**RESEARCH PAPERS** 

## Detection of a new variant of Citrus tristeza virus in Greek citrus crops

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Summary. Citrus tristeza virus (CTV), the most destructive virus of citrus, is a quarantine pathogen in Greece. Since 2000, several accidental imports of infected propagation material have been detected in the country, and while eradication measures were applied, a few disease foci still remain. CTV isolates were collected from Chania (Crete) and the "lemonwood" of Poros (Peloponnese), and their genetic variability was studied using single-strand conformation polymorphism (SSCP). One previously characterized isolate from Argolida grafted on a Mexican lime (GR3) and two Italian isolates from Calamondin were also included in the study. ELISA and RT-PCR tests confirmed CTV presence, and SSCP analysis of the virus amplified coat protein (CP) gene was used to separate either distinct virus isolates for cloning the CP gene or variants (haplotypes) for sequencing. Analyses showed that selected variants of four representative isolates clustered into three of the seven defined phylogenetic groups: groups 3b and 5 (severe isolates) and group M (mild isolates). The prevalent haplotypes detected in the CTV from lemonwood of Poros (GR9) were in group 3b, confirming previous results. However, one sequence variant was identified as a recombinant between haplotypes from groups 3b and 5. Variants of these two groups were also detected in the Italian Calamondin isolate. In the grafted Mexican lime isolate (GR3) from Argolida, only one haplotype was found which belonged to group M, while in the field isolate from Chania (GR6) the only haplotype detected was in group 5. This is the first report of variants of group 5 in Greece, suggesting an unknown virus introduction. The prevalence of severe isolates in the area is of particular concern, and implications for the future of the CTV epidemics are discussed.

Key words: CP gene, CTV isolates, nucleotide diversity, phylogenetic analysis, SSCP, recombination.

#### Introduction

*Citrus tristeza virus* (CTV) (genus *Closterovirus*, family *Closteroviridae*) is the most economically important and damaging virus of citrus trees (Moreno and Garnsey, 2010). This virus is listed among the largest positive stranded RNA viruses of higher plants; its 19.3 kb positive sense ssRNA genome is encapsidated mainly by the p25 capsid protein (CP) and the p27 proteins, in thread-like filamentous particles of size about 2000 × 11 nm (Moreno *et al.*, 2008).

CTV worldwide dispersal occurred via the movement of infected plant material, often grafted on sour orange rootstocks (Citrus aurantium L.) used to control Phytophthora root rots. Local spread of the virus is mainly by aphids in a semi-persistent manner (Moreno et al., 2008). Virus spatial and temporal spread depends on the aphid vector species present (Gottwald et al., 1999). In Europe, virus epidemics are associated with the presence of Aphis gossypii Glover (Cambra et al., 2000), while the less competent vectors A. spiraecola (Patch) and Toxoptera aurantii (Boyer de Fonscolombe) also occur (Hermoso de Mendoza et al., 1988; Cambra et al., 2000). T. citricida (Kirkaldy), the most efficient vector of CTV (Yokomi et al., 1994) is established in Asia, Australia, sub-Saharan Africa, Central and South America, and different Caribbean countries and it is now in Madeira, northern Portugal and Spain (Ilharco et al., 2005; Hermoso de Mendoza et al., 2008).

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Disease syndromes caused by CTV fall into three types depending on virus strain and rootstock-cultivar combination (Moreno et al., 2008). The so-called "severe" CTV strains are associated with (i) quick decline (QD) or tristeza of citrus species grafted onto sour orange or lemon [C. limon (L.) Burm. f.], (ii) the stem pitting (SP) syndrome characterized by the presence of elongated pits on branches of trees grown on tristeza-tolerant rootstocks, which reduces plant vigour and fruit quality, and (iii) seedling yellows (SY), that is observed by biological indexing. Mild CTV strains also occur in field trees but they cause barely detectable symptoms, usually only observed on indicator plants such as Mexican lime [C. aurantifolia (Christm.) Swing.] (Hancevic et al., 2013).

In nature, CTV exists as a mixture of sequence variants, subisolates or recombinants (Broadbent et al., 1996; Kong et al., 2000; Rubio et al., 2001; Matos et al., 2013), displaying high levels of genetic and phenotypic diversity. Biological data appeared inconsistent when analyzing virus variability, while typing methods which target the coat protein (CP) gene are more reliable (Niblett et al., 2000). Singlestrand conformation polymorphism (SSCP) is further used to estimate the genetic diversity within and between isolates (Rubio et al., 1996; Sambade et al., 2002), and worldwide occurring isolates of CTV are now grouped in seven clusters based on the analysis of their CP gene (Nolasco et al., 2009) associated with different symptoms (Hancevic et al., 2013).

