An Integrated Diagnostic Approach for Citrus Exocortis Viroid

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Abstract

Background

Citrus exocortis viroid (CEVd) is a circular single-stranded RNA pathogen consists of around 370 nucleotides and leads to a severe disease showing bark scaling symptom on citrus crops, which leads to yield decrease and economic loss. Since the absence of viroid-encoded proteins, methods for CEVd detection mainly counts on bioassays or nucleic acid-base approaches. In order to validate the CEVd disease, here we developed an integrated diagnostic protocol.

Methods

CEVd transcripts were inoculated onto two susceptible cultivars of Solanum lycopersicum L., cv. Rutgers and cv. Double-Fortune, seedings. After inoculation, total RNAs of the two tomato cultivars were extracted to detect CEVd infection by dot blot hybridization, one-step reverse transcription PCR (one-step RT-PCR) and real-time reverse transcription PCR (real-time RT-PCR). In addition, the symptom development of both cultivars was recorded weekly.

Results

The tomato cultivar Rutgers rather than Double-Fortune or others was selected as a suitable CEVd-indicator plant and the bio-index score was established based on epinasty, vein necrosis, leaf size reduction and stunting symptoms. In addition, the isolate of CEVd that collected from citrus eld could rapidly and consistently cause the index symptoms on Rutgers. As expected, CEVd could be specifically and sensitively detected in both tomato and citrus plants by dot-blot hybridization and RT-PCR technologies, including one-step RT-PCR and real-time RT-PCR. Furthermore, we found that the levels of CEVd genomic RNA or CEVd derived small RNAs are correlated to symptom severity. Conclusions

In this study, we developed an integrated detection method for CEVd and revealed potential underlying viroid-host interactions.

1. Background

As simple as solely bearing a single-stranded, circular, noncoding RNA with a range of 246-401 nucleotides in size, viroids are known the smallest plant pathogens. Given a tiny genome, viroids are difficult to be detected and thereof often neglected. So far, little is known about the importance and pathogenesis of viroids although many economic crops are reportedly susceptible for viroid infection (Daròs, Elena, and Flores, 2006; Ding, 2009; Ding and Itaya, 2007; Ding and Wang, 2009; Flores et al., 2005; Tsagris et al., 2008). Without any coding proteins to provide specific functions and not being protected in a coat protein or an envelope, viroids have gained much interest in recent years to elucidate how a non-coding RNA interacts with host factor(s) to accomplish its replication and pathogenicity, such
as survival from the cellular RNA degradation machineries, replication in specific subcellular compartments, and trafficking in a plant.

Among Pospiviroidae, seven viroids, including *Citrus exocortis viroid* (CEVd, genus *Pospiviroid*), *Hop stunt viroid* (HSVd, genus *Hostuviroid*), *Citrus bent leaf viroid* (CBLVd, genus *Apscaviroid*), *Citrus dwarfing viroid* (CDVd, genus *Apscaviroid*) (former *Citrus viroid III*, CVd-III), *Citrus bark cracking viroid* (CBCVd, genus *Cocadviroid*) (former *Citrus viroid IV*, CVd-IV), *Citrus viroid V* (CVd-V, genus *Apscaviroid*) and *Citrus viroid VI* (CVd-VI, genus *Apscaviroid*) (initially named *Citrus viroid original sample*, CVd-OS (Ito et al., 2001)), are reportedly able to infect citrus (Adams et al., 2016). Among these, *Citrus exocortis viroid* (CEVd), the causal agent of the exocortis disease of citrus plants (Semancik and Weathers, 1972), leads to a bark shelling or scaling disorder that affects trifoliate orange (*Poncirus trifoliata* (L.) Raf.), Troyer citrange, Carrizo citrange (*Citrus sinensis* (L.) Osbeck×*Poncirus trifoliata* (L.) Raf.) and Rangpur lime (*Citrus limonia* Osb.), which are all widely used as root-stocks in commercial orchards (Vernière et al., 2006). Not only citrus species, but also many different plants are reported in field grown herbaceous hosts, including tomato (*Solanum lycopersicum* L.) (Fagoaga and DuranVila, 1996; Mishra et al., 1991; Verhoeven et al., 2004), eggplant (*S. melongena* L. cv. Sonja) (Fadda et al., 2003; Fagoaga and DuranVila, 1996; Fagoaga, Pina, and Duran-Vila, 1994), carrot (*Daucus carota* L.), turnip (*Brassica napus* L.) (Fagoaga and DuranVila, 1996), broad bean (*Vicia faba* L.) (Fagoaga, Semancik, and Duran-Vila, 1995), and grape plants (García-Arenal, Pallás, and Flores, 1987).

