Comparative analysis of differentially expressed sequence tags of sweet orange and mandarin infected with Xylella fastidiosa

Alessandra A. de Souza¹, Marco A. Takita¹,², Helvécio D. Coletta-Filho¹, Magnólia A. Campos³, Juliana E.C. Teixeira⁴, Maria Luisa P.N. Targon¹, Eduardo F. Carlos¹, Juliano F. Ravasi⁵, Carlos N. Fischer⁵ and Marcos A. Machado¹

¹Centro APTA Citros Sylvio Moreira, Instituto Agronômico de Campinas, Cordeirópolis, SP, Brazil. ²Centro de Pesquisa e Desenvolvimento de Recursos Genéticos Vegetais, Campinas, SP, Brazil. ³Departamento de Biologia, Universidade Federal de Lavras, Lavras, MG, Brazil. ⁴Departamento de Genética e Fitopatologia, Escola Superior de Agricultura Luiz de Queiroz, Universidade de São Paulo, Piracicaba, SP, Brazil. ⁵Departamento de Estatística, Matemática Aplicada e Computação, Instituto de Geociências e Ciências Exatas, Universidade Estadual Paulista, Rio Claro, SP, Brazil.

Abstract

The Citrus ESTs Sequencing Project (CitEST) conducted at Centro APTA Citros Sylvio Moreira/IAC has identified and catalogued ESTs representing a set of citrus genes expressed under relevant stress responses, including diseases such as citrus variegated chlorosis (CVC), caused by Xylella fastidiosa. All sweet orange (Citrus sinensis L. Osb.) varieties are susceptible to X. fastidiosa. On the other hand, mandarins (C. reticulata Blanco) are considered tolerant or resistant to the disease, although the bacterium can be sporadically detected within the trees, but no disease symptoms or economic losses are observed. To study their genetic responses to the presence of X. fastidiosa, we have compared EST libraries of leaf tissue of sweet orange Pêra IAC (highly susceptible cultivar to X. fastidiosa) and mandarin ‘Ponkan’ (tolerant) artificially infected with the bacterium. Using an in silico differential display, 172 genes were found to be significantly differentially expressed in such conditions. Sweet orange presented an increase in expression of photosynthesis related genes that could reveal a strategy to counterbalance a possible lower photosynthetic activity resulting from early effects of the bacterial colonization in affected plants. On the other hand, mandarin showed an active multi-component defense response against the bacterium similar to the non-host resistance pattern.

Key words: ESTs, citrus variegated chlorosis, biotic stress, CitEST, phytopathogen.

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Introduction

Citrus is one of the most important crops in Brazil. The State of São Paulo is responsible for 80% of the domestic production of fresh fruit and 70% of the world production of concentrated orange juice. However, the Brazilian citrus industry faces constant challenges. The production is mostly conducted without irrigation and the industry is affected by many diseases and pests (Fernandes-Jr, 2003).

The citrus variegated chlorosis, caused by Xylella fastidiosa, is currently one of the most limiting diseases affecting the groves. The annual loss due to CVC is more than the US$100 million spent on chemical control of the vectors (sharpshooters), pruning infected branches and trees, and replacing old orchards (Fernandes-Jr, 2003). Genetic control is still not feasible because all varieties of sweet orange (C. sinensis L. Osb.) used for juice production are highly susceptible to the disease. CVC diseased plants have a strong correlation between symptoms and disturbance caused by water stress, which is in accord with the hypothesis of xylem blockage caused by X. fastidiosa (Machado et al., 1994). On the other hand, mandarins and some hybrids with sweet orange seem to be tolerant or resistant to X. fastidiosa. Genetic mapping using these hybrids has revealed a complex inheritance to the resistance (Oliveira et al., 2005). However, there is little information about mo-
Molecular mechanisms that take place in the interaction between resistant plants and the bacterium.

The use of gene tags for evaluating the response to infections in plants represents a new approach in studies of plant-pathogen interactions (Lee et al., 2004). The identification and analysis of genes that are up or down expressed in the affected plants is thus an initial step that could help in the understanding of the entire mechanism of plant defense response. This knowledge may lead to a generation of disease-resistant plants.

