Analyses of virus/viroid communities in nectarine trees by next-generation sequencing and insight into viral synergisms implication in host disease symptoms

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We analyzed virus and viroid communities in five individual trees of two nectarine cultivars with different disease phenotypes using next-generation sequencing technology. Different viral communities were found in different cultivars and individual trees. A total of eight viruses and one viroid in five families were identified in a single tree. To our knowledge, this is the first report showing that the most-frequently identified viral and viroid species co-infect a single individual peach tree, and is also the first report of peach virus D infecting *Prunus* in China. Combining analyses of genetic variation and sRNA data for co-infecting viruses/viroid in individual trees revealed for the first time that viral synergisms involving a few virus genera in the *Betaflexiviridae*, *Closteroviridae*, and *Luteoviridae* families play a role in determining disease symptoms. Evolutionary analysis of one of the most dominant peach pathogens, peach latent mosaic viroid (PLMVd), shows that the PLMVd sequences recovered from symptomatic and asymptomatic nectarine leaves did not all cluster together, and intra-isolate divergent sequence variants co-infected individual trees. Our study provides insight into the role that mixed viral/viroid communities infecting nectarine play in host symptom development, and will be important in further studies of epidemiological features of host-pathogen interactions.

Peach is one of the most widely grown fruit crops in China, and nectarine (*Prunus persica cv. nectarina*) is an important cultivar of peach. Viruses and viroids can cause significant negative effects on fruit quality and yield in peaches and nectarines. Previous reports have shown that apple chlorotic leaf spot virus (ACLSV), plum pox virus (PPV), plum necrotic ringspot virus (PNRSV), prune dwarf virus (PDV), apple mosaic virus (ApMV), plum bark necrosis stem pitting-associated virus (PBNSpaV), peach latent mosaic viroid (PLMVd), and hop stunt viroid (HSVd) are the major pathogens that infect these trees. Yu et al. (2013) performed a large-scale field survey of the major viruses and viroids that infect peach trees in China using RT-PCR and ELISA, and the results showed that only ACLSV, PNRSV, cherry green ring mottle virus (CGRMV), apricot pseudo-chlorotic leaf spot virus (APCLSV), PLMVd, and HSVd were detected. However, next-generation sequencing (NGS) approaches have opened new avenues in recent years for the identification of viruses and viroids (including novel pathogens), and this technology is well suited for large-scale pathogen surveys, not only because it can increase the speed at which a wide range of known pathogens are detected, but also because it can be used to detect newly-emerging or potential pathogens for which other diagnostic tools are not yet available. Several NGS-based strategies have been developed to overcome problems with traditional approaches, and these have resulted in the identification of known and novel viruses in peach. These include two novel luteoviruses, nectarine stem-pitting-associated virus

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were identified in sample T01. (Fig. 2A). This is also the first report of the identification of PeVD in the dominant viruses/viroids in sample T02, accounting for 63.75% (PBNSPaV) and 19.74% (PLMV) of the total respectively) were the dominant viruses/viroids identified in sample T01, while these same two pathogens were also in China.

Ampelovirus PBNSPaV in the genus Rusty mottle virus (CNRMV) in the Closterovirus as APV1 and APV3 that corresponded to segments with high sequence similarities), CGRMV, cherry necrotic ring spot virus—green mottle virus (CRGRMV), and Tymoviridae-

In addition, complex mixed infections have been found among fruit tree-infecting viruses2,5,12,13. Thus, the potential contribution of each single virus infection to the symptoms observed cannot easily be associated with a disease in the infected Prunus trees. In fact, many horticultural plants that are routinely clonally propagated are reservoirs of a large variety of viruses and viroids. The importance of the virome in mammalian biology, and the emerging concept of virome-host interactions and their relationship to host genetics was first described by Virgin (2014)14. The virome of the microbiome interactions with the host, especially in mammalian biology, has recently become a hot research topic that relies on bioinformatic tools and NGS technology14–18. However, only a limited number of studies have revealed viral communities or viromes in peach17. In this study, we used NGS technology to study the viral communities in nectarine trees with different disease phenotypes. We identified both known and novel viruses and viroids, and performed comparative analyses of the potential contribution of the pathogens to disease symptoms. Our results will extend the range and kinds of virus and viroid species that infect peach trees, and provide insight into the viral synergisms and the agents that might be associated with disease symptoms in nectarine.