In Greece, the citrus industry represents an important branch of the economy, with an annual production of 1.3 million t; citrus crops are cultivated in 58 000 ha consisting mainly of oranges [Citrus sinensis (L.) Osbeck] (70%) and lemons (17%), grafted onto sour orange rootstock (Hellenic Statistic Authority, 2006). CTV is regulated as a quarantine pathogen in the country; all areas of citrus cultivation are surveyed, and eradication measures are applied when the virus is found (Dimou and Coutretsis, 2009). CTV was firstly detected in 2000, in imported sweet orange cv. Lane Late trees grafted on Carrizo citrange, in Argolida (North East Peloponnese) (Dimou et al., 2002) and Chania (Crete) (Dimou and Coutretsis, 2009). Analyses of the nucleotide sequences of the p20 gene of these isolates showed high homology with the Spanish mild T385 isolate (Varveri, 2006; Shegani et al., 2012). A simi-

lar isolate was collected in 2005, from sweet orange trees of the cv. Washington navel and Navelina in Arta prefecture (North West Greece) (Barbarossa et al., 2007). In 2007, CTV was detected in an orchard in Scala Laconias (South Peloponnese), and subsequently in 2008, in Attiki and Chalkidiki, in greenhouses producing ornamental citrus Calamondin [Citrofortunella microcarpa (Bunge.) Wijnands]. In 2009, a more virulent strain was obtained from old lemon and mandarin trees from the "lemonwood" (an area with ca. 100-year-old lemon trees) in Poros (North East Peloponnese), and this strain clustered in group 3b (Nolasco et al., 2009) according to its CP and p20 gene sequence analysis (Malandraki et al., 2011). Although quarantine measures were applied, limited dissemination has occurred from the initial foci in Argolida and Chania (Dimou and Coutretsis, 2009).

Typing of CTV strains is considered a key element for predicting disease spread and adopting more efficient control strategies (Moreno *et al.*, 2008). In this study, SSCP analysis of the CP gene of CTV and specific sequencing of selected variants was used to characterize the population structure of CTV isolates occurring in the main disease foci in Greece.

#### **Materials and methods**

#### Virus isolates

Ten CTV isolates were initially used in the study in order to select distinct isolates for further molecular analysis. Most isolates were obtained from field plants; five from sweet orange trees in Chania, Crete (designated GR 1, 4-7) and two from lemon trees of the lemonwood in Poros (GR 8, 9). CTV isolate GR3 was obtained from a Mexican lime seedling grafted with a sweet orange isolate from Argolida that was previously characterized by Varveri (2006), and was maintained in an insect-proof greenhouse at the Benaki Phytopathological Institute (Athens, Greece). Field plants infected with these isolates were originally detected by the Greek quarantine system and sampled before eradication. Two isolates (CTV 10 and 11) from Calamondin plants originating from Italy were also included in the study. The presence of CTV in the infected plant material was confirmed in double antibody sandwich enzyme linked immunosorbent assay (DAS-ELISA) tests with polyclonal antibodies (Agdia Biofords) according to the manufacture's procedures.

# IC/RT-PCR, RT-PCR or PCR amplification of the coat protein gene

One step Reverse Transcription (RT)-Polymerase Chain Reaction (PCR) and Immunocapture (IC) RT-PCR were performed using the primers CTV1 (5'-ATGGACGACGAAACAAAGAA-3') and CTV10 (5'-ATCAACGTGTGTTGAATTTCC-3') amplifying a 672 bp product including the whole CP gene (Papayannis *et al.*, 2007).

Total RNA was extracted from fresh bark tissue (0.1 g) using TRIzol reagent (Invitrogen) according to the manufacturer's protocol. Subsequently, a single step RT-PCR was performed in 50  $\mu$ L of a reaction mixture containing 2 mg of RNA, 10 mM Tris (pH8.8), 50 mM KCl, 0.08% Nonidet P-40, 2 mM MgCl<sub>2</sub>, 200 nM each of the primers, 200  $\mu$ M of each dNTP, 1U DreamTaq DNA polymerase (Thermo Scientific), 7.5U RNAguard (Amersham Pharmacia) and 7.5U MuLV reverse transcriptase (Perkin Elmer). RT was performed for 45 min at 38°C followed by PCR amplification with 2 min at 94°C, 30 cycles at 92°C for 30 s, 52°C for 30 s and 72°C for 45 s and finally 5 min at 72°C. PCR-products were separated in agarose gel electrophoresis (1%), stained with ethidium bromide and visualized under UV light. For IC/RT-PCR, RT-PCR tests were performed in tubes previously coated with anti-CTV IgG and ELISA extracts of the CTV infected samples without the addition of total RNA in the RT-PCR reaction mixture (Nolasco et al., 2002). PCR tests were also applied to confirm the transformation of *Escherichia coli* colonies, using the same primers and conditions as described for the IC/RT-PCR and RT-PCR.

