The presence of *Citrus tristeza virus* (CTV) has previously been reported in citrus growing regions of Turkey. All serologically and biologically characterized isolates including Igdır, which was the first identified CTV isolates from Turkey, were considered mild isolates. In this study, molecular characteristics of the Igdır isolate were determined by different methods. Analysis of the Igdır isolate by western blot and BD-RT-PCR assays showed the presence of MCA13 epitope, predominantly found in severe isolates, in the Igdır isolate revealing that it contains a severe component. For further characterization, the coat protein (CP) and the RNA-dependent RNA polymerase (RdRp) genes representing the 3' and 5' half of CTV genome, respectively, were amplified from dsRNA by RT-PCR. Both genes were cloned separately and two clones for each gene were sequenced. Comparisons of nucleotide and deduced amino acid sequences showed that while two CP gene sequences were identical, two RdRp clones showed only 90% and 91% sequence identity in their nucleotide and amino acid sequences, respectively, suggesting a mixed infection with different strains. Phylogenetic analyses of the CP and RdRp genes of Igdır isolate with previously characterized CTV isolates from different citrus growing regions showed that the CP gene was clustered with NZRB-TH30, a resistance breaking isolate from New Zealand, clearly showing the presence of severe component. Furthermore, two different clones of the RdRp gene were clustered separately with different CTV isolates with a diverse biological activity. While the RdRp-1 was clustered with T30 and T385, two well-characterized mild isolates from Florida and Spain, respectively, the RdRp-2 was most closely related to NZRB-G90 and NZRB-TH30, two well-characterized resistance breaking and stem pitting (SP) isolates from New Zealand confirming the mixed infection. These results clearly demonstrated that the Igdır isolate, which was previously described as biologically a mild isolate, actually contains a mixture of mild and severe strains.

**Keywords**: *Citrus tristeza virus*, Coat protein, Phylogenetic analysis, RNA-dependent RNA polymerase, Strain identification

Tristeza disease caused by *Citrus tristeza virus* (CTV), is one of the most destructive and economically important viral diseases limiting commercial citrus production worldwide (Bar-Joseph et al., 1989; Rocha-Pena et al., 1995). The disease is now distributed in almost all citrus growing regions of the world including Turkey (Moreno et al., 2008; Rocha-Pena et al., 1995). CTV is able to infect most species, varieties and hybrids of *Citrus* as well as some close relatives of *Citrus* (Mueller and Garney, 1984). While most commercial *Citrus* varieties are sensitive to CTV, a citrus relative, *Poncirus trifoliata*, and some of its hybrids are resistant to this virus. Therefore, they were used as rootstock in some *Citrus*-growing regions including the Aegean and the Black Sea regions of Turkey. However, some isolates of CTV breaking the resistance in *P. trifoliata* were recently identified in New Zealand (Harper et al., 2009; Harper et al., 2010).

CTV is transmitted readily by grafting and several aphid species including *Toxoptera citricida*, *Aphis gossypii* and *Aphis spiraecola* are able to transmit the virus in a semi-persistent manner in nature (Roistacher and Bar-Joseph, 1987). While *T. citricida* is the most efficient vector of CTV (Yokomi et al., 1994) and widely distributed in many *Citrus*-growing regions, *A. gossypii*, acts as a major vector, in some regions including Turkey where *T. citricida* is still absent. The recent detection and spread of *T. citricida* in Portugal and Spain (Ilharco et al., 2005) has become a significant concern for many Mediterranean countries such as Turkey where the majority of citrus are grown on CTV-sensitive sour orange rootstocks.

CTV is a complex virus consisting of various strains causing a variety of symptoms in different *Citrus* hosts. Based on the symptoms inflicted on *Citrus* scions and rootstocks, CTV strains are divided into five major groups including mild (M), vein clearing in Mexican lime, seedling...
yellows (SY) in sour orange, lemon or grapefruit, quick decline (QD) in all Citrus species grafted on sour orange rootstock, and stem pitting on grapefruit (SP-G) and on sweet orange (SP-O) regardless of the rootstock. Symptoms of CTV isolates are determined experimentally using a previously established standardized set of Citrus indicator plants (Garnsey et al., 1987).

