Molecular Studies on Tomato Yellow Leaf Curl Virus in Infected Tomato and Pepper Plants

Younis, H.A¹, N. A. Zaid² and S. Deraz²

¹Department of Agricultural Botany, Faculty of Agriculture (Saba-Bacha), Alexandria University, University 21531, Egypt
²City for Scientific Research and Technology Applications, Burg El Arab, post code 43912 Egypt. Corresponding author: Z. Nehal, e-mail: nahlaz2015@gmail.com

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ABSTRACT: Tomato Yellow Leaf Curl (TYLC) is an economic disease that caused dramatic losses in tomato crops. It is caused by a monopartite whitefly-transmitted begomovirus individually or associated with Betasatellite DNAs. Recently, TYLC disease became more severe in Egypt, and it destroyed completely the tomato crop in some crop seasons during the year. Tomato and pepper samples with curling symptoms were brought from El-Nubaria region, El Beheira governorate in Egypt and tested by Polymerase Chain Reaction (PCR) with specific primers to detect the presence of the causal begomovirus and its associated DNA betasatellites and/or the presence of new genotypes between two tomato yellow leaf curl virus (TYLCV) isolates. Disease symptoms varied between yellowing, curling and cup-shaped symptoms. PCR products indicated the presence of the two Mediterranean TYLC begomoviruses strains TYLCV-IL and TYLCV-MiLD, while no new genotypes of the virus or TYLCSV isolate were found in these samples. Additionally, betasatellites were found to be associated with the virus in all infected tomato and pepper samples either the betasatellite complete genome or its defective form. In conclusion, no TYLCV new genotypes were detected in the tested samples to be responsible for the severe tomato losses in El-Nubaria region in El Beheira governorate, but TYLCV/Betasatellites association is expected to play an axial circumference in TYLC disease severity in the same region.

Keywords: Tomato Yellow leaf curl virus, Betasatellite DNA, PCR, new genotypes, El- Nubaria region.

INTRODUCTION: Family Solanaceae members are some of the most important economic crops in Egypt for both local consumption and exportation. During the last years, dramatic losses occurred for tomato plants in different localities in Egypt such as El Fayium governorate, EL-Nubaria and Wadi El Natron because of a severe infection with the tomato leaf curl virus (TYLCV) (Aboul-Ata et al., 2000; El-Dougoud et al., 2014; Rabie et al., 2017 and Sofy et al., 2017). The tomato Yellow Leaf Curl Virus (TYLCV) is a monopartite begomovirus that belongs to the Geminiviridae family, which represents the circular, single-stranded DNA plant viruses (Lazarowitz, 1992) and involves more than 380 whiteflies (Bemisia tabaci)-transmitted species other than TYLCV (Zhao et al., 2019).

Whiteflies have piercing and sucking mouthparts utilized for feeding in plant phloem (Byrne and Bellows, 1991; Gill, 1990). It is known for obligating harboring besides the primary endosymbionts with their hosts (Campbell, 1993; Thao and Baumann, 2004), and synthesizing amino acids that are in inadequate amounts in phloem sap (Buchner, 1965; Morin et al., 1999, Rosell et al., 2009 and Gorovits and Czosnek 2013). This type of viruses circulates in the whitefly body and behaves away from the midgut (MG) lumen till the hemolymph and, eventually, till the primary salivary glands (PSGs), where it is given back into the plant host pending insect feeding (Czosnek et al., 2002 and Matore, 2018). The minimum effective periods of either acquisition-access or inoculation-access are almost 10 to 20 min per each. While the latent period is about 8h at minimum starting from the acquisition is required for the insect to be able to infect a new healthy plant. Begomoviruses are thought to be not capable of replicating in their insect vectors (Rosen et al., 2015, Gilbertson et al., 2015). However, strong evidence demonstrated by He, et al., (2020) gave an exception to TYLCV and proved its ability to replicate in its vector (Bemisia tabaci) mainly in the salivary glands (Navas-Castillo et al., 2011). This monopartite genome (2.8 kb) consists of six proteins responsible for replication, transmission, pathogenesis, and movement functions (Jeske, 2009). TYLCV spreads in tropical and subtropical regions (Navas-Castillo et al., 2011) and has a wide host range as it infects numerous plant species causing considerable yield losses in many important crops either monocots or dicots; the mechanism of geminiviruses inside the plant is to reshape the intracellular environment of the host so
that create favorable conditions for replication and propagation of the virus (Mansoor et al., 2003; Mansoor et al., 2006 and Maio, 2020); The viral symptoms include plant stunting, erect shoots, small and misshaped leaflets, chlorotic, curling and cup-shaped leaves (Cohen and Nitzany, 1960). There are many TYLCV isolates globally which classified based on their origin (Cohen and Lapidot, 2007). In the Mediterranean East in particular, there are two main TYLCV isolates: TYLCV-IL from Israel and TYLCV-MID from the Mediterranean basin, besides Tomato Yellow leaf curl Sardinia virus (TYLCSV) from Italy (Kheyripour et al. 1991). Betasatellites are circular single-stranded DNAs ranged between 660-1350 nucleotides in the Nile Basin area (Briddon et al., 2003 and Idris, et al., 2002). Although the existence of betasatellites was reported in some Mediterranean countries including Egypt (Conflon et al., 2018), betasatellite/begomovirus association needs more survey and study to estimate the risk of betasatellites on the disease severity and incidence. Experimentally, TYLCV was found to be the helper of most of the various DNA betasatellites with which it is co-inoculated (Ito et al., 2009; Kon et al., 2009; Ueda et al., 2012 and Zhang et al., 2012), and the most noticeable matter, all of them enhanced its virulence.

