

Detection of "*Candidatus Liberibacter asiaticus*" using conventional PCR and nested-PCR and identification of primer sensitivity through Q-PCR in HLB infected mandarin orange and acid lime plants in Tamil Nadu.

ABSTRACT

One of the most destructive citrus diseases in the world is called citrus Huanglongbing (HLB). For effective HLB associated "*Candidatus Liberibacter spp.*" prevention, sensitive and precise diagnostics are essential. *Candidatus Liberibacter asiaticus* (Las), *Candidatus Liberibacter africanus* (Laf), and *Candidatus Liberibacter americanus* are among the "*Candidatus Liberibacter spp.*" that infect citrus. The most common of these is the species known as Las. To assess the impact of citrus greening disease in the citrus orchard, leaf samples exhibiting greening symptoms were picked up from 189 mandarin orange trees from nine localities in four districts and 202 acid lime trees from nine localities in six districts of Tamil Nadu where citrus is cultivated in a larger area. A conventional singleplex polymerase chain reaction (PCR) Nested PCR and Q-PCR (Quantitative PCR) was performed on the extracted DNA samples to detect *Ca. Liberibacter asiaticus*. Among all the mandarin orange growing localities from Kanalkadu and Thandikudi in Dindigul district and among the acid lime fields, the samples from Sankarankoil from Tirunelveli district and Kallar from Nilgiris district showed consistent amplification of 1160 base pairs fragment by using specific primers OI1 and OI2C and also 703 base pairs fragment with gene-specific primers A2 and J5 targeting beta-operon (rpiKAJL-rpoBC) gene which is a ribosomal protein gene of *Ca. Liberibacter asiaticus*. The nested PCR method significantly increased the sensitivity to identify Las up to 10 times and 100 times, respectively, compared to qPCR and conventional PCR in this study. A set of nested PCR primer pairs were examined to diagnose Las. Nested PCR was used to analyse 14 samples from 2 different citrus cultivars in 2 different districts. The findings indicate that all the samples tested positive for HLB; the samples of blotched and chlorotic leaves from the acid lime field at Sankarankoil (thirunelveli district) and mandarin orange field at kanlakaadu (Dindigul district) had the highest positive detection rate (100%). The outcomes show that the nested PCR primer pairs are capable of detecting Las in a variety of symptomatic tissues, citrus cultivars, and geographical locations. The set of nested PCR primers used in the current work will be a very helpful addition to the existing methods for Las detection. The nested PCR primers sensitivity in terms of amplification were compared with conventional PCR primers amplification through Q-PCR, and amplification efficiency (Eff) was calculated, which showed 91.21 % Eff for nested PCR second set of primers.

1. INTRODUCTION

citrus greening is one of the most devastating diseases to threaten the citrus sector in Asia, Africa, and America is citrus Huanglongbing (HLB), [15]. Citrus trees that have been severely infected by HLB exhibit yellow shoots, foliar blotchy mottle that may resemble the symptoms of zinc deficiency, vein corking that may resemble the symptoms of Citrus Tristeza virus infection, poor blooming, and stunting [6]. Citrus trees that have HLB infection are chronically foliated sparsely, have significant twig or limb die-back, and typically die within three to five years [7, 18]. Citrus with HLB infection has a 30% to 100% reduction in yield, and the degraded fruit quality [15, 6, 28]. HLB is caused by "*Candidatus Liberibacter spp.*" gram-negative, unculturable and phloem-restricted organisms that fall within the Proteobacteria division. *Candidatus Liberibacter asiaticus* (Las), *Candidatus Liberibacter africanus* (Laf), and *Candidatus Liberibacter americanus* (Lam) are *Candidatus Liberibacter spp.* linked with HLB [7]. The most common species, Las, is also the one that causes the most of the rising economic losses [22, 35]. Budding, dodder, grafting, and psyllid vectors were all used to spread HLB. HLB is spread very fast that the spread distance might reach 193 km (120 miles) each year [15, 16]. Regrettably, there is currently no cure for this illness that can be used to control it, and there are no cultivars that are resistant to this pathogen. The most common forms of HLB management are eradicating affected trees and controlling the insect vector. As a result, sensitive and precise diagnosis is a requirement for studying and managing HLB. Koch's hypotheses on HLB were not tested since it was impossible to culture the HLB bacteria [27]. Several molecular detection methods based on PCR are currently in use to detect HLB-associated bacteria, including conventional PCR [30, 31], SSR [8], droplet digital PCR [36], LAMP [26,24], immune capture-PCR [11], qPCR [18,1], and nested PCR [4]. Nested PCR has been shown to have higher sensitivity than other molecular detection methods for diagnosing disorders like HLB [21, 9], as the first round of PCR products are diluted and used as the template for the second round of amplification. Hence, in this investigation, nested PCR was employed. The F1/B1 and F3/B3 results of Las isolates from various geographic locations revealed positive, indicating that the sequence locus was preserved and shared by these Las isolates (Fig. 2A, 2B, 2C and 2D). Thus, the selected primers are substantially conserved and species-specific.