The generation and analysis of expressed sequence tags (ESTs) have been employed in genomic research, especially for research with plant species with large genome sizes. The subsequent analysis of the ESTs has become a popular method for examining the spatial or temporal pattern for gene expression in many plants (Lee et al., 2004).

To understand the plant defense mechanism activated in response to the infection of X. fastidiosa, we ran a comparative analysis of gene expression profiles of the resistant ‘Ponkan’ mandarin (Citrus reticulata) and the susceptible ‘Pêra IAC’ sweet orange (Citrus sinensis) under the presence of the pathogen.

Material and Methods

Xylella fastidiosa inoculum and monitoring of the bacterial infection

Four plants, six months of age, of ‘Ponkan’ mandarin and ‘Pêra IAC’ sweet orange grafted on Rangpur lime, were artificially inoculated by needle prick with 50 μL suspension (10^10 cells mL^-1) of X. fastidiosa strain 9a5c in PBS buffer. Negative controls of ‘Ponkan’ mandarin and ‘Pêra’ sweet orange were mock-inoculated with PBS buffer. The plants were maintained in a growth chamber (Conviron PGW36, Winnipeg, Canada) with photoperiod of 16 h. The range of temperature was from 25 °C to 28 °C.

The X. fastidiosa infection was monitored at seven-day intervals. The material was stored at -80 °C until RNA extraction. Afterwards, the material was stored at -80 °C until RNA extraction.

Leaf sampling for cDNA library

Thirty days after inoculation, a bulk of leaves was collected for cDNA library preparation from the two plants within each species that tested positive for the presence of X. fastidiosa in PCR. Young but expanded leaves were collected and immediately put in liquid nitrogen. After wards, the material was stored at -80 °C until RNA extraction.

Total RNA was extracted using Trizol reagent (Invitrogen). The Poly(A+) RNA was isolated from 1 mg of total RNA with the polyATtract mRNA Isolation System (Promega Corporation, Madison, WI). The SuperScript Plasmid System with Gateway Technology for cDNA Synthesis and Cloning (Invitrogen) was used for the library construction. All the steps were done according to the manufacturers’ instructions.

The sequencing reactions were performed using the Big Dye Terminator DNA sequencing Kit (Applied Biosystems). Sequencing was done in the ABI 3730 (Applied Biosystems).

In silico hybridization and functional annotation

For comparison of the libraries (in silico hybridization) the reads of both cDNA libraries were analyzed and assembled using the Phred and Phrap programs (Ewing and Green, 1998; Ewing et al., 1998) with the default parameters. The resulting ace file was parsed with a homemade perl script, which extracted the identification of the contig together with the composing reads. The resulting file was submitted to a web implementation (http://gbirc.rc.unesp.br/jXPAnalysis/) of the Winflat software (Audic and Claverie, 1997). The statistical cutoff was set to 95% and the sequences of the significantly differentially expressed contigs were compared with the sequences of the non-redundant database of the National Center for Biological
Information (NCBI) using the BlastX tool (Altschul et al., 1997). The functional categorization of sweet orange and mandarin putative genes was done based on similarity with Arabidopsis set of proteins v. 2.0 present at MIPS (Munich Information Center for Protein Sequence, http://mips.gsf.de/proj/funcatDB/search_main_frame.html) using the BlastX tool.

Results and Discussion

Plant inoculation

The presence and viability of X. fastidiosa artificially inoculated into the plants has been used as an approach to monitor the efficacy of the inoculation and to verify whether the plant withstands the pathogen. Here, the presence of X. fastidiosa in inoculated tissues was analyzed by PCR with specific primers and isolated on PW medium (Table 1). After two weeks, inoculation positive amplifications were detected in ‘Ponkan’ mandarin and ‘Pera’ sweet orange. Four to eight weeks after inoculation, all plants were positive for X. fastidiosa. However, positive isolation of bacteria was possible only for inoculated sweet orange plants, which showed initial CVC symptoms 5 months after inoculation. On mandarin, after inoculation, it is possible that the bacteria started growing but failed to colonize the vessels, resulting in a decrease of the population and, consequently, death. However, this hypothesis needs to be proven.