Tomato and pepper have been reported as a symptomatic TYLCV host (Moury and Verdin 2012); (El-Dougoud et al., 2014) detected TYLCV in tomato and pepper plants from different areas in Egypt using polymerase chain reaction (PCR) and degenerate primers, the targeted sequence (~530 bp) in that reaction was of the coat protein gene (CP) of Begomoviruses. The resulted DNA sequence of this investigation showed high nucleotide identities (up to 99%) to isolates of TYLCV in the GenBank; Moreover, Rabie et al., (2017) sampled TYLCV symptomatic tomato crops from different regions in Egypt during the year 2014, Israel and Mild TYLCV strains were detected in these samples by multiplex and real-time PCR. The complete sequence genome of the Egyptian isolate obtained in this study showed a high degree of identity to other reported Egyptian isolates in addition to TYLCV-IL Jordan isolate and TYLCV-IL Japan isolates.

Because of begomoviruses’ high capability to recombine, Fauquet et al, 2005, mentioned that new recombinants of begomoviruses were expected to be emerged with higher virulence and altered host range. For instance, Tomato yellow leaf curl Axarquia virus (TYLCAxV) recombiant was detected from the highly recombinogenic isolates TYLCV and TYLCSV in Spain (Garcia-Andres et al., 2006). Also, Belabess et al., 2015, reported a new recombinant named TYLCV IS76 which has been discovered in 2010 from both isolates in southern Morocco; the emergence of this recombiant coincided with the increasing use of tolerant cultivars in the 2000s. However, although so many Mediterranean countries are using tolerant tomato cultivars, TYLCV-IS76 was not detected outside Morocco (Belabess et al. 2018).

Recently, some begomoviruses in Egypt like the Squash leaf curl virus (SLCEGV) were found to be joined with a DNA molecule defined as betasatellite; for example, Abdel-Salam et al., 2017; examined betasatellites/ SLCEGV association in different crops in Egypt around 4 years by various molecular techniques. He finally concluded that the genomic mixture of SLCEGV and the possible acquisition of betasatellites by SLCEGV modified the viral virulence and fitness. Briddon et al., 2003 and Sivalingam, and Varma, 2012 proved that betasatellites had an effective role in begomoviruses accumulation and severe symptoms appearance and Conflon et al., 2018 supported the same idea by their study which illustrated that all resistant tomato plants co-infected with Cotton leaf curl Gezira betasatellites and TYLCV IL/MILD showed leaf curling and mosaic symptoms, while the same resistant cultivar inoculated only with TYLCV was asymptomatic.

The main objectives of this study is testing the introduction of TYLCV new recombinants to Egypt after the epidemic appearance of TYLCV in some tomato production areas like El-Nubaria in El Beheira Governorate and test the role of betasatellites in exhibiting symptoms in tomato and pepper plants.

MATERIALS AND METHODS

TYLCV- infected tomato sampling (first season, 2019)

Tomato symptomatic samples were collected from EL-Nubaria region, El Beheira governorate during the late summer of the year 2019. All collected samples were classified and labeled as follows: Infected tomato with clear yellowing and curling symptoms (Ts), infected tomato with unclear yellowing and curling symptoms (Tas), and infected pepper with curling symptoms (P'). These samples were tested to detect the presence of different genotypes of TYLCV including TYLCSV, Betasatellites and the native isolates TYLCV- IL and TYLCV- MiLD that are previously detected in the Mediterranean East. Samples freeze at −20 for DNA extraction and molecular analysis.