2. MATERIALS AND METHODS

In Tamil Nadu, two citrus species, Citrus (mandarin orange)-(experiment 1) and Citrus (acid lime)-(experiment 2), were surveyed and tested for the presence of *Ca. Liberibacter asiaticus*.

I. Experiment 1

Survey for Huanglongbing in Tamil Nadu (Mandarin orange)

From March 2018 to March 2021, systematic surveys were carried out in key mandarin orange-growing regions throughout Tamil Nadu, including Yercaud (Salem district), Kolli hills (Namakkal district), Conoor, Katteri, Ooty (Nilgiris district), Kanalkadu, Patlankadu, Thandikudi, and Thadiyankudisai (Dindigul

district). To prevent DNA deterioration, samples thought to have HLB infection were gathered and put right away in a refrigerated box. The gathered samples were taken to the lab and either processed right away for DNA extraction or put into cold storage at -80 °C for later use.

II. Experiment II

Survey for Huanglongbing in Tamil Nadu (Acid Lime)

comprehensive surveys were carried out Between March 2018 and March 2021 in key acid lime-growing regions in Tamil Nadu, including Pollachi (Coimbatore district), Sankarankoil (Thirunelveli district), Kolli Hills (Namakkal district), Thadiyankudisai, Ayyampalayam (Dindigul district), Coonoor, Katteri, and Ooty (Nilgiris district). Acid lime symptom manifestations were observed and noted during the survey. In order to prevent DNA deterioration, samples that were thought to have HLB infection were gathered, stored right away in a cold box, and either processed right away for DNA extraction or preserved at 80 °C for later use. Symptom manifestations on various citrus species were also seen and noted during the survey while carrying out both studies. Latitude, Longitude, and Altitude of the specific orchards and fields, together with the percentage disease incidence (PDI), were recorded. The percentage of disease incidence was calculated by the formula given by Ahmad K B. 2008 (3).

$$\% \text{ disease incidence} = \frac{\text{Total infected citrus trees}}{\text{Total number of trees evaluated}} \times 100$$

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DNA extraction

Each sample's leaf midribs were individually used to extract total DNA using the CTAB procedure suggested by Doyle *et al.*, 1991 (12). One gram of mandarin orange midribs were separated using a clean surgical blade. The midribs were then broken up into little pieces, macerated in a pestle and mortar using liquid nitrogen, and put into an Eppendorf tube. The macerated material was mixed with two millilitres of CTAB buffer and heated to 65 °C in a water bath for 20 minutes (Scientech, India). The samples were centrifuged at 10,000 RPM for 15 minutes after being inverted for a brief period of time every 10 minutes. A fresh Eppendorf tube was used to collect the supernatant. Equal quantities of 24:1 chloroform:isoamyl alcohol were then added to that and thoroughly blended by vortexing. The sample was then centrifuged for ten minutes at 10,000 RPM. A fresh Eppendorf tube was used to transfer the aqueous phase into, and the process was repeated with an equal volume of 24:1 chloroform:isoamyl alcohol. A new tube was used to collect the aqueous phase, and isopropanol was added in an equivalent volume. After three hours at -20°C, this tube was centrifuged at 10,000 RPM for 20 minutes to precipitate the DNA.