Results

Virus and viroid accumulation and pathogen communities within individual nectarine trees. In the five nectarine tree samples, T01 and T02 were collected in greenhouse #1 from the same nectarine cultivar ‘Youtao 1233’ (10-year-old trees), while T03, T04, and T05 were collected in greenhouse #2 from nectarine cultivar ‘Zhongyou 4’ (5-year-old trees). The five samples came from trees that showed different leaf and fruit symptoms (Table 1, Fig. 1).

To perform comparative analyses of the different symptoms observed in the nectarine trees, we used NGS of the sRNAs extracted from the five samples to obtain a complete survey of the virus and viroid communities infecting each tree. The Illumina reads obtained from sequencing the five cDNA libraries, which were prepared using RNA extracted from the scion parts of the grafted trees, yielded between 22,434,184 and 30,438,485 raw sRNA reads per library. From these, we obtained between 20,906,239 and 28,435,618 clean reads for samples T01 to T05 (Table S1). A virus and viroid library was constructed from virus and viroid genomes available from NCBI and was then used for mapping of the sRNA reads using the short-sequence alignment program Bowtie. The majority of the reads were 18 to 25 nt in length, with most being either 21 nt or 22 nt. De novo assembly of the sRNAs and blastn and blastx searches resulted in assemblies of 15 to 744 contigs, with lengths ranging from 33–474 nt, that were associated with known viruses/viroids. The virus/viroid-associated reads per sample ranged from 1.11% to 9.60% of the clean sRNA reads for the five samples. We found that samples T01 and T02 from greenhouse #1 had the highest number of virus/viroid-associated reads, with 5.53% and 9.60% of the clean sRNA reads, respectively, while the virus/viroid-associated reads in samples T03, T04, and T05 ranged from 1.11% to 2.71% (Table S1).

To compare the different viral communities and the relative numbers of individual viruses and viroids in each sample, we examined individual contig numbers for previously-identified viruses and viroids and calculated the percentage of individual virus or viroid-associated reads by dividing the number of virus or viroid-associated reads by the total number of clean sRNA reads (x 100).

The virus and viroid communities and the numbers of individual pathogens differed between samples T01 and T02 from trees that had different fruit symptoms. Sample T02 had the most identified viral species, and also the highest number of contigs for one viroid and nine virus genera in five families; these included PLMVd in the genus Pelamovirus, (family Avsunviroidae); two unassigned viruses NSPaV and PaLV (family Luteoviridae); BSNPaV in the genus Ampelovirus (family Closteroviridae); ACLSV in the genus Trichovirus and APVs in the genus Foveaviridae (most conspecifics were identified to correspond to segments, but a few contigs were not associated as APV1 and APV3 that corresponded to segments with high sequence similarities), CGRMV, cherry necrotic rusty mottle virus (CNRMV) in the family Tymoviridae—, and another 11 contigs that may represent a novel unknown virus in the family Tyomoviridae—which showed identities to segments from marafi-, tymo-, and maculaviruses with low sequence similarities (Fig. 2A). Excluding CGRMV and CNRMV, other seven viruses (PLMVd, NSPaV, PaLV, BSNPaV, ACLSV, APVs, PeVD) were identified in sample T01. (Fig. 2A). This is also the first report of the identification of PeVD in Prunus trees in China.

Table 1. Nectarine tissue samples used for NGS analyses of small RNAs.

| Sample | Origin        | Cultivar       | Leaf and fruit symptoms                                   |
|--------|---------------|----------------|----------------------------------------------------------|
| T01    | Greenhouse 1  | Youtao 1233    | Leaf bleaching, asymptomatic fruit                        |
| T02    | Greenhouse 1  | Youtao 1233    | Leaf bleaching, fruit pitting                            |
| T03    | Greenhouse 2  | Zhongyou 4     | Leaf and fruit asymptomatic                               |
| T04    | Greenhouse 2  | Zhongyou 4     | Leaves asymptomatic, fruits dimpled                       |
| T05    | Greenhouse 2  | Zhongyou 4     | Leaves with chlorotic mottling, asymptomatic fruit       |

(BNPVs)27,8 and peach-associated luteovirus (PaLV)9; two marafivirus, nectarine virus M (NeVM)9; and peach virus D (PeVD)10; a novel fabavirus in the family Secoviridae, peach leaf pitting-associated virus (PLPaV)11, and three very similar members of the genus Foveovirus, asian prunus virus 1 (APV1), APV2, and APV312.

