co. except for Green tea, which was obtained from the local market. Inoculum was prepared in a 0.85% (w/v) NaCl solution, adjusted turbidometrically (620 nm) to approximately 107 cfu.ml-1 and infiltrated into the leaves by a needle-free syringe. Infected plants were treated by foliar spray application of β-Aminobutyric Acid (BABA) 250 ppm, vitamin C 1.5 mM, green tea (Camellia sinensis) 0.58% (w/v), copper oxychloride 0.6% , vitamin B1 50 mM, Salicylic Acid (SA) 2 mM and for control distilled water, on the entire plant, one week after appearance of first symptoms. Lesion diameter evaluation was performed 20 days after chemicals treatment.

**Inhibitory effect of treatments on Xcc in vitro:** To determine direct antimicrobial activity of the different agents on Xcc, the well diffusion method was used (Rojas et al., 2006). Thus, 100 µL cultured strain at a concentration of 107 cfu.ml-1 was added uniformly to cooled but still liquid Nutrient Agar. After solidification, wells were punched out of the agar and 100 µL of each agent, adjusted to the above given concentration, was added into the wells. Distilled water served as a control. Plates were incubated at 28°C for 72 h. A clear growth-inhibition zone around the wells was measured as an indication for antimicrobial activity.

**Total RNA extraction from lime leaves:** Approximately 1-2 g of leaves were ground with a mortar and pestle in the presence of liquid nitrogen. The powder was transferred to a tube containing ice cold extraction buffer (100 mM Tris-HCl pH 8.5/10 mM EDTA pH 8/100 mM LiCl/1% SDS) and mixed. Tris-equilibrated phenol was added, mixed before chloroform/isoamylalcohol was added and mixed. After centrifugation, the upper aqueous phase was extracted two times with chloroform/isoamylalcohol (24:1). Addition of 10 M LiCl2 to the supernatant precipitated the RNA over night. After centrifugation, the RNA pellet was resuspended in RNase-free H2O. RNA quantity and purity was analyzed by measuring the ratios of absorption at 260/280 and 260/230 nm and RNA integrity was evaluated from the 28S and 18S rRNA bands by ethidium bromide staining after agarose gel electrophoresis.

**Preparation of cDNA:** About 1 µg total RNA was DNase treated and used to make cDNA. First-strand cDNA was prepared using RevertAid™ First Strand cDNA Synthesis Kit (Fermentas) according to the manufacturer’s instructions.

**Purification and cloning of cDNA fragment:** Specific primers were used to amplify cDNA fragments, including: F (5’-AATGTTGCTAGCATTGAGCTCC-3’) and R (5’-GCAGCATTGAAACGCATCT-3’) for Chitinase (CH1) AF090336, F(5’-TTCCGACGGATCTTTAAGTTACC-3’) and R (5’-CCACCCCCACTCTCATGATACGG-3’) for β-1,3-Glucanase (GNSL) AJ000081, F (5’-GGTCAAGCTGAGCATGC-3’) and R (5’-CATCGTACCTAGCCTTTGAGT-3’) for Elongation factor 1-alpha (EF1) AY498567. The amplified fragments were excised from an agarose gel and purified with the AccuPrep PCR Purification Kit according to the manufacturer’s instructions (Bioneer, South Korea). The purified DNA was ligated into the pTZ57R/T vector using InsT/A Clone™ PCR product Cloning Kit (Fermentas). The plasmids were
transformed into competent *E. coli* cells. Recombinant plasmids were identified by colony PCR and purified plasmids confirmed by restriction enzyme digestion. The recombinant plasmids were extracted using the AccuPrep® Plasmid extraction Kit according to the manufacturer’s instructions (Bioneer, South Korea).

**DNA sequencing:** Recombinant plasmids were sequenced in both directions by extending M13 reverse and forward primers, using the Automatic DNA Sequencer 3730Xl (Macrogene, Korea). The sequencer data were edited using Chromas software Version 1.41. The nucleotide sequence was analyzed using BLAST (Benson *et al*., 2002).