After washing the DNA pellet with 70% ethanol and centrifuging it at 5000 RPM for five minutes, the supernatant was discarded. The DNA pellet was then air-dried after being washed with ethanol, and the supernatant was discarded. The DNA pellet was finally dissolved in 50µl of sterile water and kept at -20°C. A NanoDrop Spectrophotometer was used to measure the OD values at 260 nm and 280 nm in order to evaluate the quantity and quality of the isolated DNA (NanoDrop, TNAU). For PCR analysis, DNA samples with OD values between 1.8 and 2.0 were gathered.

PCR amplification: In order to target the bacterial 16s r DNA gene in the collected samples of mandarin orange and acid lime, amplification was carried out in a Thermal cycler (Eppendorf) using the universal primers FD1 and RP2. The most specific region of the CLas genome, partial 16S rDNA, was the target of PCR using the primer pair OI1F/OI2CR (2), with an amplicon size of 1,100 base pairs, and the beta-operon (rpiKAJL-rpoBC) of the ribosomal protein gene as the target, with an amplicon size of 703 base pairs, using the primer pair A2/J5 (Fig.A and B). The reaction mixture (25 µl) contained 0.3 µl of Taq polymerase (5 units/l, GeneiTM), 0.5 l of dNTPs (10 mM), 2.5 µl of 10x buffers, 2.0 µl of forward and reverse primers (10 mM), 5 µl of DNA template (100-200 ng/l), and the remaining volume was made up of nuclease-free water. The following were the thermal cycle conditions set for the primer combination OI1F/OI2CR: Step 1: a cycle at 95 degrees Celsius for two minutes, followed by 35 cycles of 95 degrees Celsius for 40 seconds in step 2, 60 degrees Celsius in step 3, and 72 degrees Celsius in step 4, and a 72 degrees extension for 10 minutes [2]. The following are the PCR settings for the A2/J5 primer pair: Step 1: 94 °C for 3 minutes, Step 2: 94 °C for 1 minute, Step 3: 58.5 °C for 1 minute, Step 4: 72 °C for 1 minute, and Step 5: 72 °C for 10 minutes of extension. In 1.2% agarose gel with ethidium bromide in 1x Tris-acetate EDTA buffer, the amplification product was analysed. With the aid of a gel documentation system, the amplicons were seen. The PCR amplified DNA fragments were cleaned in the Eurofins lab using a DNA clean-up kit to get rid of any remaining primers, polymerase, and buffer (Bangalore, India).

NESTED PCR AMPLIFICATION

DNA preparation: Among the HLB positive samples from various districts, the kanalkadu and Thandikudi orchard mandarin orange leaf samples from dindigul and Sankarankoil acid lime field from thirunelveli district were chosen by observing the maximum PDI when compared to the remaining fields. The mandarin orange and acid lime leaves were cleaned under running water from the faucet and dried with paper towels. Midribs were removed. The CTAB method was then used to extract DNA from each sample after it had been pulverised to a 100 mg size in liquid nitrogen [24]. TE buffer (50 µl) was used to dissolve the isolated DNA. Using a NanoDrop (NanoDrop Technologies Inc.), DNA quality and concentration were examined [24]. The HLB pathogen was

identified using the Las specific primers targeting OMP (outer membrane protein) amplified fragments of 1438 bp for the first set of primers and 443 bp for the second set of primers. (Fig. C,D,E and F)

