PREVALENCE AND DIVERSITY OF CANDIDATUS LIBERIBACTER SPECIES IN MAIN CITRUS GROWING AREAS OF MALAKAND IN NORTHWEST PAKISTAN

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ABSTRACT

Malakand Division in northwest Pakistan is known for delicious sweet orange, locally called Malta (Citrus sinensis). However, both quality and quantity of the citrus produce has seen a decline over the past few decades. Among production constraints, Huanglongbing (HLB) caused by phloem-limited alpha proteobacterium Candidatus Liberibacter asiaticus (CLas) - an endosymbiont of Asian Citrus Psyllid (Diaphorina citri), is of great economic significance. Therefore, we assessed HLB incidence through symptom-based surveys in major citrus growing areas of Khyber Pakhtunkhwa (KP), Northwest Pakistan. The presence or absence of CLas in the collected citrus samples was confirmed by PCR (Polymerase Chain Reaction) using three set of primers (LSS/LAS606, OI1/OI2C “16S rDNA” and MH353/MH354 “operon nusG-rplK”). An average disease incidence of 92% and severity of 34% were recorded from main production zones in district Swat, KP. Likewise, in district Lower Dir, HLB had disease incidence and severity of 64% and 24%, respectively. Molecular diagnostics using 16S rDNA-specific primer LSS/LAS606 and OI1/OI2C, yielded characteristic bands of 500 bp and 1160 bp, from all five pooled samples from Swat district, while only two samples of Dir tested positive for both primers. Using an alternative primer MH353/MH354 (nusG-rplK) showed positive amplification for four samples from Swat and two samples of Dir with an amplicon of 631 bp. To assess the molecular diversity of the CLas, Sanger sequencing was carried out based on 16S rDNA gene. Sequence analysis of 16S rDNA gene from all samples revealed maximum similarity with C. Liberibacter asiaticus. Among studied isolates, SwKjuPk-5 (MH374503) and SwMtaPk-2 (MH374500) isolates appeared to be more divergent and grouped in separate clad as compared to remaining isolates of Swat and Dir.

INTRODUCTION

Citrus is one of the major horticultural crops of Pakistan in terms of agricultural export. Kinnow or mandarin (Citrus reticulate) is grown in Punjab (Altaf et al., 2009) while Sweet orange (Citrus sinensis L.) in Khyber Pakhtunkhwa (KP) (GOP, 2016). In Pakistan, citrus
production per unit area is much lower (9.5 tons/hectare) as compared to developed countries (25 tons/hectare) (GOP, 2016). The lower productivity can be attributed to abiotic and biotic stresses that also play a role in reducing quality of the produce (Iftikhar et al., 2009; Razi et al., 2011). Among biotic stresses, diseases like huanglongbing (HLB) (Chohan et al., 2007), Citrus tristiza virus (Grimaldi and Catara, 1989) and Citrus canker (Catara et al., 1988) are of great economic significance for citrus production in Pakistan.

HLB is distributed worldwide and has become urgently a supreme challenge for citrus researchers and producing communities in recent past. HLB affects not only productivity, but also reduce fruit quality that often ends up premature fruit drop. In citrus, about 30-100% losses have been reported due to HLB (Aubert, 1992). The poor physiology of HLB affected citrus cultivars often leads to major losses to citrus industry of the world (Cevallos-Cevallos et al., 2012). This disease is also known as yellow shoot disease or citrus greening disease and was first reported by Reinking (1919) in southern part of China (Graca, 1991).

The causal agent of HLB is fastidious phloem restricted gram-negative bacterium *Candidatus Liberibacter* (family *Rhizobiaceae*, α-subdivision Proteobacteria) that is rigid rod shaped and its size ranges from 340–540 × 600–1,500 nm (Su, 1998). The HLB pathogen is further sub-classified into three species; *C. L. asiaticus*, *C. L. americanus*, and *C. L. africanus* (Da Graça and Korsten, 2004; Zhao, 1981). In nature, the pathogen is transmitted by Asian citrus psyllid (ACP) vectors, *Diaphorina citri* and *Triozoa erytreae* (Capoor et al., 1967; McClean and Oberholzer, 1965), and also by grafting (Lopes et al., 2009). Typical symptoms of HLB are foliar mottling and asymmetrical halves. Infected fruits have lopsided strong inversion, remain green forever with distorted seeds, taste salty and bitter (McClean, 1970).