CTV is a single-stranded positive-sense RNA virus belonging to the genus Closterovirus in the Closteroviridae family. The long thread-like, flexuous, filamentous particles of CTV are about 2000 nm by 11 nm (Bar-Joseph et al., 1979; Bar-Joseph and Lee, 1990) and consist of one single-stranded positive-sense RNA molecule encapsidated with the major capsid protein (CP) and minor CP (Febres et al., 1996). Sequencing of the complete genome of CTV isolates from different geographical origins and with various biological properties confirmed that CTV has one of the largest plant virus genomes ranging from 19226 to 19306 nt. The genome of CTV was organized into 12 open reading frames (ORF) potentially encoding 17 protein products, plus the 3' and 5' untranslated regions (UTRs) (Albiach-Martí et al., 2000; Harper et al., 2009; Harper et al., 2010; Karavev et al., 1995; Mawassi et al., 1996; Pappu et al., 1994; Suastika et al., 2001; Vives et al., 1999; Yang et al., 1999). The 25 kDa protein is the major capsid protein (Sekiya et al., 1991) encapsidating about 95% of the CTV genome. Therefore, the CP gene has been used as target for various methods developed for detection of CTV and strain identification including ELISA, western blot, peptide mapping, RT-PCR, hybridization and real-time RT-PCR. In addition the CP genes of a large number of CTV isolates from different geographical regions and biological properties have been cloned and sequenced (Mawassi et al 1993; Pappu et al., 1993; Roy et al., 2003). Since a correlation between geographical origin and the biological characteristics of CTV isolates and their CP gene sequences was observed (Herrera-Isidrón et al., 2009), sequencing and the phylogenetic analysis of the CP gene was generally used for identification of newly identified and biologically uncharacterized CTV isolates (Niblett et al., 2000; Nolasco et al., 2009).

In Turkey, CTV was first detected symptomatically around Adana by Norman in 1963 and in Aegean Region by Özalp and Azari in 1967 (Baloglu and Birisik, 2009; Yılmaz, 1999). The presence of CTV was demonstrated experimentally by serological and biological assays (Baloglu, 1988) and the first Turkish CTV isolate, İlgdr, was identified (Baloglu, 1988). Since then İlgdr and a limited number of other CTV isolates from Eastern Mediterranean region were characterized by serological and biological assays and these studies indicated that only mild isolates were present in Turkey (Baloglu 1988; Korkmaz, 2002; Yılmaz and Baloglu, 1998). After the detection of CTV in the Eastern Black Sea region by ELISA and RT-PCR (Korkmaz et al., 2006), the presence of CTV in all citrus growing regions of Turkey was confirmed. Therefore, the presence and distribution of CTV isolates in different citrus growing regions of Turkey is largely known and their serological and biological properties are determined to some extent. However, there is no sequence information available for Turkish CTV isolates and their molecular characteristics are completely unknown. In this study, molecular characteristics of the first identified CTV isolate, İlgdr, was determined by molecular tests used for strain identification and sequencing and phylogenetic analysis of the CP and the RdRp genes.

Materials and Methods

Virus isolate. The İlgdr isolate of CTV originally obtained from a sweet orange tree grafted on sour orange rootstock in a commercial orchard in İlgdr village of Mersin, Province located in the Eastern Mediterranean region of Turkey was used in this study. The isolate was originally found in the 1980s, identified by ELISA (Baloglu 1988). It was later grafted and maintained on Mexican lime to date. Tissue samples of Mexican lime grafted with CTV İlgdr isolate were obtained from the Western Mediterranean Agricultural Research Institute (BATEM) in Antalya, Turkey where it is currently maintained as a reference isolate in a greenhouse. The samples of T30 and T36 used as mild and severe strains, respectively, were kindly provided by Richard F. Lee of USDA-ARS National Clonal Germplasm Repository for Citrus and Dates.

Oligonucleotide primers. The primers specific to the CP gene were designed based on the conserved sequences at the 5' end (BC24 5'-ATGGAGCCGAAAACAAGAA-3') and 3' end (BC25 5'-TCAACGTGTTGTTGAATTTC-3') of the CTV CP gene. Since the 5' and 3' ends of the RdRp gene were not conserved the primers specific to RdRp gene were designed based on conserved regions of located upstream (BC42 5'-CCTACTGAATATAAGGGTAG-3') and downstream (BC43 5'-CTCGCGAAGGGCAACAT-3') of the RdRp gene. The primers for the bidirectional PCR were previously reported (Çevik et al., 1996).