Due to the absence of viroid-encoded proteins, methods for viroid detection must rely on bioassays or nucleic acid-based approaches rather than protein-based serological ones. Diagnosis of citrus exocortis disease was initially performed by an indicator plants, Etrog citron (*Citrus medica* L.) (Calavan, 1968). Molecular detection for CEVd was then achieved by several hybridization-related methods with/without gel electrophoresis (Fonseca, Marcellino, and Gander, 1996; Guo-Ping and Shikata, 1995; Murcia et al., 2009; Nakahara, Hataya, and Uyeda, 1999). Furthermore, with higher sensitivity and efficiency, reverse transcriptase-polymerase chain reaction (RT-PCR) (Bernad and Duran-Vila, 2006; Nakahara et al., 1999; Ragozzino, Faggioli, and Barba, 2004; Yang, Hadidi, and GARNSEY, 1992), multiplex RT-PCR (Ito, Ieki, and Ozaki, 2002; Wang et al., 2009) and real-time RT-PCR (Papayiannis, 2014; Saponari et al., 2012; Tessitori et al., 2004) were also developed for CEVd detection. However, due to the high sensitivity, PCR-based methods can show false positives resulted from contamination, while false negatives also can arise from the cDNA products of unsuitable size generated from reverse transcription (Murcia et al., 2009) or the failures of primer annealing caused by the nucleotide mutations within primer-binding sites of viroids (Lin, unpublished). Additionally, for inspection, quarantine and control of CEVd, a reliable, accurate, and rapid diagnostic procedure is needed. Therefore, it is desirable to employ more than one kind of diagnostic methods and/or at least two different pairs of primers to accurately identify or detect a specific viroid. Here, we developed an integrated diagnostic protocol for CEVd, comprised of defined bio-index of indicator plants, dot blot hybridization, one-step RT-PCR, and real-time RT-PCR. Moreover, a tomato model for studying CEVd pathogenesis was established as well.
2. Methods

2.1 Plant material, growth conditions and viroid sources

Tomato cultivars *Solanum lycopersicum* L. cv. Rutgers select (from The World Vegetable Center, Tainan, Taiwan) and *S. lycopersicum* L. cv. Double-Fortune (Known-You Seed Co., Kaohsiung, Taiwan) plants were grown in pots with soil at 28°C in a 16-h light / 8-h darkness cycle for four to six weeks.

The four viroid infectious clones, pBd-ASSVd (encodes for *Apple skin scar viroid*), pBd-CCCVd (encodes for *Coconut cadang cadang viroid*), pBd-CEVd (encodes for *Citrus exocortis viroid*) and pCM20d (encodes for *Chrysanthemum chlorotic mottle viroid*; CChMVd), were constructed by head-to-tail of viral genome and cloned into pBluescript SK (+) and kind gifts from Dr. Ricardo Flores, Instituto de Biología Molecular y Celular de Plantas, UPV-CSIC as described previously (*Daros and Flores, 2004*) pGA482-PSTVd contains monomeric *Potato spindle tuber viroid* (PSTVd) in a pGA482 binary vector as described previously (*Itaya et al., 2001*), which is a kind gift from Dr. Biao Ding (Ohio State University, Columbus, OH, USA).

2.2 In vitro transcription and viroid inoculation

pBd-CEVd was firstly linearized by *Xho* I digestion prior to being used as a template for *in vitro* transcription with T7 RNA polymerase in the MAXiSript kit (Ambion, TX, USA) according to manufacturer’s instructions. Approximately 100 ng of viroid transcripts were mechanically introduced onto cotyledon of 12 day-old tomato seedlings. The negative control of plants was mock-inoculated with 20 mM sodium phosphate buffer (pH 7.0) as described previously (*Cress, Kiefer, and Owens, 1983; Hu et al., 1997; Salazar et al., 1988; Zhu et al., 2002*).

2.3 Total RNA extraction

100 mg of Rutgers or Double-Fortune leaves tissue were firstly ground with liquid nitrogen, and 1 ml of Trizol® (Invitrogen, CA, USA) was added. The homogeneous lysate was transferred to a 1.5-ml centrifuge tube and briefly vortexed. After incubation at 25°C for 5 min, the samples were centrifuged at 13,200 rpm, 4°C for 10 min. The supernatant was transferred to a new 1.5-ml centrifuge tube and mixed with 0.2 ml of chloroform. Each tube was vortexed for 15 sec and incubated at 25°C for 3 min. After centrifugation with 13,200 rpm at 4°C for 10 min, the supernatant was transferred to a new 1.5-ml centrifuge tube and mixed well with 0.5 ml of 2-propanol prior to further incubation at 25°C for 10 min. Total RNAs were spun down with 13,200 rpm at 4°C for 15 min. Then, the supernatant was removed and RNA pellets were dissolved in DEPC-treated water and stored at -80°C.