PCR conditions: The 2x Easy PCR master mix (smart prime) was used to generate the nested PCR mixture (20 µl), and the following settings were used for the amplification process: 94°C for 5 min, then 25 cycles of 30 seconds at 94°C, 30 seconds at 58°C, and 70 seconds at 72°C for the first round of PCR, and 35 cycles of 30 seconds at 94°C, 62°C, and 30 seconds at 72°C for the second round of PCR [20]. All of the findings are in line with the theory that nested PCR was more sensitive and useful for detecting Las at very low titers [47]. Compared to traditional RT-PCR, nested RT-PCR has better sensitivity and specificity. Nested PCR [30] uses two rounds of PCR, which increases the sensitivity. For bacteria to maintain their regular structure and function, the outer membrane protein (OMP) is essential. OMPs participate in interactions between plants and pathogenic microorganisms as well as exchanges with the outside environment. The nucleotide sequences of OMPs from Las showed high similarity and high species specificity (99%), and their three-dimensional architectures were largely conserved [21, 28, 6]. In HLB bacterial assays for detection, OMPs have been employed as target genes [28]. OMPs, however, were shown to vary greatly among geographical isolates and were not appropriate for identifying Las, according to earlier literature [2]. OMPs were frequently employed to make antigens and to analyse the variance between various geographic isolates. In this research, The outer membrane protein (OMP) assembly component is encoded by the conserved section of BamA, which is found in only one copy in the Las genome. This new HLB diagnosis approach uses nested PCR to amplify this area. The technique significantly increased the sensitivity in the detection of Las by up to 10 times when compared to qPCR and 100 times when compared to conventional PCR.

Identifying the sensitivity of Q-PCR for Conventional PCR, First primer set of Nested PCR, Second set of Nested PCR.

The primary objective was to quantify the DNA titre in the sample which DNA quantity was unable to identify through conventional PCR. Usually, the determination of primer sensitivity was based on different templates or different concentrations of the same template. Firstly, **three** suspected samples were amplified by conventional PCR primer (OI1/OI2) and a single sample of mandarin orange and acid lime were run through nested PCR primer pairs (F1/B1 and F3/B3). Then we detected the concentration of the conventional PCR positive product and nested PCR positive product by a NanoDrop (Bio Rad)), and adjusted to 100 ng/µl as the dilution template. The PCR product was serially diluted in a range of 1, 10¹, 10², 10³, 10⁴ and 10⁵, which represented 10⁵ to 1 pg/µl Las DNA, and served as the templates to evaluate the sensitivity among nested PCR, conventional PCR and qPCR. The PCR product from 1A line 8

and 9 was diluted as the template for conventional PCR, and the PCR product from Fig B line 3 and 6 was diluted as the templates for the nested PCR and qPCR. To evaluate the amplification efficiency of qPCR, Las DNA serially diluted in a range of 1, 10, 10², 10³, 10⁴ and 10⁵ served as the template. (Fig E, F, G, H and I). The DNA was adjusted to the same concentration for conducting the Q-PCR experiment and both sets of Nested PCR, So that the titre can be able to be quantified properly in Q-PCR. The DNA standard consisted of a 1318 bp amplicon produced by the F1/B1 primer pair (forward GGTTATGCTGCCGTAAAGTGTC and reverse B1: AACCAGCCCTTTCAGGAACAAG) that was purified by chloroform extraction followed by isopropanol precipitation, and quantified by averaging three replicate A260 absorbance determinations conducted on two spectrophotometers. A second 443 bp amplicon was produced by pairing of another primers F3/B3 (forward F3: TCTGAGGGTGAGCGTAAACAACACTG and reverse B3: TTGGGAAATAGAATGGCTGCTGAAT). The primer pair combinations of F1/B1 and the nested F3/B3 thus allowed the production of two different sized amplicons (1318 and 447 bp, respectively) using the same DNA standard dilution series.(41), The qPCR mixture (20 µl) was prepared using qPCR syber green (Takara), and amplification was proceeded using the following parameters: 94°C for 30 s and followed by 40 cycles at 94°C for 5 s and 60°C for 30 s, and followed by a melt curve (60°C to 90°C, 0.3°Cs-1). Whole Run was completed with a melting curve analysis to confirm the specificity of amplification and lack of primer dimers. Ct values were determined by the Roche software using a fluorescence threshold manually set to 0.0160 and exported into a MS Excel workbook (Microsoft office) for analysis.