DNA extraction
DNA was extracted from the tested leaf samples by the CTAB method according to Gabriadze et al., (2014) and Belabess et al., (2015) with some modifications in the speed of the centrifugation step. A sample of 20 mg of each plant leaf was ground and mixed in a sterile tube with 500 µl CTAB buffer. Tubes were incubated for 30 min in 65°C water bath and inverted three or four times during which; samples were centrifuged for 15 min at 2900 rpm then each supernatant was transferred to a new tube with 450 µl chloroform, the mixture was stirred at room temperature for 5 min followed by centrifugation for 10 min at 2900 rpm. 350 µl of the aqueous phase was mixed with 350 µl of isopropanol and the mixture was centrifuged at 2900 rpm for 15 min. The pellet was washed with 70% ethanol and after a complete drying from ethanol, the samples were re-dissolved in 50 µl of TE buffer and stored at -20 for further analysis.

Polymerase chain reaction (PCR) reactions
All the PCR reactions were performed by Cosmo PCR master mix W1020300X, Willowfort Co., following the manufacturer's instructions. PCR conditions were set up for each program individually based on primers used.

Detection of A component genome (DNA-A) of begomoviruses
DNA-A of begomoviruses was tested by specific primers according to Abd El Salam et al. (2017) which were designed to amplify the fragment of the DNA-A coat protein (580 bp). The DNA primers were AV Core 5'GCCHATRTAYAGRAAGCCMAGRAT3' and AC Core5'GGRTTDGARGCATGHGTACANGCC3'.

Detection of Betasatellites
Betasatellites complete genome and their defective genome were detected by specific primers according to Abd El Salam et al., (2017). The expected products were 1300 bp and 700-1000 bp, respectively. The DNA primers were β01 (F) 5'-GGTACCTACCTACGCAGCGACC3' and β02 (R) 5'-GGTACCTACCCCTCAGGGGTACAC3' PCR for betasatellites and DNA-A of begomovirus was conducted under the same conditions of 95 °C for 5 min as denaturation step followed by 30 cycles of 95 °C for 1 min, primer annealing step at 58 °C for 1 min, extension step at 72 °C for 1 min and final extension at 72 °C for 7 min. Electrophoresis gel was performed in 1% agarose and stained with Ethidium bromide.

TYLCV/TYLCSV recombinants (PCR1)
The degenerate primers mixture was used in this reaction to detect the presence of TYLCSV isolate and/or its recombinants TYLCV-IL/TYLCSV, TYLCV-MID/TYLCSV at locus OR (Origin of Replication). The expected products of this reaction were 405 bp, 187 bp, and 548 bp, respectively.

PCR1 primer mixture (Belabess et al., 2015): The DNA primers were Sar43R 5' - TCGTGACGCCYACTWCTTTTATCGG-3' / Mild2277F 5'-CTSWCCCCARTGABGGTG-3'/ IL2629F 5'-GTTGTCCCTCAAAGCTCTAWG-3'/ TYS2416 F1* 5'-CCCTCAGACTGAATGAGCATG-3' and TYS2416 F2* 5'-CCYTCAAYTGSATGAGAAYA-3'.

TYLCSV/TYLCSV recombinants (PCR2)
The second specific primers mixture was used in this reaction to detect the presence of TYLCV isolates IL and Mild and/or its recombinants TYLCSV/TYLCSV. The expected products of this reaction were 230 bp, 592bp and 440 bp, respectively.

PCR2 primer mixture (Belabess et al., 2015): TY78R5' GCAATTGGATCTTTAAGTGTAGSACR-3'/IL2629F5'GGTGTCCCTCAAAGCTCTAWG-3'/Mild2277F 5'CTSWCCCCARTGABGGTG-3'/TYS2416F1*5'-CCCTCAGACTGAATGAGCATG-3'/TYS2416F2*5'CCYTCAAYTGSATGAGAAYA-3' PCR1 and PCR2 were performed under the same conditions of denaturation step at 95 °C for 5 min, followed by 30 cycles of 95 °C for 45 sec, 60 °C for 45 sec, 72 °C for 45 sec, 72 °C for 10 min. Electrophoresis gel was performed in 1.5% agarose and stained with Ethidium bromide.

RESULTS AND DISCUSSION
Detection of (A) Component of Begomoviruses and Betasatellites in Tomato from El-Nobaria region

Tomato and pepper samples were collected at the end of the season at the maturity stage of their life cycle. Tomato samples from EL-Nobaria region were classified based on symptoms appearance as shown in Figure 1 (A, B and C). (A) shows tomato plants with curling and yellowing symptoms; (B) shows tomato plants with only yellowing symptoms, while severe curling symptoms on pepper plant samples illustrated in C.