HLB symptoms sometime can be confused with deficiencies of zinc, iron and magnesium and viral diseases like CTV disease (Jantasorn et al., 2007).

The disease has been new emerging threat to citrus industry of the world in general and that of Pakistan in particular (Batool et al., 2007). The incidence of HLB has been reported from the regions of Punjab, Jammu Kashmir and Rajasthan near Pakistan’s border (Ahlawat and Raychaudhuri, 1988). However, greater incidence of HLB (16-90%) was reported from orchards of KP and Punjab (Akhtar and Ahmad, 1999). It has been estimated that about 65% orchards are infected with the disease and its vector is present in Pakistan and India (Ghosh et al., 2015).

Persistent latency of the disease, nature of the pathogen, generic expression and influence of temperature on symptoms in an orchard are the factors which contribute to lack of disease management (Bové, 2006; Da Graça and Korsten, 2004; Halbert and Manjunath, 2004). In the absence of management strategies and resistant/tolerant cultivars, different Liberibacter species from Asia (*C. L. asiaticus*), Africa (*C. L. africanus*) and America (*C. L. americanus*) are evolving rapidly in the region. Therefore, we surveyed major citrus growing areas of Swat and Dir Districts to assess the current prevalence of HLB and understand the molecular systematics of the prevalent *Liberibacter* species.

**MATERIALS AND METHODS**

**Survey and sampling of HLB:** Symptom-based survey was carried out in the major citrus orchards of two districts of KP province including Lower Dir (Khanpur, Timargara, Khal, Rabat and Talash) and Swat (Jambil, Matta, Barikot, Marghozar and Kanju) from October 2016 to January 2017 as shown in Table 1 and figure 1. From each orchard of *Citrus senensis*, five different trees were selected randomly on X-pattern. From infected trees, showing characteristic symptoms of HLB, twenty-five leaves were collected for subsequent study and were stored at -20°C until further use. Disease incidence was calculated by using the following formula:

Percent Disease Incidence = \( \frac{\text{number of diseased plants}}{\text{total number of observed plants}} \times 100 \)

Disease severity index was calculated by using 0-4 rating scale; 0 = No symptom, 1 = Yellowing/ chlorosis symptoms on leaves (25%), 2 = Canopy symptoms and lopsided fruits (26-50%), 3 = Partially declined tree (51-75%), 4 = Sever dieback (>75%) (Akhtar and Ahmad, 1999). Percent disease severity was determined by using the equation given below:

\( \text{Percent Disease Severity} = \frac{\sum_{n} r}{4N} \times 100 \)

**DNA extraction:** DNA was extracted from the midrib of HLB infected citrus leaves by using CTAB-DNA extraction method of Doyle and Doyle (1987) with some modifications. Nano-drop was performed for concentration and purity of DNA using Li et al. (2006) method. Also, to check the concentration and purity of DNA, the samples were run on 1% agarose gel for 30 minutes.
Table 1. Prevalence and PCR base detection of *Candidatus* Liberibacter asiaticus in the citrus growing areas of Malakand in Northwest Pakistan.

| ID | Specific Origin | Disease index | Liberibacter Detection (PCR) |
|----|----------------|---------------|-------------------------------|
|    | District | Area | HLB Symptoms | Severity (%) | Incidence (%) | LSS/ LAS606 | OI1/OI2c | nusG-rplK |
| 1  | Swat  | Jambil | + | 45 | 100 | + | + | + |
| 2  | Matta | + | 35 | 80 | + | + | + |
| 3  | Swat  | Barikot | + | 25 | 80 | + | + | - |
| 4  | Marghozar | + | 40 | 100 | + | + | + |
| 5  | Kanju | + | 30 | 100 | + | + | + |
| 6  | Khanpur | + | 30 | 80 | + | + | + |
| 7  | Timargara | + | 20 | 80 | - | - | - |
| 8  | Dir   | Khal | + | 35 | 80 | + | + | + |
| 9  | Rabat | + | 20 | 40 | - | - | - |
| 10 | Talash | + | 15 | 40 | - | - | - |