The mathematics of Quantitative PCR was described by the protocol given by Rutledge 2003. Irrespective of many runs, Here a single run consisting of original DNA concentration and 10¹ to 10⁵ dilutions were prepared for conventional PCR, Nested PCR first set of primers and Nested PCR Second set of primers. The calculation was made with r^2 (Regression coefficient) Es (ES slope derived estimate of amplification efficiency and Nt (Number of amplicon molecules at fluorescent threshold). After the PCR product was adjusted to 100 ng/µl as the dilution template. Q-PCR at various dilutions were prepared for the conventional PCR amplified product and Ct value of first set of readings were noted and standard curve were constructed. The Ct value readings for conventional PCR is 30.96, 28.98, 26.54, 24.24, 21.28 and 18.84 which occurred through 1 to 10⁵ dilutions. The Q-PCR at various dilutions were prepared and Ct value of first set of nested PCR amplified PCR product were adjusted to 100ng and QPCR were performed with specific primers and Ct values were noted and standard curve were constructed. The Ct value readings for first set of nested PCR is 29.26, 26.21, 22.88, 19.42, 15.22 and 11.24 which is 1 to 10⁵ dilutions. After that with the same quantity of PCR product of first set of nested PCR which was used as a template DNA for second set of Nested PCR were adjusted to 100ng through Nanodrop and the second set of Nested PCR was performed, Later Q-PCR was performed, The Ct value readings for second set

of nested PCR is 26.54, 23.98, 21.00, 17.21, 13.44 and 8.98. All the samples after running the PCR were run in gel electrophoresis and purified through gel elution kit (Takara) and adjusted the DNA in nanodrop and quantified through Q-PCR (Roche instrument) at various dilutions. The primary objective of purification is to avoid the remaining the other components except DNA.

2.1 Statistical analysis

The statistical analysis was performed using OPSTAT and DMRT (Duncan Multiple Range Test) and analysed through SPSS software package. Q-PCR calculations such as Q-PCR calculations such r^2 , Eff and Nt, Mt, log factors, Average Ct and standard curves were done through MS-Excel.

3. RESULTS

3.1 Huanglongbing disease survey

The survey was carried out during different seasons, including winter (December to February), summer (March to May), monsoon season (June to September), and a post-monsoon period (October and November). The samples collected in the summer displayed more symptoms, and the concentration of CLas bacteria was higher during this time, as noted by visual observation as well as by the results of PCR and Q-PCR. The leaf samples were obtained by observing a variety of symptoms, including yellowing of the leaves and branches, the emergence of leaves that resemble rabbit ears, uneven mottling on the leaf lamina (blotchy mottle shoot with yellow areas), and symptoms similar to vitamin shortages (a regular pattern of yellowing or vein yellowing or clearing on leaf lamina).