Amplification of the CP and RdRp genes by RT-PCR. dsRNA was isolated using a previously reported dsRNA isolation method (Morris and Dodds, 1979) with minor modification. The CP and RdRp genes of the İlgdr isolate were amplified from dsRNA by a two-step reverse transcription-polymerase chain reaction (RT-PCR) assay using AMV reverse transcriptase and Pfu DNA polymerase (Promega, USA). First, 5 ml of dsRNA was denatured at 95°C for 5 min and quickly chilled on ice. cDNA was synthesized
Western blot analysis. Bark tissue from Mexican limes seedlings infected with Igdır isolate was ground to powder in liquid nitrogen using a mortar and passel. About 200 mg ground tissue was homogenized in 0.5 ml 2X extraction buffer containing 0.125 M Tris-HCl at pH 6.8, 4% SDS, 20% glycerol, and 10% 2-mercaptoethanol. The homogenate was incubated at 95°C for 5 min, cooled down to room temperature and briefly centrifuged to separate cellular debris. Supernatant was loaded onto a 12% SDS polyacrylamide gel and total protein was separated by electrophoresis, and transferred to a nitrocellulose membrane (Stratagene, USA) using a Mini Transblot (Bio-Rad, USA). The membrane was probed with the MCA13 monoclonal antibody (Promega, USA) (kindly provided by Richard F. Lee), followed by rabbit anti-mouse IgG conjugated with alkaline phosphatase (Promega, USA). The presence of the 25 kDa major CP was detected colorimetrically with a p-nitrophenol phosphate substrate (Sigma, USA).

Cloning of the CP and RdRp genes. The CP and RdRp genes amplified by RT-PCR from Igdır isolate were separated in 1% agarose gel, appropriate bands were excised from the gel and purified using the Qiaquick gel purification kit (Qiagen, Germany) according to the manufacturer's instructions. For making compatible ends for T-A cloning an adenine (A) overhang was added to the 3’ ends of the purified DNA by Taq DNA polymerase by A-tailing method. A-tailed PCR products (5 μl) were ligated into the pGEM-Teasy plasmid vector (Promega, USA) at 4°C for 16 hr using T4 DNA ligase and 1X ligation buffer (0.05 M Tris-HCl pH 8.0, 0.01 M MgCl₂, 1 mM ATP and 50 μg/ml bovine serum albumin). Competent cells of Escherichia coli strain JM109 were transformed with ligation mixture by heat shock at 42°C for 1 min. Transformed cells were plated and grown in LB medium (1% bacto-tryptone, 0.5% bacto-yeast extract, 0.5% NaCl) containing 100 μg/μl ampicillin, 25 μg/ml 5-bromo-4-chloro-indoly-β-D-galactoside (X-Gal) and 0.1 M isopropyl-beta-thio galactopyranoside (IPTG) at 37°C 16 hr. Recombinant colonies carrying pGEM-Teasy plasmid vector with the CP or RdRp genes were selected by blue/white screening on LB medium according to the manufacturer’s instructions. For each sample, at least five white colonies were screened by colony PCR and at least two colonies carrying the pGEM-T Easy plasmid with the CP and RdRp genes were identified. Plasmids were isolated from these colonies and the presence of the CP and RdRp genes in the plasmids were confirmed by EcoRI digestion.

Sequencing and sequence analysis of the CP and RdRp genes. The cloned CP and RdRp genes were sequenced in both directions by automated cycle sequencing using M13 forward and reverse primers. The sequences of the CP and RdRp genes were assembled and analyzed using Vector NTI Suite program (Invitrogen, USA). The sequences were compared with each other and other CP and RdRp gene sequences in the GenBank. Multiple sequence alignments of the full-length amino acid sequences of the CP and the RdRp genes were conducted by the AlignX Module of Vector NTI Suite. The phylogenetic analysis was performed by Clustal X2 program using neighbor joining algorithm and tested by a bootstrap analysis with 1000 replications. Constructed phylogenetic trees were visualized by TreeView program.

Results

Characterization of the Igdır isolate. Previously identi-
fied Idr isolate maintained in Mexican lime as reference mild isolate of CTV was tested by Western blot with MCA13 antibody and the BD-RT-PCR developed based on the MCA13 epitope in CP. Therefore, Western blot was used to determine if the Idr isolate react with MCA13 antibody, which is predominantly used for identification of severe, especially QD, strain of CTV. Western blot analysis of the Idr isolate was conducted along with T30 and T36, reference for mild and severe strains, respectively. Western blot analysis with MCA13 antibody showed that while T30 did not react with MCA13, T36 and the Idr isolate both reacted with MCA13 and a 25 kDa protein corresponding to the CP was detected in both samples. The result showed that the Idr isolate previously characterized as mild contains the MCA13 epitope associated with severe isolates (Fig. 1A).