2.4 Preparation of examination of sensitivity limitation by in vitro transcripts

To generate monomeric CEVd construct, we conducted the polymerase chain reaction (PCR) with PSTVd-*F-Bam*HI-88 / CEVd-*R-Bam*HI-88 primer sets (see Table 1 for details) using the pBd-CEVd as templates. Then, the amplified PCR fragment of CEVd was cloned into pBluescript II SK (-) to obtain pBmCEVd.
pBmCEVd was linearized by *Xho*I digestion and used for *in vitro* transcription with the mMMESSAGE mMACHINE® *in vitro* transcription kit (Ambion, TX, USA) according to manufacturer's instructions. CEVd transcripts were then purified by the MEGAclean® Kit (Ambion) according to manufacturer's instructions. To measure the concentration of purified CEVd transcripts, a NanoDrop spectrophotometer (NanoDrop Technology, Rockland, DE) was employed. The molecular weight of CEVd transcripts was calculated by a tool of website, Oligonucleotide Properties Calculator (http://www.basic.northwestern.edu/biotools/oligocalc.html). Based on the quotient of concentration by the molecular weight of CEVd transcripts, the copy number was calculated.

2.5 Dot blot hybridization

For dot blot hybridization, the digoxigenin (DIG)-labeled full-length viroid cDNA probes of ASSVd, CCCVd, CEVd, and CM20d were prepared by PCR amplification using pBd-ASSVd, pBd-CCCVd, pBd-CEVd, pCM20d, and pGA482-PSTVd as templates, respectively. Total RNAs were spotted on nylon membranes (Schleicher & Schuell Bioscience, NH) and then cross-linked by UV Stratalinker® 1800 (Stratagene, CA). The membranes were pre-hybridized with DIG Easy Hyb Granules (DIG Northern Start Kit, Roche) at 65°C for 1 hr. DIG-labeled probes were denatured at 100°C for 5 min and were rapidly cooled on ice. Denatured probes were then immediately added and incubated at 65°C for overnight. Membranes were washed twice with 2× SSC, 0.1% SDS at RT for 5 min and additional twice with 0.1× SSC, 0.1% SDS buffer at 65°C for 15 min. Afterward, membranes were rinsed with Maleate acid buffer (0.1 M Maleic acid, 0.15 M NaCl, pH 7.5) for 5 min. Blocking was carried out by incubation with Blocking solution (DIG Northern Start Kit, Roche) at RT for 30 min. Anti-DIG-AP Fab fragments were added into blocking solution (10000× dilution) and membranes were incubated at RT for 30 min. Membranes were washed twice with Maleate acid buffer at RT for 15 min, and then washed once with detection buffer (0.1 M Tris-HCl, 0.1 M NaCl, pH to 9.5) at RT for 5 min. Washed membranes were incubated with 1 ml of CDP-star and exposed to X-film for 3 or 15 min.

2.6 One-step reverse transcription PCR

One-step reverse transcription PCR (one step RT-PCR) was performed with One-step RT-PCR kit (Genemark, Taipei, Taiwan) according to manufacturer's instructions. 100 ng of plant total RNAs were used to conduct one-step RT-PCR analyses in a 20 μl reaction mixture with CEVd-F-96/CEVd-R-95 primer sets. After reverse-transcription at 55°C for 30 min, the PCR condition was performed at 94°C for 5 min, followed by 35 cycles of 94°C for 30 sec, 55°C for 30 sec, and 72°C for 25 sec. The reaction was completed and hold at 4°C after 10 min at 72°C. These reactions were carried out in a Veriti® 96-Well Thermal Cycler (Applied Biosystems, CA, USA). The resulting PCR products were analyzed by electrophoresis on 1.5% agarose gel and detected by ethidium bromide staining.

2.7 Real-time reverse transcription PCR

To quantitate CEVd RNAs and tomato endogenous gene expressions, a real-time reverse transcription PCR (real-time RT-PCR) was employed. Briefly, first-strand cDNAs were synthesized from 1 μg of total RNA
using Moloney Murine Leukemia Virus Reverse Transcriptase (Invitrogen) according to the manufacturer's instructions. The CEVd-R-283 or oligo-dT$_{18}$ primer was used for detection of CEVd RNAs and expressions of host genes, respectively. Then, the PCR was conducted in a 10 μl reaction mixture containing 50 ng of cDNAs, 150 nM forward primer (CEVd-F-100), 150 nM reverse primer (CEVd-R-283) and 5 μl of 2X SYBR-green Mastermix® (Ampliqon, Odense, Denmark). The program for the PCR is performed for amplification at 50°C for 5 min and pre-denaturation at 95°C for 10 min, followed by 40 cycles of 95°C for 15 sec and 60°C for 60 sec. The reaction was completed and hold at 40°C after analyses of melting curve. These reactions and analyses were carried out in a LightCycler 480 machine and software (Roche).