**Figure 1.** Map of Major Citrus growing areas in Northwest Pakistan affected with Huanglongbing (HLB).

**PCR based detection:** The selected SSR pairs of primers were used in Polymerase Chain Reaction (PCR) for the region of 16S DNA and operon (Table 2). Amplifications were performed in an automated thermal cycler, a standardized amount of reagents concentrations (1.5 µl of primer for each pair, 7 µl of master mix and 0.5 µl Platinum Taq-polymerase) were used in PCR for 12.5 ul of reaction. PCR amplification protocol was 96°C for 9 min followed by 40 cycles of 96°C for 30 sec denaturation. While the annealing temperature of primers; LSS/Las606, OI1/OI2c and MH0353/MH0354, were 53.1, 63.8 and 60.7°C respectively for one minute. The initial and final extension was performed at 72°C for 30 sec and 7 min respectively (Li et al., 2006). The
amplified products were visualized on 1.5% agarose gel, stained with ethidium bromide. **Candidatus Liberibacter sequencing and analysis for genetic diversity:** A second round of PCR was carried out for PCR positive samples and the amplified products were sequenced using Sanger Sequencer as recommended by manufacturer (Bio Product). The obtained 16S rDNA sequences were analyzed by BioEdit v7.2 (Hall, 1999). These sequences were compared with sequences retrieved from National Center for Biotechnology Information (NCBI), using Basic Local Alignment Search Tool (BLAST) algorithm (Table 3). The sequences were uploaded to MEGA-7 and aligned using Clustal W (Thompson et al., 2003). Furthermore, the phylogenetic analysis and tree was constructed by neighbor joining method using MEGA-7 software and Randomized Axelerated Maximum Likelihood (RAxML) (Kong et al., 2000).

Table 2. Sets of primers used for detection of HLB in samples collected from district Lower Dir and Swat.

| Primer Name | Region | Sequence (5'-3') | Reference | Product Size |
|-------------|--------|-----------------|-----------|--------------|
| O1I         | 16S rDNA | GCGCGTATGCAAGAGCGGCA | (Jagoueix et al., 1997) | 1160 bp |
| O12c        | 16S rDNA | GCCCTGCAGCTTCGCAACCCAT | | |
| LSS         | 16S rDNA | ACCCAACATCTAGTGAAAAACC | (Fujikawa and Iwanami, 2012) | 500 bp |
| Las606      | 16S rDNA | GGAGAGGTGAGTGGAATTCCA | | |
| NUSG-RPLK (MH0353) | nusG-rplK (Operon) | GTGTCTCTGATGTTCCGTTTCTTTTGA | (Hoy et al., 2001) | 631 bp |
| NUSG-RPLK (MH0354) | nusG-rplK (Operon) | GAACCTTCCACCACGGCATGCCCTTTCA | | |

Table 3. Various retrieved isolates Liberibacter sequences used in the study and the sequences from Swat and Lower Dir region, submitted to NCBI GenBank.

| NO. | Isolates | Isolation source | Origin | Accession No. |
|-----|----------|------------------|--------|---------------|
| 1   | Clas A4  | Mandari citrus   | China : Guangdong | CP010804 |
| 2   | Clas JXGC | Citrus          | China Jiangxi          | CP019958 |
| 3   | Clas Gxpsy | Psyllid         | China              | CP004005 |
| 4   | Clas Psy62 | nd (not determined) | USA | CP001677 |
| 5   | Clas Ishi1 | Nd               | Japan              | AP014595 |
| 6   | Clso ZC1  | Zebra chip-infected potato | USA: Dalhart, Texas | CP002371 |
| 7   | Claf PTSAPSY | Psyllid        | USA | CP004021 |
| 8   | Clam Soa Paulo | Nd             | Brazil             | EU754742 |
| 9   | Lib Cres BT0 | Babaco pedunkle sap | USA | CP010522 |
| 10  | Lib Cres BT1 | Nd              | USA | CP003789 |
| 11* | SwJmbPk-1 | Citrus sinensis leaves | Pakistan: KP, Swat | MH374499 |
| 12* | SwMtaPk-2 | Citrus sinensis leaves | Pakistan: KP, Swat | MH374500 |
| 13* | SwBktPk-3 | Citrus sinensis leaves | Pakistan: KP, Swat | MH374501 |
| 14* | SwMzrPk-4 | Citrus sinensis leaves | Pakistan: KP, Swat | MH374502 |
| 15* | SwKjuPk-5 | Citrus sinensis leaves | Pakistan: KP, Swat | MH374593 |
| 16* | DirkKhpPk-6 | Citrus sinensis leaves | Pakistan: KP, Dir | MH374497 |
| 17* | DirkKhpPk-7 | Citrus sinensis leaves | Pakistan: KP, Dir | MH374498 |