The BD-RT-PCR method previously developed for strain identification based on a single nucleotide change in the MCA13 epitope of the CP gene was used to confirm the Western blot results, suggesting that the Idr isolate may contain a severe component. The BD-RT-PCR assay was conducted for the Idr isolate along with T30 and T36, reference isolates for mild and severe strains, respectively. Using the BD-RT-PCR method, a full-length CPG (700 bp) and a 400 bp fragment specific to mild strains were produced from the mild strain T30 (Fig. 2B). On the other hand about 700 bp fragment corresponding to the full-length CP gene as well as a 300 bp fragment specific to MCA13 positive severe strains were amplified from the severe QD strain T36 and the Idr isolate (Fig. 1B). No DNA fragment was amplified from the healthy Mexican lime samples used as negative control. The results demonstrated that the BD-RT-PCR method worked properly and clearly showed that the CP of the Idr isolate contains the MCA13 epitope predominantly found in severe strains of CTV. These findings supported the Western blot data and confirmed that the Idr isolate contains the MCA13 epitope sequence associated with severe isolates.

Cloning the CP and RdRp genes. RT-PCR tests were conducted for amplification of the CP and RdRp genes of Idr isolate of CTV. While DNA fragments of about 700 bp and 1700 bp corresponding to the CP and the RdRp genes, respectively, was amplified by RT-PCR from cDNA synthesized from the dsRNA isolated from Mexican lime infected with the Idr isolate of CTV (Fig. 2), no DNA was amplified from the dsRNA isolated from uninoculated Mexican lime (Fig. 2). The results showed that Mexican lime used as maintenance host still contained the Idr isolate of CTV and primers designed based on the CP and RdRp sequences were suitable for amplification of both genes. Therefore, amplification of the CP and RdRp genes enable molecular characterization of the Idr isolate using two different genomic regions.

The CP and RdRp genes were cloned in to pGEM-Teasy plasmid vector using T-A cloning method. A number of recombinant colonies potentially carrying the CP and RdRp genes were obtained in initial blue-white screening. At least 10 colonies for each gene were selected and tested by a colony PCR and the presence of the CP and RdRp gene was confirmed in majority of these colonies. Two colonies carrying the pGEM-T easy plasmid with the CP and RdRp genes designated as pGEM-T Idr CP-1, pGEM-T Idr CP-2, pGEM-T Idr RdRp-1 and pGEM-T Idr RdRp-2 were selected and plasmid was purified and digested with EcoRI. Digestion of pGEM-Teasy plasmids carrying the CP gene revealed an about 750 bp band containing the CP gene and 3000 bp band of pGEM-Teasy plasmid in both clones (Fig.
Molecular Characterization of Igdr Isolate of CTV

On the other hand, the digestion of the pGEM-Teasy plasmids carrying the RdRp gene produced different patterns in two different clones. While digestion of pGEM-T Igdr RdRp-1 clone produced two bands of about 1700 bp and 3000 bp corresponding to RdRp gene and pGEM-Teasy respectively, the other clone produced three different bands. The 3000 bp corresponded pGEM-Teasy vector, the other two fragments (700 and 1000 bp) together corresponded to the RdRp gene suggesting that it has an internal EcoRI site (Fig. 3). The results demonstrated that the CP and RdRp genes were successfully cloned and two different clones for each gene were obtained. Different restriction patterns produced by EcoRI digest of the RdRp clones suggested that these clones contained some sequence variation and the Igdr isolate may contain a mixture of different CTV strains.

Sequence analysis of the CP and RdRp genes. The nucleotide sequence of two different clones of the CP (Igdr CP-1 and CP-2) and the RdRp (Igdr RdRp-1 and RdRp-2) genes were determined and submitted to the GenBank database under the accession KC349866, KC349867, KC349868 and KC349869 respectively.