**Expression analysis of β-1,3-glucanase and chitinase by semi quantitative PCR:** Expression of a β-1,3-glucanase and chitinase genes was analyzed by comparing the PCR band signals to the signals from the *EF1* gene. Each PCR reaction contained 0.5 μM of each primer, 2 μL of cDNA template and 10 μL of 2x PCR Master Mix (Fermentas). Cycles were programmed as follows: The initial denaturing cycle at 94°C for 5 min, followed by 35 cycles of denaturing at 94°C for 1 min, annealing at 55°C for 1 min and elongation at 72°C for 2 min, followed by a final cycle at 72°C for 7 min. To define the optimal number of PCR cycles for exponential amplification, an aliquot from each of the samples was removed at 30 cycles thereafter at every fifth additional cycle. Expression was analyzed before and after inoculation and after treatment.

**Statistical analysis:** The experiment was arranged based on a Completely Randomized Design (CRD) with ten biological replicates. Ten leaves from each replicate were evaluated for necrotic lesion diameter and the mean of them used as each replication value. The means were compared for statistical significance using Duncan’s multiple-range test (p = 0.05).

**RESULTS**

We inoculated citrus leaves with *Xcc* and after first symptoms became visible attempted to reduce their spread by spray-treatment of the inoculated leaves with BABA 250 ppm, vitamin C 1.5 mM, green tea 0.58% (w/v), copper oxychloride 0.6%, vitamin B1 50 mM, SA 2 mM. Our results showed that spray treatment with green tea and BABA reduced lesion size by about 50% as compared to the water control. Thus while water-treated leaves showed lesion sizes of about 1.2 cm, BABA and green tea treated plants had lesions of only about 0.6 cm. Treatments with vitamin C or vitamin B1 reduced lesion size by about 25%. No significant reduction in lesion size was observed following treatment with copper oxychloride or salicylic acid (Fig. 1-2).

![Fig. 1: Suppressive effect of different treatments on necrotic lesion diameter caused by *X. citri* subsp. *citri* on lime leaves. Leaves of two years old juvenile were inoculated with *Xcc* and after appearance of symptoms treated with the various compounds. The lesion diameters were measured 20 days post application of the chemicals. GT: Green Tea 0.58% (w/v); BABA: β-aminobutyric acid 250 ppm; SA: salicylic acid 2 mM; Vit B1: Vitamin B1 50 mM; Vit C: Vitamin C 1.5 mM; CO: Copper oxychloride 0.6%. Different letters (a, b, c) at each column denote significant differences between means (p = 0.05) according to Duncan’s multiple-range test. The experiment was repeated three times.](image1)

![Fig. 2: Treated leaves with BABA and Green Tea showed smaller necrotic lesions then control and other treatments. Leaves were infected by injection with 1×107 cfu.ml-1 *Xanthomonas citri* subsp. *citri*. Leaves were photographed at the indicated times post chemical application. A; BABA; B, Green Tea; C, Vit C; D, Vit B1; E, Control; F, Salicylic acid; G, Copper Oxychloride. In order to test whether the reduction in lesion size was due to a direct growth-inhibitory effect of the applied agents, we tested their effect on the growth of *Xcc in vitro*. Thus, rapidly growing *Xcc* were embedded into nutrient agar plates into which holes were punched. Into these holes, 100 μL of BABA, green tea, vitamin C and copper oxychloride were applied and the growth of *Xcc* was observed after over-night incubation. As can be seen in Fig. 3, only copper oxychloride had a direct growth-inhibiting effect on the bacteria, since all other compounds allowed uninhibited growth of the bacteria.](image2)
Fig. 3: Test for direct antimicrobial activity of the compounds suppressing lesion size. Rapidly growing Xcc bacteria were embedded in nutrient agar and incubated with the compounds indicated. Copper oxychloride showed antimicrobial activity against X. citri subsp. citri. Each compound was applied in two wells. GT; Green Tea, CO; Copper Oxychloride; w; distilled water, BABA; β-aminobutyric acid, Vit C; vitamin C. The observed reduced lesion size after green tea and BABA treatments prompted us to test whether enhanced physiological defense responses can be observed in the treated lime plants. Thus, we cloned by RT-PCR a 119 bp fragment of a β-1,3-glucanase and a 110 bp fragment of a chitinase gene from lime (Fig. 4).

Fig. 4: RT-PCR bands of β-1,3-glucanase and chitinase genes of lime. Numbers to the left of photo show marker band sizes. L; ladder 3 kb, Glu; β-1,3-glucanase, Chi; chitinase. Sequencing the cloned PCR fragments allowed to align them with the sequences from orange. The alignment showed large regions of identity (Fig. 5), confirming that we cloned the correct pathogenesis-related genes.