*This study, nd=not determined

RESULTS

**Disease incidence:** The disease incidence of HLB recorded during survey in various orchards is given in figure 2a. Disease incidence was recorded in a range of 40 to 100%. In district Lower Dir, the highest disease incidence was recorded in Khanpur (80%) and Timargara (80%). Moreover, the lowest disease incidence was recorded in Talash (40%). In orchards of Swat, the highest
incidence was recorded in Jambil (100%) and Marghozar (100%) while the lowest was recorded in Barikot (80%). The overall mean incidence of 92% was recorded in Swat and that of 64% in Lower Dir.

**Disease severity:** The disease severity in KP ranged from 15-45% (Figure 2b). In district Swat, the maximum disease severity was recorded in Jambil (45%) followed by Marghozar (40%) while the minimum disease severity was recorded in Barikot (25%). Similarly, in district Lower Dir, the maximum disease severity was recorded in Khal (35%) followed by Khanpur (25%) whereas the minimum disease severity was observed in Talash (15%). The disease severity was higher in district Swat (35%) as compared to the district Lower Dir (24%).

**Figure 2a.** Disease incidence of HLB in district Lower Dir and Swat of Khyber Pakhtunkhwa, Pakistan.

**Figure 2b.** Disease severity of HLB in district Lower Dir and Swat of Khyber Pakhtunkhwa, Pakistan.

**PCR amplified product:** Polymerase chain reaction (PCR) used primer LSS/Las606 (16S rDNA), amplified a band of 500 bp for all samples of Swat region; Jambil (S1), Matta (S2), Barikot (S3), Marghozar (S4) and Kanju (S5). Similarly, samples collected from district Lower Dir, the same primer amplified a band of 500 bp for Khanpur (D1) and Khal (D3). However, samples collected from Timargara (D2), Rabat (D4) and Talash (D5) were tested negative for *Candidatus* Liberibacter species while there was no amplification for negative control (Figure 3a).
Primer OI1/ OI2c (16r DNA) amplified a band of 1160 bp for the samples of district Swat; Jambil (S1), Matta (S2), Barikot (S3), Marghozar (S4) and Kanju (S5). Similarly, the same primer amplified two samples of Lower Dir; Khanpur (D1) and Khal (D3) whereas, other three symptoms based samples from Timargara (D2), Rabat (D4) and Talash (D5) were found negative in PCR (Figure 3b).

Primers MH353/ MH354 (631bp) amplified operon (nusG-rplK) region of Candidatus Liberibacter spp in the samples of district Lower Dir and Swat. Primer (MH353/ MH354) amplified four samples from district Swat i.e. Jambil (S1), Matta (S2), Marghozar (S4) and Kanju (S5). However, samples from Barikot indicated negative response for the primer. Likewise, in district Lower Dir: Khanpur (D1) and Khal (D3) samples were observed positive for HLB while other three samples from Timargara (D2), Rabat (D4) and Talash (D5), were negative for Candidatus Liberibacter spp. detection (Figure 3c).

Figure 3a. Amplification of a band of 500 bp for Candidatus Liberibacter asiaticus using primer LSS/LAS606 (16S rDNA) from samples collected from various location of district (A) Swat and (B) Lower Dir. Amplicons were separated on 1.5% agarose gel. M: 100bp plus DNA ladder, NC: negative control.