3. On the other hand, the digestion of the pGEM-Teasy plasmids carrying the RdRp gene produced different patterns in two different clones. While digestion of pGEM-T Igdr RdRp-1 clone produced two bands of about 1700 bp and 3000 bp corresponding to RdRp gene and pGEM-Teasy respectively, the other clone produced three different bands. The 3000 bp corresponded pGEM-Teasy vector, the other two fragments (700 and 1000 bp) together corresponded to the RdRp gene suggesting that it has an internal EcoRI site (Fig. 3). The results demonstrated that the CP and RdRp genes were successfully cloned and two different clones for each gene were obtained. Different restriction patterns produced by EcoRI digest of the RdRp clones suggested that these clones contained some sequence variation and the Igdr isolate may contain a mixture of different CTV strains.

Sequence analysis of the CP and RdRp genes. The nucleotide sequence of two different clones of the CP (Igdr CP-1 and CP-2) and the RdRp (Igdr RdRp-1 and RdRp-2) genes were determined and submitted to the GenBank database under the accession KC349866, KC349867, KC349868 and KC349869 respectively.

Fig. 3. Restriction digestion of the pGEM-Teasy plasmid vector with cloned CP and RdRp genes of Igdr isolate with EcoRI. M: DNA size marker. The sizes of the cloned DNA fragment after EcoRI digestion are indicated by arrows.

Fig. 4. Amino acid sequence alignment of two different clones of the CP (A) and the RdRp (B) genes of Igdr isolate. The positions of amino acids are indicated above the sequence. The identical and similar amino acids are highlighted in yellow and green, respectively. The GenBank accession numbers for the Igdr CP-1, CP-2, RdRp-1 and RdRp-2 are KC349866, KC349867, KC349868 and KC349869, respectively.
and KC349869. The nucleotide and the deduced amino acid sequences of two different clones of the CP gene from the Idr isolate were 100% identical (Fig. 4A, Table 1). However, two different clones of the RdRp gene from the Idr isolate shared only 90% nucleotide sequence identity and only 91% amino acid sequence identity with each other (Fig. 4A, Table 2). The amino acid sequence comparisons of different clones of the RdRp and CP genes of the Idr isolates is shown in Fig. 4. These results showed that while two clones of the CP genes were identical, the RdRp clones were different indicating that two RdRp clones were derived from mixed infection of more than one isolate. To determine the phylogenetic relationships and possible biological activity of the potential isolates in the mixed infection, the deduced amino acid sequences of the CP and RdRp genes from the Idr isolate were compared with respective genes from well-characterized CTV isolates from other Citrus growing regions of the world available in the GenBank databases.

Table 1. Percent amino acid sequence identity of the RdRp gene of Idr isolates with other CTV isolates from different Citrus-growing regions of the world

| Isolates       | IGDIR-2 RdRp | Mexico RdRp | NUAGA RdRp | NZ 18 RdRp | NZ M16 RdRp | NZRB G90 RdRp | QAHA RdRp | T30 RdRp | T318A RdRp | T36 RdRp | T385 RdRp | VT RdRp | NZRB TH30 RdRp | IGDIR-1 RdRp |
|----------------|--------------|-------------|------------|------------|------------|--------------|-----------|----------|------------|----------|----------|---------|----------------|--------------|
| B165 RdRp      | 91           | 86          | 94         | 98         | 95         | 93           | 88        | 94       | 94         | 90       | 94       | 92      | 92             |              |
| IGDIR-2 RdRp   | 89           | 91          | 88         | 90         | 97         | 91           | 91        | 91       | 93         | 91       | 91       | 96      | 90             |              |
| MEXICO RdRp    | 87           | 84          | 86         | 91         | 95         | 87           | 87        | 96       | 87         | 87       | 91       | 86      |                |              |
| NUAGA RdRp     | 92           | 96          | 93         | 89         | 96         | 100          | 90        | 96       | 96         | 99       | 92       | 95      |                |              |
| NZ B18 RdRp    | 92           | 91          | 92         | 86         | 92         | 92           | 88        | 92       | 92         | 92       | 90       | 90      | 90             |              |
| NZ M16 RdRp    | 92           | 88          | 96         | 96         | 90         | 96           | 97        | 93       | 94         | 93       | 94       | 92      | 94             |              |
| NZRB G90 RdRp  | 94           | 94          | 93         | 95         | 94         | 93           | 95        | 94       | 93         | 98       | 92       |         |                |              |
| QAHA RdRp      | 89           | 89          | 98         | 89         | 98         | 89           | 89        | 94       | 88         |          |          |         |                |              |
| T30 RdRp       | 96           | 91          | 100        | 96         | 96         | 96           | 93        |          |            |          |          |         |                |              |
| T318A RdRp     | 91           | 96          | 99         | 93         | 95         | 90           |          |          |            |          |          |         |                |              |
| T36 RdRp       | 91           | 91          | 95         | 90         | 97         | 96           | 93        |          |            |          |          |         |                |              |
| T385 RdRp      | 96           | 93          | 95         | 90         | 97         | 96           | 95        |          |            |          |          |         |                |              |
| VT RdRp        | 92           |             |            |            |            |              |           |          |            |          |          |         |                |              |
| NZRBTH30 RdRp  |              | 92          |            |            |            |              |           |          |            |          |          |         |                |              |