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virus and viroid-associated reads. In addition, CGRMV (5.77% of the total virus-associated reads) and CNRMV (5.77% of the total virus-associated reads) were only detected in sample T02 (Fig. 2B,C). The number of contigs in T01 and T02 that mapped to the PBNSPaV genomes in GenBank was different, and indicated the presence of divergent sequence variants in the two samples (Table 2). We also found that the number of contigs that mapped to several isolates of PBNSPaV (Phm-WH-3, WH-1, PR258-2) collected from plum trees with disease symptoms19,20 were significantly increased in sample in T02 (Table 2). By comparing the number of assembled viral contigs and the percentages of virus-associated reads between samples T01 and T02, we found that the T02 sample collected from fruit pitting tree had the higher levels reads of the viruses PBNSPaV, CGRMV, and CNRMV and the higher mapping number of contigs of PBNSPaV isolate from sample with disease symptoms.

The viral communities and the numbers of the individual virus varied significantly in the three samples with diverse disease phenotypes (T03, T04 and T05). In the asymptomatic sample T03, we only identified contigs associated with two known viruses/viroids (PLMVd and NSPaV), while in sample T04, which had dimpled fruits, we identified contigs from another two viruses (ACLSV and APV2) in addition to PLMVd and NSPaV (Fig. 3A). Figure 3(B,C) shows that PLMVd and NSPaV accounted for the highest percentages of the individual pathogen-associated reads and were the dominant pathogens in samples T03 and T04. From these results, we can infer that higher levels of ACLSV and APV2 co-infection or their interactions with PLMVd or NSPaV may be associated with the fruit dimpling symptoms seen in sample T04. Sample T05 with chlorotic mottle leaf symptoms contained mainly PLMVd sequences (14 assembled contigs, 2.27% of the total clean reads), and one contig that showed identity to sequences related to a segment (68/84 nt) of GRGV in the genus Maculavirus, (family Tymoviridae), but the read number was very low (0.0012% of the total clean reads) and this will need to be further confirmed by subsequent RT-PCR to determine whether the low level of reads observed resulted from sample contamination (Fig. 3).

Confirmation of identified viruses and viroids by RT-PCR. In order to determine whether the viruses and viroids identified by NGS were actually present in the five nectarine tree samples, we conducted RT-PCR using the primer pairs specific for the individual viruses/viroids (Table S2). We found that it was very difficult to distinguish between very closely related viruses such as APV1, APV2, and APV3, and CGRMV and CNRMV, and to identify the presence of viruses that had very few and short contigs with low read numbers in the NGS data. The RT-PCR results obtained, followed by Sanger sequencing, indicated that excluding APV3 and grapevine red globe virus (GRGV), the samples were positive for all other viruses identified by NGS, suggesting that these viruses were actually present in the sRNA extracted from the nectarine tree samples (Fig. 4). In order to further confirm the results of the NGS screen, we also detected individual viruses/viroids in 36 samples collected from different greenhouses and cultivars using RT-PCR assays, and these results are summarized in Table S4. RT-PCR detection indicated that the other test samples (T01-type: N8, N8-2; T02-type: N9, N9-2; T05-type: P12, P13, P14; T03-type: P25, P26, P27; T04-type: P20, P21, P23, P24) with disease symptoms similar to those from the five individual trees (T01, T02 and T03, T04, and T05), respectively, had uniform virus and viroid communities that
agreed with the NGS data. The different mixed virus/viroid communities which were detected in the different cultivars were also confirmed.

**Phylogenetic analysis of the identified viruses/viroids.** We identified PLMVd infection in all five tested nectarine trees, and 17 complete genome sequences of PLMVd isolates were obtained from the five samples by RT-PCR and cloning. A combined phylogenetic analysis of the genomic sequences of the PLMVd isolates from this study with some PLMVd sequences in GenBank gave three major phylogroups, and we found that the PLMVd sequences associated with symptomatic and asymptomatic trees did not all cluster together, except for two previously-reported peach calico (PC) isolates which clustered alone in Group II (Fig. 5). Further pairwise comparisons of the 17 PLMVd genome sequences showed that the PLMVd isolates from the nectarine samples are quite divergent, and share between 80.0 and 99.7% nucleotide identity. Of the five sampled trees, three sequences