#### SSCP analysis of RT-PCR or PCR products

SSCP analysis was performed on the amplicons obtained by RT-PCR (of infected tissues) and on the PCR products (of recombinant *E. coli* colonies), in order to separate distinct virus variants. The amplified product (1 to 3  $\mu$ L) was mixed with 9  $\mu$ L of denaturing buffer [95% formamide, 20 mM EDTA, pH 8, 0.05% bromophenol blue) placed for 5 min at 90°C and then chilled on ice. Denatured products were electrophoresed in a non-denaturing 8% polyacrylamide gel (Bio-Rad Mini-Protean II, Bio Rad Laboratories) for 3 h at 200 V, 4°C, using TBE (89 mM Tris-Borate, 2 mM EDTA, pH 8) as buffer (Rubio *et al.*, 1996). The gels were stained with GelStar (Lonza Bioscience) and visualized under UV light. PCR products displaying different SSCP patterns were considered different genomic variants (haplotypes) (Kong *et al.*, 2000).

#### **Cloning and sequencing**

amplicons were TA-ligated into Selected pTZ57R/T vector (InsTAclone PCR Cloning Kit, Thermo Fisher Scientific Inc.) and competent E. coli cells (Mach1 -T1, Invitrogen) were transformed according to manufacturer's instructions. Transformed colonies were selected by  $\alpha$ -complementation on plates supplied with X-gal and ampicillin, according to standard procedures (Sambrook et al., 1989). Transformation was confirmed by PCR amplification of the white colonies using primers CTV1/CTV10, and distinct clones were identified by SSCP analysis of at least 15 colonies per CTV isolate. Selected colonies were transferred to liquid medium and the plasmids harbouring selected variants were purified using the GeneJET Plasmid Miniprep Kit (Thermo Fisher Scientific Inc.) according to manufacturer's instructions.

Sequencing reactions were carried out at CC-MAR, Universidade do Algarve, using the forward and reverse primers. Sequence alignment and clustering were performed using MEGA5 software package (Tamura et al., 2011). Searches for recombination events among the sequences were carried out using the software package RDP (Martin et al., 2010), which implements several algorithms for detecting recombination. The obtained sequences (Table 1) were analysed with the following reference sequences of the CP gene from the GenBank: 19-121, Portugal (AF184114); T36, Florida (M76485); VT, Israel (U56902); SY568, California (AF001623); and T30, Florida (AF260651). Sequences of B249, Venezuela and T3, Florida were provided by C.L. Niblett. These sequences are representative of each of the CTV phylogenetic groups proposed by Nolasco et al. (2009).

#### Results

#### IC/RT-PCR, RT-PCR and SSCP analysis of CTV isolates

IC/RT-PCR and RT-PCR of all samples resulted in amplicons of the expected size (672 bp) (results not shown). The preliminary assessment of these amplicons showed four different SSCP patterns (Figure 1a). No variability was observed among field isolates



**Figure 1.** SSCP patterns obtained from the RT-PCR amplicons of (a) infected plants (GR3, GR6, GR9, CTV11) and (b) their haplotypes subsequently sequenced. *Citrus tristeza virus* (CTV) isolates: GR3, Mexican lime grafted with the isolate from Argolida (haplotype 11); GR6, Sweet orange from Chania (haplotypes 23, 30); GR9, Lemon from Poros (haplotypes 28, 16, 15, 17); and CTV11, Calamondin from Italy (haplotypes 27, 18).

originating from the one area and for each isolate in the patterns of the amplicons produced by either IC/ RT-PCR or RT-PCR (results not shown).