Fig. 5: Alignment of amplified lime (Citrus aurantifolia) cDNA with orthologous sequences from orange (Citrus sinensis). Two fragments (110 bp for chitinase and 119 bp for glucanase) were amplified by RT-PCR from lime and sequenced. The obtained nucleotide sequence aligned with A) >99% identity β-1,3-glucanase and B) 97% identity chitinase to the orthologous orange sequences. Using the same primers, we investigated the expression pattern of the two genes 24 h after infection with Xcc and a further 24 h after treatment with BABA or green tea by semi-quantitative reverse transcription PCR. As can be seen in Fig. 6, infection up-regulated glucanase expression. Further treatments did not appreciably influence expression of glucanase or chitinase.

Fig. 6: Expression analysis of lime PR genes after Xcc inoculation and subsequent treatment with chemicals. Semi-quantitative reverse-transcription PCR was used to compare the relative transcript amounts of the PR genes β-1,3-Glucanase (Glu) and Chitinase (Chi) after treatment of Xcc-infected lime leaves with BABA and Green Tea. Glu; β-1,3-glucanase, Chi; chitinase, EF1; Elongation Factor 1-alpha, C0; control (time 0), M24hpi; non infected plant Mock 24 h post inoculation, 24hpi; Infected plants 24 h post inoculation, C24hpt; non treated Inoculated plant (control 3) 24 h post treatment; B24hpt; Treated plant with BABA 24 h post application, GT24hpt; Treated plant with Green Tea 24 h post application. Application of treatments were performed one weak after infection.
DISCUSSION

Citrus canker is a disease causing great economic damage to the afflicted regions. Effective treatments contain copper that limits its application to a few per year due to its general toxic effects. Our study aimed at identifying agents that do not have a general toxicity effect, yet limit pathogen damage on the plant. A function of vitamin B group in disease resistance has been described (Dong and Beer, 2000). Treatment with some members of vitamin B group protected tobacco (Nicotiana tabacum) and Arabidopsis plants from fungal and bacterial infections without inhibiting pathogen growth (Dong and Beer, 2000). Our results indicate that among the different treatments, BABA, green tea and vitamin C significantly reduced canker lesions in lime leaves. In vitro assays indicated that none of these agents had direct antimicrobial activity on Xcc confirming previous results (Cohen et al., 1999; Reuveni et al., 2003). Only copper oxychloride at 0.6% had a direct negative effect on the growth of the bacteria. Therefore, the reduced lesions obtained after BABA or green tea treatments cannot be explained by direct antimicrobial characteristics. We wondered whether BABA or green tea could raise the activation of a highly coordinated biochemical and structural defense system. The protection of plants from bacterial invasion by activation of the plant’s defense system is an accepted concept (Lawton et al., 1996; Schroder et al., 1992). The mode of action of BABA in inducing resistance against Xcc is not yet fully understood. There are reports showing the induction of PR genes after BABA treatment (Cohen, 1994). Green tea is full of components with anticancer, antiviral and anti pathogenic characteristics like polyphenols (catechins), fragrance components, amino acids (theanine), saponins and γ-amino butyric acid (GABA). Thus, any of these constituents or a combination of them might induce increased pathogen defense. In plants, GABA accumulation was observed in response to biotic and abiotic stress (Roberts, 2007; Shelp et al., 1999). Here, we report for the first time a partial mRNA sequence of a β-1,3-glucanase and chitinase genes (PR) of lime. Analysis of the steady-state mRNA levels of these genes by semi-quantitative reverse transcription PCR showed that one PR gene (β-1,3-glucanase) showed enhanced expression in response to Xcc infection (Fig. 6). We cannot see the reported effects of BABA on PR-1, chitinase and β-1,3-glucanase accumulation in pepper, tomato and tobacco (Cohen, 1994; Cohen et al., 1994; Hwang et al., 1997). But understanding of the signaling pathways of BABA and green tea in resistance of lime to Xcc clearly deserve more investigation.

CONCLUSION

Application of BABA or green tea in citrus possesses promising potential in the control of the disease and could serve as an alternative to traditional chemical compounds which are harmful to the environment.

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