![Figure 2. Confirmation of the identified viruses and viroids by RT-PCR. RT-PCR was used to amplify partial sequences of (A) APV1, (B) APV2, (C) APV3, (D) ACLSV, (E) CGRMV, (F) CNRMV, (G) GRGV, (H) PBNSPaV, (I) PeVD, (J) NSPaV, (K) PaLV, and (L) PLMVd using virus or viroid-specific primer pairs from nectarine tree RNA samples T01, T02, T03, T04, and T05. Amplified DNA fragments were examined by electrophoresis on 1.5% agarose gels and stained with EtBr. M, DNA size 2000 marker; N, negative control.](image)

| Genome             | Host            | Symptoms                                | Origin   | Contig numbers |
|--------------------|-----------------|-----------------------------------------|----------|----------------|
| KJ792853.1 isolate Plm-WH-3 | Plum            | Trunk gummosis, cracking, necrosis (19) | China    | 24 47          |
| KJ792852.1 isolate WH-1 | Peach           | Trunk gummosis, cracking, necrosis, stem pitting (19) | China    | 29 49          |
| KC590346.1 isolate FR258-2 | GF305 (Peach)   | Peach red marbling disease (20)         | France   | 49 52          |
| HG917400.1 isolate VC1 | Apricot         | Unknown                                 | Italy    | 6 15           |
| EF546442.1  | *Prunus domestica* | Small, chlorotic, distorted leaves | USA      | 13 18          |
| KC590345.1 isolate Pair-2 | GF305 (Peach)   | Unknown (20)                            | France   | 13 10          |
| KC590344.1 isolate Tatao25 Q-375-02 | Peach | Unknown (20) | China    | 10 6           |
| KJ792854.1 isolate GS-3 | Peach | Asymptomatic (19) | China    | 13 9           |

Table 2. Number of contigs that map to PBNSPaV genomes in GenBank using NGS.
from the T03 isolate (asymptomatic sample) that clustered in subgroup IA shared a high degree of nucleotide identity (98.5–98.8%), whereas sequences from the other four sampled trees were highly variable, and clustered into different subgroups, IA to IF (Fig. 5). These results suggest that intra-isolate, genetically distinct sequence variants in individual nectarine trees were present in four of the tree samples (T01, T02, T04, and T05) based on the distribution of sequences in different phylogroups. We conclude that divergent sequence variants that co-infect individual trees is common.

In the same way, we constructed two phylogenetic trees based on one nearly complete genome sequence (excluding the 5’ and 3’ terminal sequences) and 11 RNA-dependent RNA polymerase (RdRp) gene sequences from NSPaV isolates T01, T02, T03, and T04 (Fig. 6A). Three NSPaV RdRp gene sequences from T02 grouped with isolate SK from a nectarine in South Korea22. Other sequences including RdRp and nearly complete genome sequences of the T01, T03, T04 NSPaV isolates were found to be closely related to isolate NSPaV/12P42 derived from a nectarine in the USA5, but three NSPaV RdRp gene sequences from T04 were not consistant, and the T04-1 sequence was detected as a recombinant. Six coat protein (CP) gene sequences from PBNSPaV isolates from T01 and T02 are closely related to the known isolates Plm-WH-3 and WH-119 from peach in China. One PBNSPaV CP gene sequence (T01-1) is closely related to the known isolate GS 19, also from peach in China (Fig. 6B). Four homologs of the heat shock protein 70 (HSP70h) gene sequences from PBNSPaV isolates from samples T01 and T02 are also closely related to the known isolates Plm-WH-3 and WH-1, and three T01 HSP70h gene sequences are also closely related to the known isolates GS (Fig. 6B). Seven CP sequences from the ACLSV...
Figure 4. Identification and quantification of the pathogen-associated sequencing reads for the viruses/viroids infecting tree samples T03, T04, and T05 using sRNA sequencing. (A) The numbers of assembled viral contigs associated with known viruses and viroids in samples T03, T04 and T05 from greenhouse #2. The percentage of virus and PLMVd-associated reads (B,C) in each sample was calculated as described in Fig. 3 for samples T03, T04, and T05.