The CP genes of the CTV isolates GR3, GR6, GR9 and CTV11, representing different SSCP patterns (Figure 1a), were subsequently cloned. A clear predominance of a single SSCP pattern consisting of two conspicuous bands was obvious in all isolates; for isolate GR3 this was the sole haplotype detected (GR3-11). CTV isolate GR6 consisted of two haplotypes; GR6-30 represented the one observed for the 87% of the analysed clones, while GR6-23 was observed in only 13% of them. Clones of isolate GR9 consisted of 60% of the haplotype GR9-15, 2% of the GR9-28, while one haplotype (6.5%) for each of the GR9-16 and GR9-17 was also present. Isolate CTV11 was difficult to clone; among 40 clones tested only four were successfully transformed showing two haplotypes CTV11-27 (75%) and CTV11-18 (25%) (Figure 1b).

## Sequences and comparative analysis of the CTV coat protein

From each isolate one clone representing each haplotype obtained was chosen for sequencing. Nine sequences obtained were submitted to the GenBank with the accession numbers: KF196264, KF196265, KF196266, KF196267, KF196268, KF196269, KF196270, KF196271, KF196272 (Table 1). After excising the two

terminal parts 20 bases long that corresponded to the primers, search for evidence of recombination was carried out among the new sequences, and the representative international isolates that were considered for comparisons. A strong recombination signal was obtained with several of the algorithms implemented in the RDP software for sequence GR9-16 in the stretch between positions 440 and 630. According to RDP, the backbone of GR9-16 derives from GR9-17 (group 3b) and the recombination stretch is close to the group 5 Greek sequences. Except for GR9-16, the remaining CTV sequences were aligned, their pairwise distances determined according to the Kimura 2 parameters model, and the resulting distance matrix was used to reconstruct a phylogenetic tree (Figure 2). GR9-16 was not included, as this evolutionary model is not suitable for calculating distances among sequences evolving through recombination.

All sequences matched with the expected sequence of the CTV coat protein. However, they were distributed clearly among three CP gene clusters, with high bootstrap values. The only SSCP variant detected in isolate GR3 (GR3-11) was grouped with the reference sequence from phylogenetic group M, while both variants of isolate GR6 (GR6-23, GR6-30) were clustered in group 5. The CP sequence variants of isolate CTV11 (CTV11-27, CTV11-18) were grouped into two different clusters (groups 5 and 3b), while those of isolate GR9 (GR9-15, GR9-17, GR9-28) were in group 3b.

**Table 1.** The GeneBank accession numbers and geographical locations in Greece and Italy of nine isolates and variants of *Citrus tristeza virus* (CTV).

CTV isolate	Host	Geographic origin	Haplo- type	Accession No.
GR3	Mexican lime	Argolida	11	KF196264
GR6	Orange	Chania	23	KF196269
			30	KF196272
GR9	Lemon	Poros	15	KF196265
			16	KF196266
			17	KF196267
			28	KF196271
CTV11	Calamondin	Italy	18	KF196268
			27	KF196270

#### Discussion

CTV epidemiology in a region is highly influenced by the susceptibility of the cultivated citrus varieties, the composition and the population dynamic of the aphid vector fauna, and the predominant virus isolates (Moreno et al., 2008). In Greece, the prevalent use of sour orange rootstock provides very many CTV susceptible trees, while the aphid vectors A. gossypii, A. spiraecola and T. aurantii are widespread (Tsitsipis et al., 2007). Therefore, the strain profile of CTV is a key factor for predicting future spread of the disease. In this study, SSCP analysis was employed to assess the population structure and sequence variability of the CP gene of three Greek and one Italian CTV isolates. Due to the guarantine status of CTV in Greece, a limited number of isolates were available; however the genetic diversity of the virus was still obvious.

Two of the field isolates analysed, Poros (GR9) and the Italian (CTV-11) isolate, had typical quasispecies structures consisting of sequence variants, with the one being predominant (Holland *et al.*,



**Figure 2.** Neighbour-joining tree obtained from the matrix of pair-wise distances (nucleotide, Kimura 2-parameter) between the CP gene sequences of nine isolates of *Citrus tristeza virus* (CTV) from Greece and Italy. Bootstrap values obtained from 1000 replications are presented. Nodes with bootstrap values lower than 70% are not individualized. Group names as proposed by Nolasco *et al.* (2009) are indicated on the right. Nucleotide distance is represented in the horizontal bar.