For each run, dedicated software of Real Time instrument plotted the fluorescence intensity against the number of cycles to give the threshold cycle (Ct) value using the automatic method. Standard curves were obtained by linear regression analysis plotting Ct values versus the logarithm of the starting CEVd copy number, enabling estimation by interpolating the individual Ct values in the standard curve.

3. Results

3.1 Quantification of CEVd-infected symptoms of tomato plants

Initially, we sought for establishing a susceptible bio-indicator for CEVd infection and symptom development rather than using its typical exccortis symptom-expressing citrus hosts. Due to different sensitivity and tolerance to CEVd (Matoušek et al., 2007), various tomato cultivars were tested for suitability as CEVd bioindicators (data not shown). We produced dimeric RNA transcripts of CEVd via in vitro transcription for inoculation. Then, we maintained CEVd-inoculated tomato plants in growth chambers at high temperature and light intensity to enhance symptoms development of viroid-infected plants as described in Materials and Methods (Grasmick and Slack, 1985; Handley and Horst, 1988; Harris and Browning, 1980; Skoric et al., 2001). Among these, Rutgers (a sensitive tomato cultivar) and Double-Fortune (a local tomato cultivar in Taiwan) invariantly showed systematic symptoms (see below) and, thereafter, were chosen for further comparison on viroid symptomatic development and pathogenesis.

The early stage of CEVd-infected symptoms exhibited epinasty and leaf curl inward on the upper leaves of CEVd-infected Rutgers at 3 weeks post-infection (wpi) and showed further severity at 4 wpi (Fig. S1). At 6 wpi, in addition to the severe epinasty, leaf deformation and size reduction were observed on the upper leaves of CEVd-infected plants (Fig. 1A). Moreover, the lower leaves also showed epinasty and vein necrosis (Fig. 1B and S2), and the whole plant was prominently stunted (Fig. 1A left sides). On the contrary, CEVd-infected Double-Fortune did not often showed severe CEVd-infected symptoms in comparison with Rutgers (Fig. 1 right sides). Moreover, less reduction in the plant height induced by CEVd-infection is observed in Double-Fortune compared to that in Rutgers (Fig. 1A). These results indicate that Rutgers is more susceptible to CEVd-infection than Double-Fortune.
To quantitate the severity of CEVd-infected tomato plants, the CEVd-infected symptoms displayed at 6 wpi were classified into five categories, such as stunting, leaf size reduction, epinasty, vein necrosis, and chlorosis (Fig. 2A) for bio-indexing. To this regard, the symptom severity of stunting and leaf size reduction were resulted from the reduction of height, length and width; on the other hand, the symptom severity of epinasty and vein necrosis were derived from the percentage of total leaflets showing symptoms. There were 11 levels (0 to 10) for scoring the severity level of index symptoms, and only 2 levels (0 and 1) for chlorosis.

Based on the results of indicator symptoms, the epinasty, leaf size reduction, vein necrosis and stunting of CEVd-inoculated Rutgers exhibited significant discrepancy compared with the mock plants, whereas Double-Fortune did not recapitulate the differences upon CEVd infection (Fig. 2B). Accordingly, it will be useful to enlist Rutgers as a bioindicator for CEVd diagnosis in the inspection and quarantine protocol.

### 3.2 Rutgers as an indicator plant for CEVd infected citrus crops

To test the applicability of Rutgers tomato as a CEVd bio-indicator for viroid-infected citrus plants in the farming field, same amounts of total RNAs extracted from the exocortis symptom-displaying or symptomless citrus plants (the citrus samples were collected from the citrus field in Douliu, Taiwan) (Fig. 3A) were inoculated to indicator Rutgers tomato plants. After 4 wpi, Rutgers started to display certain index symptoms (Fig. 3B). These results confirm that Rutgers can be employed as an indicator plant for diagnosing CEVd-infected citrus plants.

### 3.3 An easy and nano-scaled diagnosis for CEVd by dot blot hybridization

We generated and tested the DNA probes specific and sensitive for dot blot hybridization of CEVd to provide an easily molecular diagnosis. Firstly, we demonstrated that the probe displayed reasonably high specificity among different viroids (Fig. 4A). The sensitivity for the CEVd probe in dot blot hybridization was 0.41 ng. The result indicates that dot blot hybridization with this probe employed in this study provides a high specificity and sub-nanogram sensitivity for detection of the tested CEVd.