Figure 5. Phylogenetic analysis of PLMVd isolates. The phylogenetic tree was constructed by the maximum-likelihood (ML) method using MEGA6 software. Bootstrap confidence values (1,000 replicates) are given at the branch nodes. Branches corresponding to partitions reproduced in <75% of bootstrap replicates are collapsed. The PLMVd genome sequences obtained in this study are marked with a red triangle.
isolates from samples T01, T02, and T04 in our study all grouped together with the known isolates Z1 and Z3 from peach trees in China (Fig. 6C). Six CP sequences from the APV1 isolates from T01 and T02 are closely related to each other and to the known isolates D2363 and D2367 (Fig. 6D). The three CP sequences from APV2 isolates from sample T04 are closely related to the APV2 isolate Bonsai12 from Japan (Fig. 7E). The three CGRMV CP sequences from isolate T04 are closely related to isolate F9 from China (Fig. 7F). The three CP sequences from CNRMV isolates from sample T02 are closely related to isolates Pe-WH-18 and Pe-WH-1 from China (Fig. 7G). For the few available sequences in GenBank for PaLV and the newly reported PeVD sequences, we analyzed their phylogenetic relationships using an outgroup. As expected, six RdRp and three CP sequences from PaLV and five PeVD replicase polyprotein gene sequences from isolates T01 and T02 were found to be closely related to the isolates in GenBank (Fig. 7H,I).

Sequence comparisons using the T01-T05 datasets showed that the NSPaV and PBNSPaV sequences in this study are highly variable, with the nucleotide sequence identities among the NSPaV RdRp gene sequences ranging from 89.4 to 99.6%, with comparable values of 93.2 to 99.5% for the PBNSPaV CP gene sequences. This contrasts with other gene sequences from ACLSV, APV1, APV2, CGRMV, CNRMV, PaLV, and PeVD that are highly homologous (96.7 to 100%) in identified isolates.

Figure 6. Phylogenetic analysis of the identified viruses. The phylogenetic trees were constructed as for PLMVd in Fig. 5. The viral gene sequences obtained in this study are marked with a red triangle. (A) The nearly full genome and RdRp gene of NSPaV; (B) the CP and HSP70h genes of PBNSPaV; (C) the CP gene of ACLSV; (D) the CP gene of APV1.
NGS technologies provide a powerful way to detect and identify viral pathogens with no prior knowledge of virus genome sequences\(^3,26\). This technology is finding increased applications in revealing the viromes that contribute to host phenotype and also the recent evolutionary history of RNA viruses\(^14\text{--}18,27,28\).

In this work, we studied co-infecting virus and viroid communities using NGS technology in five individual trees of two nectarine cultivars associated with different disease phenotypes, and then confirmed the identified viruses/viroids using RT-PCR in 36 samples from four cultivars. Identifying comparable plant materials with the same genetic background and cultivation history associated with the different disease symptoms can be difficult. In this study, some valuable samples were collected from trees in two small greenhouses that were grown under the same environmental conditions, with the same cultivation methods, pesticides, and fertilizers, which allowed for a direct comparison of the effects of co-infecting virus and viroid communities in trees with different disease phenotypes. The results indicated that the viral species present in viral communities isolated from the four different cultivars were diverse. A single tree, T02 of cultivar ‘Youtao 1233’ that showed symptoms of fruit pitting, harbored most of the nine different viruses/viroids from five families including PLMVd, NSPaV, PaLV, PBNSPaV, ACLSV, APV1, CGRMV, CNRMV, and PeVD, while T01, that did not show fruit pitting, harbored seven viruses/viroids from the above group in addition to CGRMV and CNRMV. To our knowledge, this is the first report that most identified viral species form co-infections in a individual peach tree, and also the first report of PeVD in peach in China. These results confirmed and extended the utilization of NGS to detect fruit tree viruses and to provide further insight into the complex multiple infections in individual trees and between different cultivars.

**Figure 7.** Phylogenetic analysis of the identified viruses. The phylogenetic trees were constructed as for PLMVd in Fig. 5. The viral gene sequences obtained in this study are marked with a red triangle. (E) The CP gene of APV2; (F) the CP gene of CGRMV; (G) the CP gene of CNRMV; (H) the CP and RdRp genes of PaLV; (I) the CP gene of PeVD. NSPaV and barley yellow dwarf virus (BYDV) were used as outgroups in the PaLV tree. Grapevine rupestris vein feathering virus (GRVFV) and maize rayado fino marafivirus (MRFV) were used as outgroups in the PeVD tree.

**Discussion**

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Combined genetic variation and sRNA data analyses of the co-infecting viruses/viroids in this study implied that viral synergisms and divergent sequence variants play an important role in determining disease symptoms. This includes synergisms among a few genera in the viral families Betaflexiviridae, Closteroviridae, and Luteoviridae and the possible effects on facilitating divergent virus sequence variants in nectarine symptom expression, increasing the titers of pathogenic viral genotypes.