1982). The prevalent haplotype in the Poros isolate was classified in group 3b, in agreement with previous studies (Malandraki et al., 2011). However, we also detected, at a low concentration, a recombinant that implies the natural presence of variants of group 5, as well. The 100-year-old lemonwood may represent a distinct environment for CTV strain variability and spread. Infected lemon plants grafted on sour orange remain symptomless with low CTV titre and uneven distribution of the virus (Moreno and Garnsey, 2010). Long standing CTV presence within individual trees may therefore remain unnoticed, favouring reinfection and changes such as mutation or recombination (Rubio et al., 2001; Martin et al., 2009; Roy and Brlansky, 2009; Melzer et al., 2010; Biswas et al., 2012). On the other hand, the poor colonisation of lemon by A. gossypii (Barbagallo et al., 2007) and its low susceptibility to aphid transmission of CTV (Hermoso de Mendoza et al., 1988) may be the reason for the restricted virus spread within the lemonwood population (Malandraki et al., 2011).

In Chania, CTV was introduced (Dimou and Coutretsis, 2009) with propagation material infected with a strain closely related to the Spanish T385 (Varveri, 2006; Shegani et al., 2012), which clusters in group M (Nolasco et al., 2009). In our analysis, however, a field isolate from the same area (GR6) consisted of only haplotypes of group 5, suggesting that this variant was either overlooked in previous studies or represents another virus introduction. Severe CTV strains can be masked by milder ones (Broadbent et al., 1996; Brlansky et al., 2003). In this area, natural spread of CTV was reported (Dimou and Coutretsis, 2009; Shegani et al., 2012) suggesting virus adaptation to the local aphid vectors (Moreno et al., 2008). Aphid mediated transmission may alter strain profile as CTV subisolates differ in their transmissibility (Broadbent et al., 1996; Ayllón et al., 1999; Brlansky et al., 2003; Huang et al., 2005; Nolasco et al., 2008). The A. gossypii that occurs in the area has been associated with long-range dissemination of the virus (Gottwald et al., 1999) that may maximise this bottleneck (Nolasco et al., 2008) and may have contributed to the emergence of haplotypes of group 5 in preference to the mild strains previously reported.

The slow evolution of CTV has resulted in low strain diversity, despite the extensive global exchange of budwood (Silva *et al.*, 2012). Sequence variants from groups M and 3b are dominant in the Mediterranean Basin, reflecting the major CTV sources

of Spain for group M, and Israel for 3b (Djelouah et al., 2009). In agreement with this, the original isolate introduced with Spanish budwood in Argolida and Chania was previously included in the Spanish T386 isolate (Varveri, 2006; Dimou and Coutretsis, 2009; Shegani et al., 2012) that clusters in group M with mild isolates (Nolasco et al., 2009). In our studies of the isolate from Argolida (GR3) we detected only a variant belonging to group M. However, in the isolate from Chania we only detected variants of group 5, which are not reported by other recipients of Spanish propagation material (Lbida et al., 2004). Additionally, the predominant variant of the Italian Calamondin isolate (CTV-11) was also clustered in group 5, although only isolates corresponding to groups 3b and M were previously identified in Italian plant material (Barbarossa and Savino, 2004; Davino et al., 2005; Abou Kubaa et al., 2012). In the Mediterranean basin, strains clustering in group 5 are present in the field in Croatia, Cyprus, Morocco, Portugal, Syria and Tunisia (Djelouah et al., 2009; Abou Kubaa et al., 2012). The presence of these variants in Greece may reflect a different CTV introduction from a location at which diverse CTV isolates occur.

Overall, our analysis of the three Greek and one Italian isolates pointed to the predominance of variants clustered in groups 3b and 5, which harbour severe SP and SY isolates, as do isolates VT from Israel and B249 from Venezuela (Nolasco et al., 2009). The presence of these variants, even at low incidence, may explain the citrus decline symptoms observed in Greece, in areas where only mild strains were previously reported (Dimou and Coutretsis, 2009). For CTV, the structure of a complex virus population, and occasionally the major (Sambade et al., 2002) or minor (Cerni et al., 2008) components of the mixture, determine field symptoms. On the other hand, the presence of potentially aggressive strains may be critical for the development of future epidemics, as these strains show greater fitness and become predominant in mixed infected plants (Moreno et al., 2008), while their spread seems to be favoured by T. citricida over the mild strains (Rocha-Rena et al., 1995; Niblett et al., 2000; Halbert et al., 2004; Matos et al., 2013).

In Greece, quarantine measures are constantly applied in order to limit CTV spread. However, the spread of *T. citricida* will challenge the ability of our agricultural systems to quickly shift to resistant root-stocks, and particularly the efficiency of our quar-

antine and certification programmes against the increasing threats posed by severe CTV strains.