Table 2. Percent sequence identity of the CP gene of Idr isolates with other CTV isolates from different Citrus-growing regions of the world

| Isolates       | IGDIR-IGDIR-1 CP | MEXICO CP | NUAGA CP | NZ B18 CP | NZ M16 CP | NZRB G90 CP | NZRB TH30 CP | QAHA CP | T30 CP | T318A CP | T36 CP | T385 CP | VT CP |
|----------------|------------------|-----------|----------|-----------|-----------|-------------|---------------|---------|--------|----------|--------|---------|-------|
| B165 CP        | 97               | 96        | 96       | 98        | 98        | 98          | 96             | 96      | 96     | 96       | 96     | 96      | 97    |
| IGDIR-1 CP     | 100              | 96        | 96       | 97        | 96        | 98          | 96             | 95      | 96     | 96       | 95     | 96      | 96    |
| IGDIR-2 CP     | 96               | 96        | 97       | 97        | 96        | 98          | 96             | 95      | 96     | 96       | 96     | 96      | 96    |
| MEXICO CP      | 95               | 96        | 95       | 95        | 100       | 96           | 96             | 96      | 96     | 96       | 96     | 96      | 94    |
| NUAGA CP       | 100              | 97        | 97       | 97        | 95        | 96           | 99             | 95      | 96     | 97       | 96     | 97      | 97    |
| NZ B18 CP      | 98               | 98        | 98       | 97        | 96        | 99           | 96             | 96      | 96     | 96       | 96     | 97      | 97    |
| NZ M16 CP      | 96               | 96        | 95       | 97        | 95        | 96           | 97             | 95      | 97     | 96       | 96     | 97      | 96    |
| NZ RB G90 CP   | 96               | 95        | 96       | 97        | 95        | 96           | 94             | 96      | 96     | 96       | 96     | 96      | 96    |
| NZ RB TH30 CP  | 95               | 95        | 96       | 95        | 94        | 95           | 94             | 95      | 96     | 96       | 96     | 96      | 96    |
| QAHA CP        | 96               | 96        | 100      | 96        |          |              |                |          |        |           |        |         |       |
| T30 CP         | 96               | 96        | 96       | 95        | 97        | 96           | 94             |          |        |           |        |         |       |
| T318A CP       | 96               | 96        | 97       | 95        | 95        | 96           | 94             |          |        |           |        |         |       |
| T36 CP         | 96               | 96        | 97       | 96        | 97        | 96           | 95             |          |        |           |        |         |       |
| T385 CP        | 96               | 96        | 97       | 96        | 97        | 95           | 94             |          |        |           |        |         |       |
identity the CP gene of the Igdr isolate was most similar with the CP gene of NZRB-TH30 (Table 1), a resistance breaking amino acid isolate from New Zealand (Harper et al., 2009; Harper et al., 2010). This isolate is able to break the CTV resistance and replicate in P. trifoliat a and also cause severe stem pitting on sweet orange. Interestingly, the CP gene of the Igdr isolate was the least similar to that of two well-characterized mild isolates, T30 from Florida and T385 from Spain with 95% sequence identity (Table 1). The results clearly demonstrated that the Igdr isolates contains a severe component and both clones of the CP gene were derived from this severe component of the mixed infection.