Two functional classes of viral genes, a transcription-related RdRp gene and a structural CP gene, are often used for phylogenetic classification of plant viruses. The HSP70h genes are used for phylogenetic classification of plant viruses in the family Closteroviridae. A study by Qu et al. showed that the evolutionary relationships of global PBNSPaV isolates can be reliably inferred using HSP70h sequences.

Phylogenetic analyses of the identified 10 viruses and one viroid in the five nectarine trees using whole genome sequences, nearly complete genome sequences, and CP, RdRp, and HSP70h gene sequences showed that PLMVd, NSPaV, and PBNSPaV gene sequences were more divergent, and presented different sequence variants or recombinants. However, gene sequences from ACLSV, APV1, APV2, CGRMV, CNRMV, PaLV, and PeVD from the five individual trees showed much less variability. It should be noted that full genome sequences of the identified viruses could not be obtained in this study; however, the phylogenetic relationships were analyzed using two functional classes of viral genes that might be involved in viral synergisms.

Stem pitting has been diagnosed in peaches infected by tomato ringspot virus (ToRSV) and PBNSPaV, and in nectarine infected by NSPaV. The PBNSPaV peach isolate WH-1 causes trunk gummosis, cracking, necrosis, and stem pitting symptoms. Based on the results of our phylogenetic analysis, CGRMV and CNRMV are most similar to apple stem pitting virus, but the effects on fruit pitting in Prunus have not been reported.

Compared with sample T01 (no fruit pitting), CGRMV and CNRMV were only detected in the fruit pitting sample T02, and their CP sequences clustered into a single group with known peach isolates from China. Also, for PBNSPaV, the number of PBNSPaV-specific reads in T02 was higher than in T01, and the number of contigs in T02 that mapped to the known PBNSPaV isolate WH-1 was significantly higher. The NSPaV isolates in T02 and T01 were clustered into different groups, confirming that they represent divergent sequence variants. Taken together, this result indicates that the fruit pitting observed in the T02 nectarine tree could be the result of interactions between PBNSPaV in family Closteroviridae, CGRMV and CNRMV in family Betaflexiviridae, and NSPaV in family Luteoviridae. Therefore, we can speculate that CGRMV and CNRMV might possibly serve as “helper” viruses in the involvement of specific variants of either NSPaV or PBNSPaV or both in the expression of fruit pitting symptoms. Synergisms have been reported to occur between the members of the genera Closteroviridae and Betaflexiviridae, and the p10 silencing suppressor from grapevine virus A in the family Betatextviridae enhances the infectivity of a Closterovirus, beet yellow virus, but the synergisms of several viral genera reported in this study are first discovered.

Again, samples T03, T04, and T05 showed different disease phenotypes; we detected PLMVd and NSPaV in the asymptomatic tree T03, PLMVd, NSPaV, APV2, and ACLSV in tree T04 with dimpled fruits, and only PLMVd in tree T05 with mottled leaves. In addition, of the five sampled nectarine trees, APV2 was only found in sample T04. This result was also confirmed in an additional five sampled trees with dimpled fruits using an RT-PCR screen. These results suggest that fruit dimpling at least may be related to the increase in APV2 and ACLSV titers. Previous studies have shown that, based on the amino acid sequences of the CP, ACLSV isolates have been classified into the types Z1/Z3 and Ta Tao5 from peach samples, and P205 and B6 from apple samples. In our study, seven ACLSV CP sequences from T01, T02, T03, and T04 were all closely related to isolate Z1. This result would seem to exclude a contribution of ACLSV to the dimpled fruit symptoms in T04. According to a previous report, APV2 infection could contribute to leaf symptoms in the GF305 peach indicator, but we have not found documented fruit dimpling symptoms. We have stated that NSPaV sequences in this study are more divergent, and that the T04-1 sequence, one of three RdRp gene sequences for NSPaV, was found to be a recombinant. Taken together, we infer that dimpled fruit symptoms also result from the synergistic effects of co-infection with NSPaV and APV2. As previously reported, some luteoviruses, such as groundnut rosette assistor virus, serve as “helper” viruses for the transmission of other viruses that cause disease.