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### Literature cited

- Abou Kubaa R., A.M. D'Onghia, K. Djelouah, V. Savino and M. Saponari, 2012. Characterization of *Citrus tristeza virus* isolates recovered in Syria and Apulia (southern Italy) using different molecular tools. *Phytopathologia Mediterranea* 51, 496–504.
- Ayllón M.A., L. Rubio, A. Moya, J. Guerri and P. Moreno, 1999. The haplotype distribution of two genes of *Citrus tristeza virus* is altered after host change or aphid transmission. *Virology* 255, 32–39.
- Barbagallo S., G. Cocuzza, P. Cravedi and S. Komazki. 2007. IPM case studies: deciduous fruit trees. In: *Aphids as Crop Pests* (H.F. van Emden, R. Harrington, ed.), CAB International, Wallingford, Oxon, UK, 651–676.
- Barbarossa L. and V. Savino, 2004. Genotype characterization of Apulian Citrus tristeza virus isolates. Journal of Plant Pathology 86, 309.
- Barbarossa L., G. Loconsole and C. Vovlas, 2007. Virus and virus-like diseases of citrus in Epirus. *Journal of Plant Pathology* 89, 273–276.
- Biswas K.K., A. Tarafdar, S. Diwedi and R.F. Lee, 2012. Distribution, genetic diversity and recombination analysis of *Citrus tristeza virus* of India. *Virus Genes* 45, 139–148.
- Brlansky R.H., V.D. Damsteegt, D.S. Howd and A. Roy, 2003. Molecular analyses of *Citrus tristeza virus* subisolates separated by aphid transmission. *Plant Disease* 87, 397–401.
- Broadbent P., R.H. Brlansky and J. Indsto, 1996. Biological characterization of Australian isolates of *Citrus tristeza virus* and separation of subisolates by single aphid transmissions. *Plant Disease* 80, 329–333.
- Cambra M., M.T. Gorris, C. Marroquín, M.P. Román, A. Olmos, P.C. Martínez, A.H. Hermoso de Mendoza, A. López and L. Navarro, 2000. Incidence and epidemiology of *Citrus tristeza virus* in the Valencian community of Spain. *Virus Research* 71, 85–95.
- Cerni S., J. Ruscic, G. Nolasco, Z. Gatin, M. Krajacic and D. Skoric, 2008. Stem pitting and seedling yellows symptoms of *Citrus tristeza virus* infection may be determined by minor sequence variants. *Virus Genes* 36, 241–249.
- Davino S., L. Rubio and M. Davino, 2005. Molecular analysis suggests that recent *Citrus tristeza virus* outbreaks in Italy were originated by at least two independent introductions. *European Journal of Plant Pathology* 111, 289–293.