On the other hand, multiple alignment and comparison of deduced amino acid sequences of two different clones of the RdRp from Igdr shared different sequence identity with the RdRp of other CTV isolates. While the RdRp-1 showed 88 to 97% identity, the RdRp-2 shared 86 to 95% amino acid sequence identity with RdRps from other CTV isolates. The RdRp-1 was 95% identical to five isolates with different biological characteristics, but it was most similar to T30 and T385, two well-characterized mild isolates from Florida and Spain, respectively (Table 2). With 86 and 88% identity it was the most different from a QD isolates from Mexico and QAHA from Egypt, respectively (Table 2). In contrast, with 96 and 97% identity the RdRp-2 was most similar to NZRB-G90 and NZRB-TH30 two characterized RB and SP causing isolates from New Zealand (Table 2). It was the least similar to an orange SP isolate NZ-B18 from New Zealand and a QD isolate from Mexico, with 88 and 89% sequence identity, respectively (Table 2). The result revealed that two different clones of RdRp from the Igdr isolates show homology with the RdRp of CTV isolates with very different biological activity one being M and the other being the most severe RB isolates of CTV from different citrus growing regions.

Phylogenetic relationships of the Igdr isolate with previously characterized CTV isolates from different citrus growing regions was determined by comparing the amino acid sequences of the CP and RdRp genes. The phylogenetic analysis of the CP gene revealed four main isolates groups with different biological activity supported by high bootstrap values. The Igdr isolate were clustered with NZRB-TH30, a RB isolate from New Zealand (Fig. 5). The results showed that two different clones of the CP gene with identical amino acid sequences were closely related with a severe stem pitting isolate enable to break CTV resistance in P. trifoliat a confirming that the Igdr isolate contains a severe component. On the other hand, although the a similar phylogenetic tree was constructed using the RdRp genes of different CTV isolates, two different clones of the RdRp
genes of Igldr isolates were clustered separately with different CTV isolates having diverse biological activity. While the RdRp-1 was clustered with T30 and T385, two well-characterized mild isolates from Florida and Spain, respectively, the RdRp-2 was most similar to NZRB-G90 and NZRB-TH30 two well-characterized RB and SP causing isolates from New Zealand (Fig. 4B). These result clearly demonstrated that the Igldr isolate biologically described as a mild isolate actually contains mild and severe strains as mixture.

Discussion

Citrus tristeza virus is reportedly present in Turkey for about fifty years but it did not cause any widespread epidemics and economical losses in citrus mainly grown on CTV-sensitive sour orange rootstocks (Baloglu and Birisik, 2009; Yilmaz, 1999). Epidemiology and destructiveness of CTV depends on the type of scion rootstock combination used for production, the presence of vector species and type(s) of strains found in a specific region. Therefore, besides knowing the alternative rootstocks and keeping efficient vectors out, the characterization of strains is very critical for prevention of CTV epidemics in Turkey. Although CTV isolates found in main citrus growing regions of Turkey were biologically and serologically characterized (Baloglu, 1988; Korkmaz, 2002; Yilmaz and Baloglu, 1998) the molecular characteristics of CTV isolates of Turkey is mainly unknown. Here we report molecular characterization of the first identified CTV isolates, Igldr, using sequences of the CP and the RdRp coding regions of its genome.

Molecular analysis of the Igldr isolate with the MCA13 monoclonal antibody (Pappu et al., 1990) and bi-directional PCR developed based on the MCA13 epitope (Çevik et al., 1996) revealed that it contains MCA13 epitope indicating the presence of severe strains of CTV. The MCA13 epitope was previously characterized and while MCA13 reactive severe strains contain phenylalanine (F) at position 124 of the CP, mild isolates unrecognized by MCA13 have tyrosine (T) at the same position (Pappu et al., 1993) Sequence analysis of the CP of the Igldr isolate showed that it contained F at position 124 of the CP confirming the presence of MCA13 epitope. Since presence of the MCA13 epitope predominantly indicates severe strains such as QD, SP and SY, these analyses clearly showed the presence of severe strains in the Igldr isolate. Although we previously reported the presence of MCA13 reactive strains in Turkey (Korkmaz et al., 2008), finding of this study further confirms and clearly demonstrates the presence of the MCA13 epitope by sequencing of the CP gene, in addition to a positive antibody reaction, in an isolate previously described as mild.