PLMVd is the only viroid shared by all five nectarine trees that displayed three leaf disease phenotypes (asymptomatic, bleached, and chlorotic mottle) in this study. Some studies have shown that PLMVd infection can be associated with albinism (peach calico, PC) and green mosaic symptoms, and revealed a close association between the albino phenotype and variants containing an insertion of 12–14 nt, folding into a hairpin capped by a U-rich loop in the proposed PLMVd branched secondary structure. It is worth noting that we did not find this 12–14 nt insertion in this study. However, evolutionary analysis of PLMVd shows that the PLMVd sequences associated with symptomatic and asymptomatic trees do not all cluster together, and that there are intra-isolate divergent sequence variants present in every individual symptomatic tree. It has been suggested that divergent sequence variants of PLMVd co-infecting a single tree could contribute to host phenotype.

Biological data on the direct association of disease symptoms with the viruses identified in this study in fruit trees are scant in the literature. The main reasons for this are the fact that alternative herbaceous hosts are difficult to identify, and viral particles are very difficult to obtain in purified form for fruit tree viruses. Mixed viral communities that co-infect individual trees could increase viral genotypic complexity with implications for consequences to host pathology. However, increased application of NGS technology has resulted in the recent discovery of a large number of co-infecting viral communities in plants. Typically, this technology is used to identify candidate pathogens that may be associated with disease symptoms in plants. In the present study, several valuable field samples with disease phenotype differences combined with NGS data and genetic analyses supported the association between multiple co-infecting viruses and the disease symptoms observed. The data reported in this study will be important for further study of the biological and epidemiological features of virus/viroid interactions in plant hosts.
and 6-mer random primers in a 10 μL (also in Daliang) were also collected. The area of the three greenhouses is approximately 0.5–1.0 Chinese mu (1 mu = 666.7 square meters). The thirty-six symptomless and symptomatic tissue samples from greenhouses #1, #2, and #3 were collected and stored at −80 °C prior to use in the experiments. Of the 36 samples, the several different classes of symptoms were observed simultaneously on trees in greenhouses #1 and #2, which share the same environmental conditions as well as cultivation methods, pesticides, and fertilizers, and this allowed us to further study the etiological agent(s) associated with the different disease symptoms. Thus, five tissue samples from five different trees (two ‘Youtao 1233’ from greenhouse #1 and three ‘Zhongyou 4’ from greenhouse #2) with or without disease symptoms on leaves and fruits were screened for viruses by sequencing the small RNAs using NGS as shown in Table I and Fig. 1.

Viral communities identified by NGS in the small RNA libraries. Total RNA was extracted from each sample, and the small-RNA libraries were constructed using the NEB Multiplex Small RNA Library Prep kit (NEB, USA) following the manufacturer’s recommendations. Unique index codes were added to attribute the individual sequence reads to each sample library. The libraries were size-selected in 6% polyacrylamide gels prior to sequencing on an Illumina HiSeq 2500 SE50 instrument (Biomarker Technologies Co., Ltd), and paired-end reads were generated.

The raw read data in fastq format were initially processed using in-house perl scripts. In this step, clean reads were obtained by removing reads containing adapters, reads containing multiple Ns (unknown bases), and low quality reads from the raw data. The reads were trimmed and cleaned by removing sequences smaller than 18 nt or longer than 35 nt. The phred quality scores (Q20 and Q30), GC-content, and sequence duplication level were calculated for the clean data. All downstream analyses were conducted with high quality clean data.

The sequence reads were assembled de novo into contigs using Velvet Software with k-mer = 1737–38. The contigs obtained were subsequently annotated by BlastN and BlastX searches of the Genbank virus and viroid Reference Sequence Database. sRNA reads that mapped to individual viral genomes were also tabulated to identify candidate viruses present in the analyzed nectarine samples.