- Dimou D. and P. Coutretsis, 2009. Citrus tristeza virus (CTV) in Greece: historical review. In: Citrus Tristeza Virus and Toxoptera citricidus: a Serious Threat to the Mediterranean Citrus Industry. (A.M. D'Onghia, K. Djelouah, C.N. Roistacher, ed.), CIHEAM-IAMB, Bari, Italy, 2009, 69–72, Options Méditerranéennes: Série B. Etudes et Recherches 65.
- Dimou D., J. Drossopoulou, E. Moschos, C. Varveri and F. Bem, 2002. First report of *Citrus tristeza virus* in Greece. *Plant Disease* 86, 329.
- Djelouah K., A.M. D'Onghia, G. Nolasco, S. Cerni, F. Fonseca, C. Santos and G. Silva, 2009. Diversity of the coat protein gene of *Citrus tristeza virus* (CTV) in the Mediterranean region. In: *Citrus Tristeza Virus and Toxoptera citricidus: a Serious Threat to the Mediterranean Citrus Industry*. (A.M. D'Onghia, K. Djelouah, C.N. Roistacher, ed.), CIHEAM-IAMB, Bari, Italy, 2009, 159–163, *Options Méditerranéennes: Série B. Etudes et Recherches* 65.
- Gottwald T.R., G.J. Gibson, S.M. Garnsey and M. Irey, 1999. Examination of the effect of aphid vector population composition on the spatial dynamics of *Citrus tristeza virus* spread by stochastic modeling. *Phytopathology* 89, 603–608.
- Halbert, S.E., H. Genc, B. Cevik, L.G. Brown, I.M. Rosales, K.L. Manjunath, M. Pomerinke, D.A. Davison, R.F. Lee and C.L. Niblett, 2004. Distribution and characterization of *Citrus tristeza virus* in south Florida following establishment of *Toxoptera citricida*. *Plant Disease* 88, 935–941.
- Hancevic K., S. Cerni, G. Nolasco, T. Radic, K. Djelouah and D. Skoric, 2013. Biological characterization of *Citrus tristeza virus* monophyletic isolates with respect to p25 gene. *Physiological and Molecular Plant Pathology* 81, 45–53.
- Hellenic statistic authority, 2006. http://www.statistics.gr
- Hermoso de Mendoza A., J.F. Ballester-Olmos and J.A. Pina, 1988. Comparative aphid transmission of a common citrus virus isolate and a seedling yellows-citrus tristeza virus isolate recently introduced in Spain. In: *Proceedings*, 10th Conference of the International Organization of Citrus Virologists, 1986, Riverside, CA, USA, 68–70.
- Hermoso de Mendoza A., A. Alvarez, J.M. Michelena, P. Gonzáles and M. Cambra, 2008. Spread, biology and natural enemies of *Toxoptera citricida* (Kirkaldy) (Hemiptera, Aphididae) in Spain. *Boletín de Sanidad Vegetal Plagas* 34, 77–87 (in Spanish).
- Holland J.J., K. Spindler, F. Horodyski, E. Grabau, S. Nichol and S. VandePol, 1982. Rapid evolution of RNA genomes. *Science* 215, 1577–1582.
- Huang Z., P.A. Rundell, X. Guan and C.A. Powell, 2005. Evaluation of the transmission of different field sources of *Citrus tristeza virus* and the separation of different genotypes by single brown citrus aphids. *Horticultural Science* 40, 687–690.
- Ilharco F.A., C.R. Sousa-Silva and A. Alvarez Alvarez, 2005. First report of *Toxoptera citricidus* (Kirkaldy) in Spain and continental Portugal (Homoptera, Aphidoidea). Agronomia Lusitana 51, 19–21.
- Kong P., L. Rubio, M. Polek and B.W. Falk, 2000. Population structure and genetic diversity within California *Citrus tristeza virus* (CTV) isolates. *Virus Genes* 21, 139–145.
- Lbida B., F. Fonseca, C. Santos, M. Zemzami, A. Bennani and G. Nolasco, 2004. Genomic variability of *Citrus tristeza vi*-

rus (CTV) isolates introduced into Morocco. *Phytopatholo*gia Mediterranea 43, 205–210

- Malandraki I., E. Marouli and C. Varveri. 2011. New isolates of *Citrus tristeza virus* naturally occurring in old lemon and mandarin trees in Greece. *New Disease Reports* 23, 2.
- Martin D.P., P. Lemey, M. Lott, V. Moulton, D. Posada, P. Lefeuvre, 2010. RDP3: a flexible and fast computer program for analyzing recombination. *Bioinformatics* 26, 2462–2463.
- Martin S., A. Sambade, L. Rubio, M.C. Vives, P. Moya, J. Guerri, S.F. Elena and P. Moreno, 2009. Contribution of recombination and selection to molecular evolution of *Citrus tristeza virus*. *Journal of General Virology* 90, 1527–1538.
- Matos L.A., M.E. Hilf, X.A. Cayetano, A.O. Feliz, S.J. Harper and S.Y. Folimonova, 2013. Dramatic change in *Citrus tristeza virus* populations in the Dominican Republic. *Plant Disease* 97, 339–345.
- Melzer M.J., W.B. Borth, D.M. Sether, S. Ferreira, D. Gonsalves and J.S. Hu, 2010. Genetic diversity and evidence for recent modular recombination in Hawaiian *Citrus tristeza virus. Virus Genes* 40, 111–118.
- Moreno P. and S.M. Garnsey, 2010. Citrus tristeza diseases-A worldwide perspective. In: *Citrus Tristeza Virus. Complex and Tristeza Diseases* (A.V. Karasev, M.E. Hilf, ed). APS Press, MN, USA, 27–52.
- Moreno P., S. Ambrós, M.R. Albiach-Martí, J. Guerri and L. Peña, 2008. *Citrus tristeza virus:* a pathogen that changed the course of the citrus industry. *Molecular Plant Pathology* 9, 251–268.
- Niblett C.L., H. Genc, B. Cevik, S. Halbert, L. Brown, G. Nolasco, B. Bonacalza, K.L. Manjunath, V.J. Febres, H.R. Pappu and R.F. Lee, 2000. Progress on strain differentiation of *Citrus tristeza virus* and its application to the epidemiology of citrus tristeza disease. *Virus Research* 71, 97–106.
- Nolasco G., Z. Sequeira, C. Soares, A. Mansinho, A.M. Bailey and C.L. Niblett, 2002. Asymmetric PCR ELISA: increased sensitivity and reduced costs for the detection of plant viruses. *European Journal of Plant Pathology* 108, 293–298.
- Nolasco G., F. Fonseca and G. Silva, 2008. Occurrence of genetic bottlenecks during *Citrus tristeza virus* acquisition by *Toxoptera citricida* under field conditions. *Archives of Virology* 153, 259–271.
- Nolasco G., C. Santos, G. Silva and F. Fonseca, 2009. Development of an asymmetric PCR-ELISA typing method for *Citrus tristeza virus* based on the coat protein gene. *Journal of Virological Methods* 155, 97–108.
- Papayiannis L.C., C. Santos, A. Kyriakou, T. Kapari and G. Nolasco, 2007. Molecular characterization of *Citrus tristeza virus* isolates from Cyprus on the basis of the coat protein gene. *Journal of Plant Pathology* 89, 291–295.
- Rocha-Pena M.A., R.F. Lee, R. Lastra, C.L. Niblett, F.M. Ochoa-Corona, S.M. Garnsey and R.K. Yokomi, 1995. *Citrus tristeza virus* and its aphid vector *Toxoptera citricida*: threats to