To mimic the actual RNA composition, a mixture of CEVd RNA transcripts and healthy tomato total RNAs at a ratio of 1:100 was used to establish a standard curve of the dot blot hybridization for CEVd RNA copy number. The result suggests that $10^9$ copies of CEVd RNAs is the detection limit for detecting CEVd in the dot blot hybridization (Fig. 4B).

To test the reliability which is more appropriate for diagnostic techniques of the dot blot hybridization for detecting CEVd in CEVd-infected plant samples, total RNAs from the CEVd-infected tomato and the CEVd-infected *Gynura aurantiaca* were isolated and then subjected to the dot blot hybridization. We obtained specific CEVd signals with limitations at 31 and 12 ng of total RNAs from the CEVd-infected tomato and the CEVd-infected *Gynura aurantiaca*, respectively. (Fig. 4C). In light of the standard curve performed above (Fig. 4B), the converted CEVd copy numbers for both infected plant samples are $4.8 \times 10^7$ and $9.2 \times 10^7$ per ng of citrus and *Gynura aurantiaca* RNA, respectively.
Total RNAs isolated from healthy or suspected CEVd-infected citrus samples displaying exocortis symptom collected in the farming field were also subjected to dot blot hybridization (Fig. 4D). We determined that the CEVd copy number in the tested citrus was around \(1.4 \times 10^7\) in per 1 ng of citrus total RNAs.

### 3.4 One-step RT-PCR provides a fast, sensitive and accurate way for CEVd detection

To date, PCR-based diagnostic approaches are commonly used to provide fast, sensitive, and accurate diagnostic methods for detecting viral pathogens. We hereof developed one-step RT-PCR and real-time RT-PCR for CEVd detection. Using \textit{in vitro} transcribed CEVd dimers as templates, the RNA titration of one-step RT-PCR shows that the sensitivity of CEVd RNA transcripts was less than \(10^8\) copies (Fig. 5A). To test the applicability of the one-step RT-PCR in the detection of CEVd-infected plants, the total RNAs extracted from CEVd-infected or mock tomato plants were subjected to the one-step RT-PCR analysis as described in Materials and Methods. We demonstrated that CEVd could be detected from CEVd-infected tomato plants by the one-step RT-PCR whereas the mock control showed negative signal (Fig. 5B). The detection limitations for both CEVd-infected tomato and \textit{Gynura aurantiaca} plants were as low as 10 ng of total RNA (Fig. 5C) and the copy number per ng of the total RNAs is calculated as \(4.8 \times 10^7\) and \(2.0 \times 10^7\) copies, respectively. Citrus samples displaying typical exocortis symptoms also showed positive results in the one-step RT-PCR with a calculated concentration of \(4.5 \times 10^7\) copies per ng of total RNA (Fig. 5C). Further sequence analysis showed that the AD3 CEVd strain found in Taiwan shares 98% identity to the CEVd strain used in this study (M34917-G74\(\Delta\); \textit{(Daros and Flores, 2004)}), differing in 6 nucleotides (Fig. 6).

### 3.5 Quantitative detection of CEVd by real-time RT-PCR

Next, we established a standard curve based on the amplification curves and the melting peaks of the real-time RT-PCR for CEVd transcripts as described in the Materials and Methods, which enables directly and absolutely measuring the quantity of CEVd (Fig. 7A). In the standard curve, a linear correlation was observed for six logarithmic concentrations with an R square value of 0.995, indicating an intimate reliability for calculating CEVd’s quantities within the range of the examined concentrations. Purified total RNAs from tomato and citrus were subjected to the test. The results of the RNA titration real-time RT-PCR suggest that the detection limits are \(5.9 \times 10^6\) and \(3 \times 10^6\) copies in per 1 ng of total RNA for tomato and citrus plants, respectively (Fig. 7B). Together, these data indicate that real-time RT-PCR confers more sensitivity than other methods, such as one-step RT-PCR and dot blot hybridization, for detecting CEVd.