In silico analyses and functional annotation

A total of 19,200 EST sequences was used in the bioinformatics analysis. Of these, 17,695 met satisfactory quality standards, corresponding to 350 nucleotides of Phred quality score of 20 or above. For a score of 20, the accuracy of the base calling is 99% (Ewing and Green, 1998). The sequences were subsequently used in clustering, generating 7,814 contigs and 4,635 singlets after assembly. Through an in silico differential display, we found 152 genes to be significantly differentially expressed (p ≤ 0.05). In this comparison, 81 and 71 putative genes showed higher expression in mandarin and sweet orange infected plants at 30 days after inoculation (dai), respectively.

We categorized the significant contigs into functional groups based on the MIPS Arabidopsis gene functional categories. The functional categories and distributions of the genes differentially expressed in mandarin and sweet orange are shown in Figure 1. Evaluation of mandarin and sweet orange ESTs showed distribution of putative genes in 15 functional categories with emphasis in five, namely: i. Metabolism; ii. Energy; iii. Cellular transport, transport facilitation and transport routes; iv. Cell rescue, defense and virulence; and v. Proteins with unknown function.

Table 1 - PCR and recovery of X. fastidiosa from artificially inoculated plants of 'Pera' sweet orange and 'Ponkan' mandarin.

| Plant inoculation | PCR results after inoculation | X. fastidiosa recovery |
|------------------|-------------------------------|-----------------------|
|                  | 2 weeks | 4 weeks | 8 weeks | P. fastidiosa recovery |
| Ponkan 1         | ++      | ++      | ++      | +                       |
| Ponkan 2         | -       | ++      | ++      | -                       |
| Ponkan 3         | ++      | ++      | ++      | -                       |
| Ponkan 4         | ++      | ++      | ++      | -                       |
| Ponkan mock      | -       | -       | -       | -                       |
| Pera 1           | -       | ++      | ++      | ++                      |
| Pera 2           | -       | ++      | ++      | ++                      |
| Pera 3           | ++      | ++      | ++      | +                       |
| Pera 4           | ++      | ++      | ++      | +                       |
| Pera mock        | -       | -       | -       | -                       |

1 Xylella-specific amplification products from plant inoculated samples collected at two, four and eight weeks after inoculation: (+) positive amplification; (-) negative amplification.
2 Recovery of Xylella fastidiosa, from plant inoculated tissues, on PW medium at 11 weeks after inoculation: P - successful bacteria recovery; N - unsuccessful bacteria recovery.

Figure 1 - Number of significant contigs distributed in functional categories of the mandarin (black) and sweet orange (gray) libraries infected with X. fastidiosa.
Functional categorization and comparative analysis of Sweet Orange and Mandarin

The functional category of putative genes related to energy had the highest number of differentially expressed genes in sweet orange compared to mandarin. This category includes mainly genes associated with photosynthesis (Table 2). No typical chlorosis could be observed in the leaves of plants infected with X. fastidiosa at 30 DAL. This could eventually explain the increase in the expression of genes associated with photosynthesis as a response to chlorophyll degradation. Nevertheless, studies on the physiology of sweet orange plants infected with X. fastidiosa showed that even without development of typical symptoms of chlorosis it was possible to verify early effects of the bacteria on photosynthesis, with a reduction in the activity of the photosynthetic apparatus (Ribeiro et al., 2003). Our results showed that the infection by X. fastidiosa, besides blocking the xylem vessels in the later phase of the disease, induced the expression of several genes responsible for synthesis of important components of the photosynthetic apparatus (Table 2). The higher expression of such genes could be correlated with a de novo synthesis of redox photosynthesis machinery as a mechanism to counterbalance the lower photosynthetic activity resulting from the early effects of the bacterial colonization in susceptible plants. In later stages of the infection, chlorosis is observed in sweet orange and a decrease in transcription of photosynthesis related genes is observed (Souza et al., this issue). This reinforces the idea that the expression of these genes in early stages of the infection is an attempt to maintain the physiological functions at an adequate level.

In mandarin, the most significant functional category was Cell rescue, defense and virulence (Figure 1). Because this species is resistant to X. fastidiosa, the expression of these genes in this organism suggests that they may have an important role in resistance. The genes significantly expressed in mandarin that are possibly associated with defense response to the pathogen are listed in Table 3.