citrus production in the Caribbean and Central and North America. *Plant Disease* 79, 437–445.

- Roy A. and R.H. Brlansky, 2009. Population dynamics of a Florida *Citrus tristeza virus* isolate and aphid-transmitted subisolates: Identification of three genotypic groups and recombinants after aphid transmission. *Phytopathology* 99, 1297–1306.
- Rubio L., M.A. Ayllon, J. Guerri, H. Pappu, C. Niblett and P. Moreno, 1996. Differentiation of *Citrus tristeza closterovirus* (CTV) isolates by single-strand conformation polymorphism analysis of the coat protein gene. *The Annals of Applied Biology* 129, 479–489.
- Rubio L., M.A. Ayllon, P. Kong, A. Fernandez, M.L. Polek, J. Guerri, P. Moreno and B.W. Falk, 2001. Genetic variation of *Citrus tristeza virus* isolates from California and Spain: evidence for mixed infections and recombination. *Journal* of Virology 75, 8054–8062.
- Sambade A., L. Rubio, S.M. Garnsey, N. Costa, G.W. Müller, M. Peyrou, J. Guerri and P. Moreno, 2002. Comparison of viral RNA populations of pathogenically distinct isolates of *Citrus tristeza virus*: application to monitoring crossprotection. *Plant Pathology* 51, 257–265.
- Sambrook J., E.F. Fritsch and T. Maniatis, 1989. Molecular Cloning: A Laboratory Manual. 2nd ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, USA.
- Shegani M., D. Tsikou, A. Velimirovic, H. Afifi, A. Karayanni, A. Gazivoda, K. Manevski, I. Manakos and I.C. Livieratos, 2012. *Citrus tristeza virus* on the island of Crete: a survey and detection protocol applications. *Journal of Plant Pathol*ogy 94, 71–78.
- Silva G., N. Marques and G. Nolasco. 2012. The evolutionary rate of *Citrus tristeza virus* ranks among the rates of the slowest RNA viruses. *Journal of Virology* 93, 419–429.
- Tamura K., D. Peterson, N. Peterson, G. Stecher, M. Nei and S. Kumar, 2011. MEGA5: Molecular evolutionary genetics analysis using Maximum Likelihood, Evolutionary Distance, and Maximum Parsimony Methods. *Molecular Biol*ogy and Evolution 28, 2731–2739.
- Tsitsipis J.A., N.I. Katis, J.T. Margaritopoulos, D.P. Lykouressis, A.D. Avgelis, I. Gargalianou, K.D. Zarpas, D.C. Perdikis and A. Papapanayotou, 2007. A contribution to the aphid fauna of Greece. *Bulletin of Insectology* 60, 31–38.
- Varveri C., 2006. Molecular detection and characterization of *Citrus tristeza virus* in Greece. *Phytopathologia Mediterranea* 45, 68 (abstract).
- Yokomi R.K., R. Lastra, M.B. Stoetzel, V.C. Damsteegt, R.F. Lee, S.M. Garnsey, T.R. Gottwald, M.A. Rocha-Pena and C.L. Niblett, 1994. Establishment of the brown citrus aphid (Homoptera: Aphididae) in Central America and the Caribbean Basin and transmission of *Citrus tristeza virus*. *Journal of Economic Entomology* 87, 1078–1085.

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