Figure 3b. Amplification of a band 1160 bp for CLas, using primer pair OI1/ OI2c from samples collected from various location of district (A) Swat and (B) Lower Dir. Amplicons were separated on 1.5% agarose gel. M: 100 bp plus DNA ladder, NC: negative control.
Sequencing of the causal agent of HLB: To further investigate the genetic diversity of *Candidatus* Liberibacter asiaticus on 16S rDNA gene, PCR positive samples were sequenced and then analyzed. During NCBI BLAST, the obtained sequences from GenBank showed 85-99% similarity with our sequences (Table 3). In this study, DirKhpPk-6 (MH374497) and DirKhlPk-7 (MH374498) isolates had close similarities with retrieved sequences from GenBank (Accession number CP001677, CP010804 and CP019958) whereas; isolate SwBktPk-3 (MH374501) had resemblance with Clas Gxpsy (CP004005). Likewise, isolates collected from Swat; SwjmbPk-1(MH374499) and SwMzrPk-4 (MH374502) were also similar to Clas Ishi-1 (AP014595). Among Swat and Lower Dir isolates; SwMtaPk-2 (MH374500) and SwKjuPk-5 (MH374503) showed diversity and produced an isolated cluster. The phylogenetic tree indicates that Clam Soa Paulo (EU754742), CLaf PTSAPSY (CP004021), Clso ZC1 (CP0002371) and other isolates Lib cres BT0 (CP010522) and Lib cres BT1 (CP003789) formed a separate clade from Liberibacter asiaticus isolates. In precise, multiple sequence alignment and phylogenetic analysis, based on 16S rDNA revealed that KP-isolates reported in present study are mostly similar to *Candidatus* Liberibacter asiaticus having accession numbers (CP010804, CP019958, CP004005, CP001677 and AP014595), compared to others C. Liberibacter spp. The homology of isolates; DirKhpPk-6 (MH374497), DirKhlPk-7 (MH374498), SwBktPk-3 (MH374501), SwjmbPk-1(MH374499) and SwMzrPk-4 (MH374502) were about 97-99% with Clas sequences in NCBI database. However, isolates SwMtaPk-2 (MH374500) and SwKjuPk-5 (MH374503) showed 85-90% homology with the Liberibacter species retrieved from NCBI database (Figure 4).

DISCUSSION
Citrus has been one of the major export products of Pakistan (GOP, 2016), however, the citrus orchards in Pakistan are increasingly coming under severe pressure from HLB (Hussain et al., 2019; Khan et al., 2014; Qasim et al., 2018; Razi et al., 2014). Among several biotic stresses, HLB is one of the major contributing factors toward quantitative and qualitative losses of fruit in the region. HLB is one of the diseases that not only reduce fruit yield but also alters fruit quality i.e. bad flavor, reduced sugar content, high acidity, distorted fruit shape etc (Spann and Danyluk, 2010). The present study was, therefore, designed to assess the prevalence of HLB and diversity of *Candidatus* Liberibacter asiaticus in the Malakand Division of Northwest Pakistan. The study showed that district Swat has been severely affected by HLB disease. The maximum disease incidence and severity were observed in Swat as...
compared to district Lower Dir of the country. HLB affects all varieties and cultivars of citrus like mandarin, sweet orange, grapefruit and tangelo being the most susceptible while pummel, lime and trifoliate orange are the least susceptible (Knapp et al., 2004). Typical HLB symptoms as described by Ahlawat (1997) were noticed in the orchards. It has been reported that most of the orchards (lemon, grapefruit, orange and mandarin) of Punjab and KP showed HLB symptoms (Catara et al., 1991; Munir et al., 2019). The causal agent could even be detected in the asymptomatic citrus trees (Ghosh et al., 2003). Accordingly, most growers start grafting only asymptomatic trees to eradicate the pathogen. While the asymptomatic trees still have the Clas pathogen which hampers the sanitation and eradication practices. There is a delicate balance among temperature, host and the pathogen which provides favorable condition to HLB. The other main factors that contribute to the disease are large numbers of citrus susceptible varieties and host range in the region (Razi et al., 2014). In Malakand Division, it may be the possible reason that nurseries are the initial source of inoculum, since most of the growing stocks in nursery are usually infected with HLB disease (Akhtar and Ahmad, 1999; Munir et al., 2018).

Figure 4. Phylogenetic analysis based on 16S rDNA showing variation for HLB isolates of Swat and Lower Dir of Khyber Pakhtunkhwa-Pakistan. The Bold isolates represent “Liberibacter asiaticus sp” obtained from Swat and Lower Dir, KP, Pakistan.