### 3.6 The correlation between the level of CEVd RNAs/sRNAs and CEVd-infected symptoms

The discrepancy on the symptom expression of CEVd-infected tomato plants was noticed between Rutgers and Double-Fortune as documented in Fig 1. To explore the potential culprit leading to the discrepancy, we first examined the viroid genomic RNA level in the both CEVd-infected tomato plants. However, there is no significant difference on the accumulative viroid genomic RNA level and the
expression of CEVd-derived small RNAs (sRNAs) between CEVd-infected Rutgers and Double-Fortune by Northern hybridization and real-time PCR (data not shown). These results indicated that CEVd enables successfully replicated in both tomato cultivars despite Double-Fortune might be more tolerant than Rutgers in response to CEVd infection. In addition, symptoms on CEVd-infected Rutgers are prominent but diverse at 6 wpi (Fig. 8A). The leaves exhibiting various degrees of symptoms (mild and severe, also being indicated by the index described above) at the same position of CEVd-infected Rutgers were collected and were subjected to comparison of the levels of CEVd genomic RNAs and sRNAs. As a result, the leaves with severe symptoms appeared to have accumulated higher levels of CEVd genomic RNAs and sRNAs than the leaves with mild symptoms (Fig. 8B). In addition, the level of CEVd sRNAs is also associated with level of the viroid genomic RNA (Fig. 8C). Interestingly, a positive correlation between CEVd sRNAs and symptom severity was found. Taken together, our results implicate that the severity of CEVd-induced symptoms may be associated with higher amount of replicative CEVd genome and/or accumulative viroid-derived small RNAs.

4. Discussion

4.1 Bio-index and molecular methods for detection

The initial attempt of this study is to seek an indicator plant for CEVd inspection and quarantine uses. Although CEVd readily infects tomato plants, distinguishable symptoms among various tomatoes cultivars, for example, on both Rutgers- and Double-Fortune tomato plants, exhibit statistical discrepancy on the severity and progression of the disease (Fig. 1 and 2). Our finding not only established a bioindicator for CEVd symptom expression, but also provided a basis for understanding the molecular mechanism underlying CEVd pathogenesis. Hence, in need of a reliable standard of protocol for CEVd
detection/diagnosis, we first established an integrated diagnostic approach, including bio-index in a
tomato cultivar Rutgers, dot blot hybridization with CEVd molecular probes, one-step RNA and real-time
RT-PCR, as a reliable, accurate, and rapid method used for CEVd detection.

4.2 Tomato plant as a bioindicator for CEVd

Among susceptible hosts, the suitability of tomato plant on serving as an indicator plant for CEVd is
attributed by several advantages as shorter growth time, small size, rapidness for symptom expression
(4-6 weeks), and a model organism for plant science. Firstly, it takes shorter time, i.e. 4-6 weeks, to display
CEVd-related symptoms on tomato plants than on citrus plants for more than 6 months. Secondly, it is
easier to obtain more information from tomato due to its property as a model plant; lastly, there are also
more genetic or molecular tools for studies of tomato plant so that studies of the interactions between
viroid and tomato in the future can be more easily achieved.

After inoculation of CEVd transcripts, cv. Rutgers showed typical symptoms of PSTVd or CEVd, which
were consistent with previous studies (Harris and Browning, 1980; Owens and Hammond, 2009; Qi and
Ding, 2003; Sano et al., 1992). Although various variants of CEVd might be limited to specific hosts, in our
case, CEVd isolated from citrus in field could infect tomato cv. Rutgers systemically and then caused
epinasty, vein necrosis, and stunting at 4 to 6 wpi. Hence, CEVd-sensitive cv. Rutgers may be a great
indicator plant host for CEVd diagnosis, amplification of CEVd quantities, and further studies of viroids,
such as distinguishing viroid variants of different severity and/or investigate the pathogenic nucleotides
and domains of viroids (Morris and Wright, 1975; Owens et al., 1996; Qi and Ding, 2003; Sano et al.,
1992). On the other hand, the fact that Double-Fortune is not a good indicator plant for viroids potentially
makes it a better candidate for studying essential factors involved in tolerance to viroids. Also, the
tolerant cultivar could be bred with other high-value or sensitive cultivars to obtain new cultivars that
might be more commercially profitable with tolerance to viroid infection.

4.3 CEVd isolated from citrus in the farming citrus field

The sequences were not fully identical between CEVd used in this study (M34917-G74Δ) and CEVd from
AD3 citrus in field (Fig. 6); however, the nucleotide change of primer-binding sites might not be obtained
unless another set of primers is used. Because the symptoms of tomato plants infected by M34917-
G74Δ and AD3 were mostly the same, it might indicate that the nucleotide changes did not affect the
pathogenicity of these two CEVd variants on tomato host. In addition, CEVd from two tomato plants
infected by AD3 citrus total RNA were identical to the initial inoculum, AD3, at least in the sequenced
region. In addition, based on nearly 98% sequence identities between AD3 and the Chinese variant
DQ431991, Citrus exocortis viroid isolate CEVd-XNM-ZK Guangdong-YC (data not shown), the AD3
variant might be imported from China or vice versa.