Regardless the etiology of this differentiation in symptoms degrees, it may be attributed to plant stage at the time of infection. Brown (2000) reported that TYLCV infection of tomato plants in the early stage causes severe symptoms. Short-
developed leaves after infection are cupped downward; while later-developed leaves are prominently chlorotic and deformed in addition to upward rolled-leaf margins and curling between the veins.

(A): Clear yellowing, curling and cup-shaped symptoms on tomato leaves (Ts).

(B): Unclear yellowing and curling symptoms on tomato leaves (Tas).
Detection of Begomoviruses and Betasatellites in Tomato and Pepper

The results showed that both clear and unclear symptomatic tomato plants have a DNA A fragment of begomovirus (~580bp) clearly appeared in the two frequencies of each (Figure 2). Similarly, the betasatellites complete genome (~1300bp) or its defective form (700-900bp) was observed at the same samples. However, the complete genome appeared in a high concentration in the (Ts) tomato samples with its defective form, while it was appeared in a lower concentration in the (Tas) tomato samples with a more concentrated band for its defective form (Figure 2). PCR test for pepper proved the presence of A component genome (DNA A) of begomovirus in the two pepper sample frequencies coincided with the presence of Betasatellite DNAs and its defective form in the same frequencies but in low concentration (Figure 3). This begomovirus/betasatellite association was reported previously by (Dry et al., 1997; Briddon et al., 2003; Idris et al., 2005; Idris et al., 2011 and Fiallo-Olivé et al., 2012, Abdel Salam et al., (2017) found that betasatellite DNA association with Squash leaf curl bipartite begomovirus led to increasing the biodiversity of the virus through modifying its virulence and fitness.

TYLCV degenerate primers used in this experiment indicated that the two Mediterranean TYLCV strains IL and MiLD were detected in tomato, while only IL strain was found in pepper (Figure 4). Noteworthy, Conflon et al., (2018) mentioned that TYLCV accumulation increased in the mixed infection of betasatellite with TYLCV-Il, while those of TYLCV-Mld/betasatellite infection decrease this accumulation in plants.

Usually, severe or moderate TYLCV infection is determined by two interfering factors: Efficiency of Ty1 gene-bearing cultivars, which play an important role in TYLCD severity with a variable immune mechanism such as encoding RNA-dependent RNA polymerase or viral transcriptional gene silencing by allelic genes Ty-1 and Ty-3 (Verlaan et al., 2013; Caro et al., 2015); or TYLCV resistance locus Ty-2, which was genetically linked to the gene TYNBS1 that encodes an NB-LRR (Nucleotide-Binding domain and Leucine-Rich Repeat) protein a type of plant immune receptor (Yamaguchi et al., 2018). Nevertheless, many recent suggestions refer that these resistance cultivars no longer being effective against TYLCV because of new present or coming breaking strains of the virus under field conditions (Verlaan et al., 2013; Butterbach et al., 2014; Belabess et al., 2016; Ohnishi et al., 2016). The second factor is betasatellites, which play an axial circumference in TYLCV severity. Generally, betasatellites are considered pathogenicity determinants (Voorburg et al., 2020) because of their ability to produce the single protein BC1 which takes part in virus pathogenicity Saeed, (2010). This BC1 protein was proved to increase symptoms expression (Cui et al., 2004; Saunders et al., 2004), works as post transcriptional gene silencing (PTGS) suppressor (Amin et al., 2011;
Cui et al., 2005) and in the systemic movement of the virus in plants (Briddon and Stanley 2006). Voorburg et al. (2020) showed in a study performed on ty-1 transformants of Nicotiana benthamiana and tomato plants that TYLCV accumulation was increased when plants co-infected with the virus and betasatellites compared to plants only infected with TYLCV; also, Conflon et al., (2018) reported that betasatellites overcame Ty-1 resistance gene in TYLCV-resistant tomato cultivars which are commonly used in the Mediterranean countries.

Fig. 2: Detection of A component (580 bp) of begomoviruses and β DNAs CG (Complete genome ~1300 bp) and β DNAs DF (Defective form 700-1000 bp) in symptomatic and asymptomatic tomato.

Fig. 3: Detection of A component of begomoviruses and betasatellites in symptomatic pepper.
Fig. 4: Detection of TYLCV IL (230 bp)/ TYLCV MiLD (592 bp) strains with degenerate primers in tomato and pepper.