Virus and viroid detection with RT-PCR. Total nucleic acids were extracted from each sample using the RNAprep Pure Plant Kit (Tiangen Biotech (Beijing) Co., Ltd). Seventeen specific primer pairs (Table S2) were designed to amplify genomic regions corresponding to the CP, RdRp, or HSP70h genes from APV1, APV2, APV3, ACLSV, CGRMV, CNRMV, PBNPSaV, PeVD, NSPaV, PaLV, and GRGV, and also the complete genome sequence of PLMVd. The specific primer pairs to amplify nearly the complete genome of NSPaV are listed in Table S3. Reverse transcription (RT) was performed at 42 °C for 1 h using 1 μL of total RNA and 1 μL of oligo (dT) primer and 6-mer random primers in a 10 μL reaction volume containing Maloney murine leukemia virus (M-MLV) reverse transcriptase (Promega, Madison, WI, USA), according to the manufacturer’s protocol. Following RT, PCR assays were performed in 25 μL reaction volumes containing 1.5 μL of the RT reaction, 12.5 μL of 2X Taq Mix [Tiangen Biotech (Beijing) Co., Ltd ], 9.0 μL distilled water, and 1.0 μL (10 pmol) of the forward and reverse primers. The thermocycling conditions were as follows: an initial denaturation step of 5 min at 94 °C, followed by 35 cycles of 30 s at 94 °C, 30 s at 52°C–55 °C, and 90 s at 72 °C, with a final extension step of 10 min at 72 °C. RT-PCR products were purified using a PCR purification kit (AXygen), and the resulting DNA fragments were then cloned into the pMD18-T vector (Takara) for sequencing by the Sanger sequencing method. At least three clones of each amplified fragment were sequenced. Sequence reads were assembled using DNAMAN 6.0 (Lynnon Biosoft, Quebec, Canada).

Phylogenetic analyses of the identified viruses/viroids. We amplified complete genome sequences for PLMVd, nearly complete genome sequences for NSPaV, and full-length or partial sequences of the CP, RdRp, and HSP70h genes for the identified viruses from the five tree samples T01, T02, T03, T04, and T05 using RT-PCR with specific primers (Tables S2 and S3). The PCR products were cloned and sequenced, with at least three clones sequenced from each sample tree. All CP, RdRp, and HSP70h gene sequences from the identified viruses were aligned and the flanking sequences in the amplified fragments were removed to obtain the full or partial-length gene sequences for each virus. In total, 17 PLMVd genomes (336 to 338 bp), one nearly complete NSPaV genome (4.578 bp), 11 NSPaV RdRp genes (987 bp), seven ACLSV CP genes (582 bp), six PBNPSaV partial CP genes (963 bp) and six partial HSP70h genes (587 bp), six APV1 CP genes (1,206 bp), three APV2 partial CP genes (1,182 bp), six PaLV partial RdRp genes (981 bp) and three CP genes (647 bp), five PeVD genes (695 bp), three CGRMV CP genes (807 bp), and three CNRMV CP genes (804 bp) were used in the phylogenetic analyses. We download other known complete genome, CP, and RdRp gene sequences from GenBank (www.ncbi.nlm.nih.gov) to determine the phylogenetic relationships with known viruses/viroids. If there are too many virus/viroid sequences deposited in GenBank except for recently-identified PeVD, PaLV, and NSPaV sequences, after filtering partial sequences, we only retrieved complete genome sequences homologous to each virus and a few representative sequences (with different disease symptoms) from the NCBI nucleotide database to use in phylogenetic tree construction. We aligned the genome or gene sequences using the ClustalW multiple alignment program and calculated a sequence identity matrix using MEGA with the default parameters. The aligned sequences were checked for potential recombination events using RDP13,40. After sequence alignment, a phylogenetic tree was
constructed using the maximum-likelihood (ML) method as implemented in MEGA6, with confidence levels for the phylogroups estimated using 1,000 bootstrap replicates. The evolutionary history was inferred using the ML method based on the GTR + G model.

Data Availability

The sequences from the virus/viroid isolates using in the phylogenetic analyses in the present work have been deposited with NCBI (https://www.ncbi.nlm.nih.gov) under the following accession numbers: 17 PLMVd genomes: MH974826-MH974842, 1 NSPaV nearly complete genome: MK361454, 11 NSPaV partial RdRp gene sequences: MK361443-361453, 6 PBNSPaV partial CP gene sequences: MK361484-361489, 7 PBNSPaV partial HSP70h gene sequences: MK361490-361496, 7 ACLSV CP genes: MK361436-361442, 6 APV1 CP genes: MK361455-361460, 3 APV2 partial CP gene sequences: MK361461-361463, 3 CGRMV CP genes: MK361464-361466, 3 CNRMV CP genes: MK361467-361469, 3 PaLV partial CP gene sequences: MK361470-361472, 5 PeVd partial CP gene sequences: MK361473-361477.

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**Author Contributions**

Conceived and designed the experiments: M.L. Performed the experiments: Y.X., L.Y. and M.L. Analyzed the data: M.L., Y.X. Contributed reagents/materials/analysis tools: S.L., C.N. Contributed to the writing of the manuscript: M.L., Y.X. All authors reviewed the manuscript.

**Additional Information**

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