4.4 Implications for CEVd pathogenesis
We found a positive association in the accumulation of CEVd gRNAs with the symptomatic severity of CEVd-infected Rutgers tomato plants (Fig. 8). In contrast, it insignificantly recapitulated these differences between the two tomato cultivars, Rutgers and Double-Fortune (data not shown) although they exhibited distinguishable symptomatic expressions (Fig. 1 & 2), which could result from the genetic variation of two tomato cultivars. One study has indicated that varied genotypic tomato plants render different susceptibility to viroid infection, which is dispensable at the levels of viroid replication in sensitive and tolerant plants (Matoušek et al., 2007). It is thus desirable to further identify other host factors which may lead to the noticeable differences in symptom expression on these two cultivars.

5. Conclusions

For the diagnostic purpose, we statistically assessed the CEVd-infected symptoms in tomato cultivars Rutgers and Double-Fortune in search of a suitable bio-indicator for CEVd disease. Base on the selected five bio-indexes comprised epinasty, vein necrosis, stunting, and reduction of leaf length and width, tomato cv. Rutgers was demonstrated to serve as a better indicator plant for CEVd in comparison with Double-Fortune. In addition, combination with molecular detections of CEVd RNAs, including dot blot, one-step RT-PCR and real-time RT-PCR, we were able to monitor the existence, distribution, amount as well as even pathogenicity and pathogenesis of CEVd. Among these, the sensitivities of three molecular methods were compared and showed that real-time RT-PCR conferred the best sensitivity of $10^7$ copies of CEVd. On the other hand, indicator plants and bio-index could help recognition of different severity of viroids and were good models for further studies in the mechanism of viroid pathogenesis. To sum up, for further researches of viroid pathogenesis and viroid-host interactions, indicator plants and real-time RT-PCR were more appropriate; for larger scale of detection work of quarantine, dot blot and one-step RT-PCR seem to be more suitable, due to the lower cost of time and money.

Abbreviations

ASSVd: Apple skin scar viroid

CBCVd: Citrus bark cracking viroid

CBLVd: Citrus bent leaf viroid

CCCVd: Coconut cadang cadang viroid

CChMVd: Chrysanthemum chlorotic mottle viroid

CDVd: Citrus dwarfing viroid

CEVd: Citrus exocortis viroid

CVd-V: Citrus viroid V
Declarations

Ethics approval and consent to participate
Not applicable

Consent for publication
Not applicable

Availability of data and materials
Not applicable

Competing interests
The authors declare that they have no competing interests.

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Authors’ contributions
CH Lin, I Hu and CY Lin carried out the experimental work under the supervision of TH Hung and TL Shen. TH Hung, ML Wu, TL Shen participated in the experiment design and analyzed data. CH Lin and TL Shen wrote the manuscript. All authors have read and approved the final manuscript.

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Figures
Figure 1

Symptoms of susceptible tomato plants infected by Citrus exocortis viroid (CEVd). (A) The symptoms of CEVd-inoculated tomato plants (S. lycopersicum cv. Rutgers and Double-Fortune) at 3 or 6 week post-inoculation (w.p.i). The scale bar = 10 cm. (B) The upper leaves of CEVd-infected symptoms on tomato plants. The scale bar = 5 cm.
Figure 2

Symptoms on CEVd-infected Rutgers tomato plants as a bio-index. (A) Categories of CEVd-infected symptoms showed on Rutgers tomato plants and could be used as a bio-index for CEVd infection. (B) Quantification of CEVd-infection symptoms on Rutgers and Double-Fortune at 6 wpi. Error bars indicated ± standard deviation of the mean of at least three times experiments. Statistically significant differences
for values were determined by Student’s t test (* P < 0.05, ** P < 0.01, *** P < 0.001). M: mock-inoculated tomato plants; I: CEVd-inoculated tomato plants. The scale bar = 5 cm.

Figure 3

A.

Symptomless citrus

Citrus exocortis

B.

mock exocortis citrus RNA-inoculated

mock exocortis citrus RNA-inoculated

Figure 3

Symptoms of citrus and Rutgers tomato plants caused by CEVd in the field. (A) Exocortis and symptomless citrus from the field. (B) Field-collected citrus samples were inoculated on Rutgers index
plants. Right panel displays a close-up view of the upper leaves. The photographs were taken at 4 wpi. Scale bar left: 10 cm; left: 5 cm.

Figure 4

A.

| H₂O | 41 | 4.1 | 0.41 (ng) | PSTVd | ASSVd | CCCVd | CChMVd |
|-----|----|-----|---------|--------|-------|--------|---------|
| CEVd probe |

B.