The disease incidence and severity were very high in Swat. The possible reason behind this alarming situation is the abundant population of vectors in the region which easily multiply the pathogen in their bodies. Other possible reasons may be susceptible varieties and too-old citrus trees in the region which enhanced the possibility of the pathogen’s attack and gradual climatic changes in the region are supportive factors. All citrus
varieties and cultivars were found susceptible to the disease (Chung and Bransky, 2005). Although some tolerance has been reported in some species of citrus (Rawat et al., 2017) as commercially grown citruses are generally susceptible to the disease. The present findings are similar to those of Akhtar and Ahmad (1999) who found the incidence of HLB between 16-66% and 90% in Punjab and KP, respectively. For identification and elucidating molecular diversity, three primers were used for detection of HLB including LSS/LAS606, OI1/OI2C and MH353/MH354, which amplified 16S rDNA and operon (nusG-rplK) region. All samples from district Swat were positive for two pairs of primers (LSS/LAS606 and OI1/OI2C) and two out of five samples from Lower Dir were amplified by primers (LSS/LAS606 and OI1/OI2C). However, some symptom based collections were PCR negative during the study, that require further investigations if other divergent species are present in the study area (Teixeira et al., 2008). Furthermore, the citrus greening symptoms are sometimes confused with other diseases and nutritional deficiencies (Bové, 2006; Munir et al., 2019). Additionally, an uneven distribution and overlapping of nutrient deficiencies in field causes problem to differentiate the disease, hence molecular tools are useful to distinguish them. The majority of asymptomatic trees may have the pathogen, but the titer of pathogen was not sufficient for detection. Further investigations using new molecular tools such as LAMP PCR, may further shed light on the prevalence and severity of the disease (Choi et al., 2018).

The results of the present study are in agreement with those of Kunta et al. (2014) as it is difficult to distinguish between HLB symptoms and nutrient deficiencies (Munir et al., 2018). MH353/MH354 primer showed less sensitivity for CLas pathogen. It may be due to the presence of inhibitors in the citrus leaf tissues that can affect PCR result (Hocquellet et al., 1999; Li et al., 2006) or the presence of divergent species. Primers LSS/LAS606, OI1/OI2C and MH353/MH354 detected “Candidatus Liberibacter spp.” from various samples of citrus. It was observed that CLas is responsible for HLB in citrus (Ruangwong and Akarapisan, 2006; Yaqub et al., 2017) and a pair (O11/O12C) of primers was used for the detection of “CLas” (Munir et al., 2019). The primer amplified a product of 1160 bp but no amplicon was detected for healthy samples. Study suggests that LSS/LAS606 is a robust primer for HLB detection which has high sensitivity at temperature ranges from 50-64°C. Phylogenetic relationship among various species of Liberibacter was elucidated from other countries on the basis of 16S rDNA. In the current study, the species isolated from Swat and Lower Dir isolates were closely related to Candidatus Liberibacter asiaticus (CLas) than other species of Liberibacter (CLam, CLaf and CLso) (Figure 4). Our study is in line with the findigns of Bastianel et al. (2005) and Subandiyah et al. (2000) who differentiated “CLas” from other species of “C. Liberibacter” on the basis of 16S rDNA sequences and stated that isolate from Thailand (99.4-100%), Nepal (100%) and India (98.8%) have closer identity with CLas. These results have similarity with those reported by Ghosh et al. (2003) who confirmed that Indian isolates had 99.9% similarity with those of China and Thailand. The study also revealed that two isolates of HLB (SwMtaPk-2 and SwKjuPk-5) from Swat region have minor divergence in genetic makeup of “CLas” reported from other countries. A geographical distribution and maximum use of pesticides in the region may be a possible reason for mutation in gene. Lack of quarantine inspection for imported rootstock (HLB infected) also promotes genetic variation.

CONCLUSION
The prevalence of HLB disease (incidence and severity) was high in district Swat as compared to Lower Dir. Primers used in the study confirmed the presence of HLB (CLas) in the region. Also, LSS/Las606 and OI1/OI2c primers were more specific for detection of the pathogen. Phylogenetic relationship indicates that KP-Pakistan isolates are relatively more similar to C. L. asiaticus as compared to C. L. africanus and C. L. americanus. Pakistan is one of the largest producers of Kinnow with unique taste throughout the world so, it is required to control HLB disease to promote export of the fruit to rest of the world. Molecular tools are requirement of time to identify the infected root stock in nurseries, and quarantine department to prevent propagation of the disease. Detailed analysis of HLB is needed to identify other emerging strains and species. Therefore, this study will be helpful to develop management strategies for HLB.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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