In mandarin we found one gene encoding an NBS-LRR-like disease resistance protein and a second coding for multidomain cyclophilin type peptidyl-prolyl cis-trans isomerase. It has been recently demonstrated that the later protein activates the avirulence protein AvrRpt2 from Pseudomonas syringae in incompatible interaction with Arabidopsis thaliana, leading to RPS2-mediated resistance and providing evidence for the ‘guard hypothesis’ (Coaker et al., 2005). These proteins are normally involved in responses in which virulence proteins of the pathogens are involved. However, neither homologues of avr genes nor their type III secretion apparatus were identified in the X. fastidiosa genome. A possible explanation for the inducible mechanisms activated in mandarin in response to X. fastidiosa can be given from a perception of general elicitors ‘non-self’ or PAMPs (pathogen-associated molecular pattern). This is the basis for the non-host resistance, where general elicitors derived from pathogens activate signaling pathways leading to gene expression and consequently, resistance (Ingle et al., 2006).

Table 2 - Up-regulated genes in sweet orange infected with X. fastidiosa observed in the most relevant functional category.

| MIPS functional category | p-value1 | Best blast match | NCBI number2 |
|--------------------------|----------|------------------|--------------|
| Energy                   | 0.01     | Pyruvate decarboxylase-1 Pdc1 (Arabidopsis thaliana) | gb|AAM674591 |
|                          | 0.01     | Ribose 5-phosphate isomerase (Arabidopsis thaliana) | gb|AAF049051 |
|                          | 0.003    | Photosystem II polypeptide (Arabidopsis thaliana) | gb|AAM201941 |
|                          | 0.004    | 33 kDa polypeptide of water-oxidizing complex of photosystem II (Nicotiana tabacum) | emb|CAA457011 |
|                          | 0.002    | 22kD-protein of photosystem II (Spinacia oleracea) | emb|CAA485571 |
|                          | 0.001    | Oxygen-evolving enhancer protein 2, chloroplast precursor (OEE2) (23 kDa subunit of oxygen evolving system of photosystem II) (OEC 23 kDa subunit) (23 kDa thylakoid membrane protein) (OEC23) (Cucumis sativus) | sp|Q9SLQ8 |
|                          | 0.0009   | Photosystem I subunit V precursor (Arabidopsis thaliana) | gb|AAK003741 |
|                          | 0.00002  | Rubisco small subunit (Coffee arabica) | gb|AA611201 |
|                          | 0.000002 | Rubisco small subunit (Coffee arabica) | emb|CAD119911 |
|                          | 0.01     | Chlorophyll a/b-binding protein CP24 precursor (Vigna radiata) | gb|AAD278822 |
|                          | 0.006    | Chlorophyll a/b-binding protein precursor (Citrus limon) | gb|AAO458851 |
|                          | 0.002    | Chlorophyll a-b binding protein, chloroplast precursor (LHCI type II CAB) (Petunia x hybrida) | sp|P13869 |
|                          | 0.0001   | Phosphoribulokinase, chloroplast precursor (Mesembryanthemum crystallinum) | sp|P27774 |
|                          | 0.0      | Plastidic aldolase (Nicotiana paniculata) | dbj|BAA776031 |

1Probability of having the putative gene differentially expressed only by chance (significant for p < 0.05).
2http://www.ncbi.nlm.nih.gov/.
A gene encoding lipoxygenase (LOX) was induced by *X. fastidiosa* in mandarin. LOX is the key enzyme in the synthesis of jasmonic acid (JA), catalyzing the production of JA from linoleic acid, which stimulates the expression of defense-related genes functioning as secondary signals that activate a subset of defense genes. We also found one gene encoding S-adenosyl-L-methionine:salicylic acid methyltransferase. This gene is responsible for the synthesis of methyl salicylate, which is a plant-signaling compound of the salycilic acid (SA) pathway that activates defense responses (Deng et al., 2005). The presence of two different precursor enzymes of two signal compounds, JA and SA, may indicate crosstalk among regulatory pathways controlling different cellular processes in mandarin during the interaction with *X. fastidiosa*. Some genes associated with defense and related to the oxidative burst were also detected in our study. Among them there are genes that encode antioxidant enzymes, such as peroxidase, Fe-superoxide dismutase and annexin. The identified annexin is similar to the *Arabidopsis thaliana* annexin 1, which possesses peroxidase activity, limiting excessive levels of reactive oxygen species during the oxidative burst (Gorecka et al., 2005). Oxidative Burst is a common mechanism that occurs both in abiotic and biotic stresses. Genes involved with abiotic stress were found to be up-regulated in mandarin infected with *X. fastidiosa*, including those encoding DnaJ and RD22 proteins. DnaJ is a chaperone that functions in protein folding and marks incorrectly folded proteins for turnover (Bohnert et al., 1995). Despite the increase in their expression that occurs during the abiotic stresses, induction in the synthesis of chaperones has also been observed in responses to pathogen attack as well (Jelitto-Van Dooren et al., 1999). The RD22 protein acts in detoxification of oxidative stressors produced under water deficit (Abe et al., 1997; Gillet et al., 1998).