Detection of Parental Isolates and new genotypes of TYLCV in Tomato and Pepper:
The two multiplex PCR tests performed in this study were used to test the presence of new recombinants of TYLCV in El-Nubaria region. Often, having a new strain or recombinant from a viral genome happens due to genetic selection through mutation or recombination actions (Fleischmann, 1996). Lefeuvre et al., (2010) found that the epidemiology of TYLCV variants in the western Mediterranean and the Middle East resulted from recombination. The results of our study illustrate that there is no new genotypes were found in tomato and pepper samples either from TYLCSV/TYLCV or TYLCV/TYLCSV recombination (Figure 5);

Fig. 5: Detection of TYLCV genotypes, no new genotypes or TYLCSV isolate were found in tomato and pepper samples, while only the native isolates of TYLCV (IL and MiLD) were detected.

Additionally, it illustrates the absence of the parental isolate TYLCSV. So, there were no new genotypes have been introduced to this area from any new imported seeds or seedlings. Likewise,
what happened before in different countries; for instance, Mabvakur et al., (2016) indicated that the epidemic distribution of TYLCV in Australia and China originated by the introduction of the virus from Eastern-Asia around Japan and Korea. Moreover, the author found that the New Caledonian epidemic was imported by a cultivar from the Mediterranean West region and the Mauritian epidemic was a result of an imported cultivar from the neighboring island of Reunion. By contrast, IS76 recombinant that has been detected in Morocco by Belabess et al., (2015) was not detected in any other neighboring countries outside Morocco (Belabess et al., 2018).

CONCLUSION

There were no imported genotypes to El Nubaria region in El Beheira governate during the last years; but both IL (~230 pb) and Mid (~592) strains of TYLCV appeared in tomato and pepper plants associated with DNA betasatellites that has been reported to strongly affect the virus severity on plants.

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Conflicts of interest/Competing interests

The authors of this study declare that they all accept the publication of these data and they have no conflict with each other or any other external scientist could prevent the publishing this work.

Availability of data and material

The data used to support the findings of this study are included within the article.

Code availability

The coding of the data is available from the corresponding author upon reasonable request.

Authors’ contributions

The first author is the owner of the main idea of the study. The second author provided some materials and implemented the practical work of the idea. The third author provided the lab supplies and facilities.

Ethics approval

The study have been conducted with no opposition with the scientific ethics.

Consent to participate

All authors accept the publication of these data.

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الملخص العربي
دارسات جزيئية على فيروس تجد الإوراق الأصفر في الطماطم على نباتي الطماطم والبصل
حسن علي يونس1 و نهال عبدالمجيد زيد2 و سحر دارز2
1- قسم النبات الزراعي، كلية الزراعة – ساها باشا، جامعة الأسكندرية، مصر
2- مدينه الابحاث العلمية والتطبيقات التكنولوجية برج العرب، مصر

يعتبر تجد الإوراق الأصفر مرض اقتصادي بسبب خسائر فادحة لمحصول الطماطم. يتسبب هذا المرض عن بيجوموفيروس أحادي الجزيء، وينقل للنبات عن طريق الذرية البيضاء متفردا أو مصوما بالحمض النووي لجزيء البينستالايت. وقد أصبح مرض تجد الإوراق الأصفر طريدا أكثر شراسة في مصر، وقد دمر محصول الطماطم كليا في بعض المواسم الزراعية خلال السنة. وتم الحصول على عينات من نباتي الطماطم والبصل المصابة بأعراض تجد الإوراق الأصفر من خلال البوادر المنحتفة (PCR) ومنطقة النيبارية في مصر، وتم اختبارها بواسطة التفاعل المتماسك للذبابة النباتية، وبوادي واسعة الدائرة للكشف عن وجود البيجوموفيروس المسبب، أو البينستالايت المرتبطة به أو أي مؤلفات جديدة لهذا المرض، والتي قد تكون استقامت إلى TYLCV أو السلالة الأوروبية TLYCSV وTYLCV-CL أو السلالة الأفريقية TYLCSV-IL و TYLCV-MiLD، وعولاوة على ذلك، فقد وجد أن البيجوموفيروس يمتلك الطماطم فلسف وميزة TYLCV-CL و TYLCV-MiLD، بينما لم توجد أي مؤلفات جديدة من الفيروس أو السلالة الأوروبية له البينستالايت مرتبطة بالفيروس في كل عينات الطماطم والبصل المصابة في صورة الجينوم الكامل للبينستالايت أو الجينوم المنقوص. ومع ذلك، لم يتم اكتشاف أي مؤلفات جديدة من الفيروس قد يكون لها تأثير على الخسائر الشديدة للطماطم في منطقة النيبارية، ولكن من المتوقع أن أرضي فيروس تجد الإوراق الأصفر مع البينستالايت قد يلعب دوراً محوريا في الشدة المرضية لمرض تجد الإوراق الأصفر في الطماطم في نفس المنطقة.