Healthy tomato RNA

| 10¹² | 10¹¹ | 10¹⁰ | 10⁹ | 10⁸ | 10⁷ | 10⁶ | 10⁵ | 10⁴ | 10³ | 10² | 10 | 0 | + | H₂O | H₂O |
|------|------|------|-----|-----|-----|-----|-----|-----|-----|-----|----|---|---|-----|-----|

C.

| CEVd-inoculated tomato plant | CEVd-infected Gynura aurantiaca plant |
|-------------------------------|---------------------------------------|
| 1000 500 250 125 63 31 16 8  | 200 100 50 25 12 6 3 1 (ng)          |

D.

Citrus samples from the field

| AH1 | AH2 | AH2 | AD3 | AD4 | DG | H₂O | + |

Figure 4

Dot blot hybridization of CEVd. (A) Probe specificity test for CEVd dot blot hybridization detection. (B) Dot blot hybridization sensitivity test for CEVd detection. (C) Detection of CEVd-infected tomato and Gynura aurantiaca RNAs by dot blot hybridization. (D) Detection of CEVd-infected citrus from the farming field by
dot blot hybridization. +: CEVd plasmid; DT: CEVd-infected tomato plant; DG: CEVd-infected Gynura aurantiaca; AH1-AD4: field samples showing no symptoms (AH) or suspected symptoms (AD).

Figure 5

**A.**

Healthy tomato RNA

|                | +  | +  | +  | +  | +  | +  | +  | +  | +  | M  | 1012 | 1011 | 1010 | 109  | 108  | 107  | 106  | 105  | 104  | 103  | 102  | 101  | -    | +    | H2O  |
|----------------|----|----|----|----|----|----|----|----|----|----|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|
| bp             | 500| 250| 500| 250| 500| 250| 500| 250| 500| 250| 500  | 250  | 500  | 250  | 500  | 250  | 500  | 250  | 500  | 250  | 500  | 250  | 500  | 250  | 500  | 250  |

**B.**

| CEVd-inoculated tomato plant | CEVd-infected Gynura aurantiaca plant |
|-------------------------------|---------------------------------------|
| bp                            | M 100 10 1 0.1 100 10 1 0.1 100 10 1 0.1 |
| 500 400 300                   | + H2O CEVd: 370 bp                    |

**C.**

Citrus samples from field

| bp | AH1 AH2 AD4 AD3 Transcripts | H2O M |
|----|-----------------------------|-------|
| 500|                            |       |
| 250|                            |       |

**Figure 5**

CEVd detection by one-step RT-PCR. (A) One-step RT-PCR sensitivity test for CEVd detection. (B) Detection of CEVd-infected tomato plants by one-step RT-PCR. Elongation factor 1 alpha (EF-1α) was used as an internal control. (C) Detection of CEVd-infected citrus from the farming field by one-step RT-PCR. +: CEVd
plasmid; DT: CEVd-infected tomato plant; DG: CEVd-infected Gynura aurantiaca; AH1-AD4: field samples showing no symptoms (AH) or suspected symptoms (AD).

Figure 6

Sequence comparison between CEVd M34917-G74Δ strain and a CEVd isolate from the field. Secondary structures of the minimum free energy at 24°C of CEVd strain (M34917-G74Δ) were predicted by the Mfold program (Walter et al., 1994; Zuker, 2003). CEVd-F96 and CEVd-R95 primers were used in one-step RT-PCR and sequencing. The nucleotide variations between M34917-G74Δ and a CEVd isolate from citrus farming field were indicated. TL (Terminal left), P (Pathogenicity), C (Central), V (Variable) and TR (Terminal right) refer to the structural domains of the viroid secondary structure (Keese and Symons, 1985).
Detection of CEVd-infected samples by real-time RT-PCR. (A) Threshold cycle (Ct) and amplification curve established the standard curve for CEVd real-time RT-PCR analyses with ranging from 10^7 to 10^12 copy numbers of CEVd transcripts. (B) Ct values were obtained from real-time RT-PCR amplification using 50 ng of tomato and citrus total RNAs. H: healthy plant; D: diseased CEVd-infected plant. Error bars indicated ± standard deviation of the mean of three times of experimental replicate.
Figure 8

Symptom development on CEVd-infected Rutgers is associated with the levels of CEVd genomic RNA (gRNAs) and CEVd-derived small RNAs (sRNAs). (A) Varied degrees of CEVd-infected symptoms on Rutgers tomatoes. The photographs were taken at 6 w.p.i. Scale bar = 5 cm. (B) CEVd viroid genomic RNA and small RNA were analyzed by denaturing northern blot. U6 snRNA was used as an internal control. (C)
The relative expression levels between CEVd gRNAs and sRNAs. (D) The gRNAs and sRNAs of CEVd were plotted against index of symptom severity of the two CEVd-infected Rutgers samples.

**Supplementary Files**

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- Fig.S1LinCHetal.pdf
- Fig.S2ALinCHetal.pdf