Genes encoding cytochrome p450 were also induced in mandarin by *X. fastidiosa*. Plant P450s are involved in the biosynthesis of secondary products (phenylpropanoids, alkaloids, terpenes, lipids, cyanogenic glycosides and glucosinolates), which give important properties to plants, such as flavor, color and herbivore/pathogen resistance (Bolwell et al., 1994; Schuler, 1996). Moreover, plant P450s are also essential in the biosynthetic pathways of plant growth regulators, including gibberellins, brassinosteroids and jasmonic acid, and in detoxification processes. Many of the best-characterised P450s are inducible enzymes, involved in xenobiotics detoxification, and have relatively broad substrate specificity (Paquette et al., 2000). Since one of the virulence factors of *X. fastidiosa* is the production of toxins and in its genome were found genes encoding toxins and antibiotic production (Simpson et al., 2000), it is reasonable to suppose that this protein could be actively working in the detoxification of these factors.

Several contigs sharing homology with unclassified *Arabidopsis* genes and others for which no similarities could be found with sequences from any database were found to be induced within the mandarin plants after infection with *X. fastidiosa*. This finding may help to elucidate the function of the proteins encoded by them.

The activation of all these genes in mandarin may be part of the defense response mechanisms against pathogens. Moreover, several genes related to cellular transport, transport facilitation and transport routes, specific proteins and transcriptional factors were also induced in mandarin plants inoculated with *X. fastidiosa*.

The response of mandarin to *X. fastidiosa* seems to include features of the non-host resistance, with activation of the gene expression through the recognition of non-specific elicitors. This activation includes two different signaling pathways, the jasmonate and the salycilic acid pathways, which ultimately leads to synthesis of resistance components. In the responses, the production of reactive oxygen species (ROS) leading to the production of detoxifying proteins appears to exist. Another characteristic of this resistance seems to be the production of proteins that are responsible for fighting the toxins or antibiotics probably
produced by the bacterium and that could cause damage to the plant. Therefore, our study suggests that different defense responses are taking place in mandarin in the interaction with *X. fastidiosa* in a multifactor antipathogen response. This hypothesis should be verified by further functional experiments.

In silico analysis of the species-specific genes associated with *X. fastidiosa* response

The *in silico* hybridization using two different species, *Citrus reticulata* and *Citrus sinensis*, can generate clusters that are specific to each of the species. This can affect the analysis of differential gene expression which is based on clustering of ESTs. Therefore, the clusters containing reads from only one of the species were further analyzed by searching for homologues in either *Citrus sinensis* or *Citrus reticulata*. This analysis was performed using the Blastn tool (Altschul et al., 1997). As a result, homologues for all analyzed genes were found in the comparing species, showing that only the expression levels in the different libraries were different.

Some changes in gene induction/repression were observed when comparing the susceptible and resistant Citrus species. Many genes were observed to be up- or down-regulated upon *X. fastidiosa* infection. An examination of a wider range of hybrid between *Citrus sinensis* and *Citrus reticulata* could answer the question of whether our findings represent a general feature of the Citrus-*X. fastidiosa* interaction, which might also lead to development of helpful markers for Citrus breeders. In order to complete the information on the function and nature of the genes that respond to *X. fastidiosa* infection, studies on the protein level as well as on silencing and over-expression of candidate genes will be helpful.

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