Phylogenetic analysis of amino acid and nucleotide sequences of viral genes has been used for strain identification and molecular characterization. Phylogenetic analyses of CTV isolates were mainly conducted using the CP gene sequences. Comparison of the CP gene of biologically and geographically different CTV isolates revealed that the CP gene sequence correlated with biological activity or geographical origin of isolates (Herrera-Islidrón et al., 2009; Mawassi et al., 1993; Pappu et al., 1993b; Roy et al., 2003). Therefore, it is commonly used for identification of newly discovered strains and genetic relationship among strains of CTV from different citrus growing regions. On the other hand, genome sequences of CTV strains demonstrated that while the 3' half of the genome is highly conserved, significant sequence variation was observed in the 5' half of the genome (Albiach-Marti et al., 2000a; Mawassi et al., 1996; Suastika et al., 2001; Vivees et al., 1999; Yang et al., 1999). Therefore, two genomic regions, the CP and RdRp, representing respectively the 3' and 5' half of the genome were utilized for molecular characterization of Igldr isolate. Comparison of sequences obtained from two randomly selected clones of the CP and the RdRp genes revealed that two cloned CP genes were identical but the RdRp genes were significantly different from each other suggesting a mixed infection with two different strains. Comparison and phylogenetic analysis of the CP and RdRp genes with respective genes of biologically and geographically different CTV isolates available in the GenBank databases revealed that the CP gene and the one of the RdRp gene were most similar to and grouped with severe SP isolates including NZRB-TH30, a RB isolate from New Zealand (Harper et al., 2010) but the other RdRp gene was more closely related with mild isolates and phylogenetically grouped with mild T30 and T385. This finding suggested that the Igldr isolate contains both mild and severe components. Similarly, isolates containing mixed infection of mild and severe strains in the same tree was previously reported in different citrus growing regions (Harper et al., 2009; Sentandreu et al., 2006). Particularly one Spanish isolate contain both mild and severe strains which were separated by aphid transmission and host passage and characterized in detail (Ayllon et al., 2006; Sentandreu et al., 2006). Similarly, some CTV isolates from New Zealand were sequenced and genomes of different strains were obtained from the same isolates (Harper et al., 2009). These and other findings clearly showed that mixed infections are common and the component of mixed infection can be separated by aphid vectors experimentally as well as naturally in the field.

Therefore, from epidemiological and control point of view it is important to know the biological, serological and molecular characteristics of CTV isolates in the region. Biological characterization of the Igldr isolate was previ-
ously completed and based on the reaction in indicator plants it was considered a mild isolate (Baloglu, 1988; Yılmaz and Baloglu, 1998; Korkmaz, 2002). The finding in this study showed that the İğdır isolate actually contains mixed infections of mild and severe SP isolates. Since the biological indexing of the İğdır isolate was done by graft inoculation all components of the mixed infection graft inoculated and strains in the isolate were not separated. Therefore, the mild nature of the İğdır isolate in the indicator plant can be explained by interaction of different strains and/or cross protection of severe component by the mild strains found together. Determination of this has to await separation of mixed strains by aphid transmission.

Citrus tristeza virus is present in all in main Citrus-growing regions of the Mediterranean coast for many years (Baloglu, 1988; Yılmaz and Baloglu, 1998; Korkmaz, 2002) and recently reported in other Citrus-growing region of Turkey (Korkmaz et al., 2008). Studies indicated that virus is not spreading constantly from tree to tree but rather found sporadically (Baloglu, 1988; Yılmaz and Baloglu, 1998; Korkmaz, 2002; Korkmaz et al., 2008). This is probably due to the absence of the efficient vector T. citricida. Current predominant vector is A. gossypii, which unable to transmit CTV efficiently (Roistacher and Bar-Joseph, 1984; 1987). However, possible introduction of T. citricida may change the situation. It has been well documented in Florida that the predominant strain profile was dramatically changed after introduction of T. citricida and severe SP isolates become more prevalent (Halbert et al., 2004).

This study provides serological and molecular data as well as sequence information for the presence of severe strain component in the previously identified Turkish CTV isolate İğdır, which is known as a mild isolate. Therefore, it is certain that the severe strains of CTV are actually present in citrus producing areas of Turkey. These strains are presumably transmitted but are not separated from mild strains by transmission by A. gossypii in nature of the absence of the most efficient vector, T. citricida. A possible introduction of T. citricida, which is currently found in Spain and Portugal, into Turkey may cause efficient transmission of severe components in so-called mild isolates and may change the predominant strain profile. Newly transmitted predominant severe strains could result in epidemics in citrus growing regions of Turkey where the majority of citrus is grafted on CTV-sensitive sour orange rootstocks. Therefore, the CTV isolates and/or strains found in Turkey should be detected, identified characterized in genetic diversity and epidemiological point of view and CTV resistant rootstocks and scions should be developed in preparation for the inevitable results of T. citricida introduction into Turkey.

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