3.2 Detection of CLas in mandarin orange:

The CLAs in acid lime and mandarin orange were found through the course of two different tests. One task was to identify samples from mandarin oranges, and the other involved acid lime samples. All the samples of mandarin oranges collected that were thought to be infected with putative CLAs isolates initially produced an amplicon of about 1500 bp DNA fragment after a PCR experiment using the universal primers FD1 and RP2 targeted the 16s r RNA gene. These DNA fragments were then confirmed as uncultured CLAs by sequencing (Fig A and B). Additionally, samples were taken from the Nilgiris, Nammakal, Dindigul, and Salem districts for the purpose of detecting CLAs in mandarin oranges and from the Nilgiris, Nammakal, Dindigul, Salem, Coimbatore, and Tirunelveli districts for the purpose of detecting CLAs in acidLime samples were examined utilising pairs of OI1 and OI2C primers for amplification of a 1160 base 16s rDNA fragment. The outcomes demonstrated that all samples collected from distinct mandarin orange orchards had amplified partial 16S r RNA (the most conserved portion of the CLAs bacteria genome) at position 1160. The mandarin orange orchards that displayed higher PDI also displayed steady CLAs amplification. Gene-specific primers A2 and J5, which demonstrated amplification of the 703 bp targeting (rpiKAJL-rpoBC) beta operon gene, further verified the CLAs infection. Each field's samples overwhelmingly tested positive for CLAs. Mandarin orange sequences for the 16 s r RNA gene and the beta operon gene were entered into the NCBI GenBank database and given accession numbers. Tables 1 and 3 list the primers used and their accession numbers. Mandarin orange samples were drawn at random from each field during the study and combined. Samples from Kanalkadu displayed the highest incidence of HLB illness among the various localities (100%). Out of the 25 samples, every one accurately displayed clinical symptoms and used particular primers to amplify the CLAs 1160bp 16s rDNA fragment. The gene-specific primer pair A2 and J5, which amplified at a 703 bp nucleotide fragment, was used to further confirm the samples from Kanalkadu (100% PDI) and Thandikudi (92% PDI), which showed the highest disease incidence (Fig. C, D and G). The sequenced amplified DNA samples were entered into the NCBI database to receive accession numbers. (Table.1), samples taken from the Patlankadu orchard from the Dindigul district also showed PDI of 91.66% which is followed by Kanalkadu and Thandikudi .

3.2 Detection of CLAs in acid lime:

Acid lime samples from the three investigated localities Kallar in Niligirs district, Amayapalem in Dindigul district, and Sanakrankoil in Tirunelveli district showed the highest frequency of HLB illness (100%). Out of the 25 samples, every one amplified 1160 bp fragments of 16s r DNA using particular primers and displayed accurate symptoms. Also, the samples were amplified using the particular A2/J5 primer pair, yielding a 703 bp DNA fragment (Fig. E , F and G). The sequenced amplified DNA samples were entered into the NCBI database to get accession numbers after being sequenced. 25 samples were taken during the survey of the 18 fields in the Tirunelveli district, and all 25 samples tested positive for CLAs. The majority of them are from the Tirunelvelidistrict's

Sankarankoil field. In the Kallar region of the Nilgiris district and the Amayapalam region of the Dindigul District, all 25 samples (out of 25) were positive. This is the first report of CLAs being found in the midribs of acid lime and mandarin oranges in Tamil Nadu. By employing mid-ribs, this technique may demonstrate that the CLAs inoculum can transfer from all sources, including source and sink through phloem.

3.3 Comparing nested PCR amplification and conventional PCR amplification

The nested PCR method greatly enhanced the sensitivity to detect Las up to 10 times and 100 times compared to qPCR and conventional PCR. The PCR product was serially diluted in a range of 1, 10, 10^2 , 10^3 , 10^4 and 10^5 , which represented 10^5 to 1 pg/ μ l of Las DNA, and served as the templates to evaluate the sensitivity among nested PCR, conventional PCR and qPCR. The PCR product from **Sankarankoil** was diluted as the template for conventional PCR (Fig 2C), and the PCR product from **kanalkaadu** was diluted as the templates for the nested PCR (Fig 2D) and qPCR. To evaluate the amplification efficiency of qPCR, Las DNA (Fig 2B) serially diluted in a range of 10, 10^2 , 10^3 , 10^4 and 10^5 served as the templates. Analysis of the dilution series of the DNA showed that nested PCR (Table 3 and 4) detected much lower template DNA concentrations than conventional PCR (Table 5). The amplification efficiency (AE) of qPCR for second set of nested PCR was therefore estimated to be 0.91 based on the equation $AE = [10^{-1/\text{slope}} - 1]$ [40]

3.4 Identifying the sensitivity of Q-PCR for Conventional PCR, First primer set of Nested PCR, Second set of Nested PCR.

The purified sample was performed with Q-PCR and the CT value obtained at various dilutions were as follows 25.680, 20.248, 16.440, 12.280, 10.440, 8.982. The r^2 (Regression coefficient) Es (ES slope derived estimate of amplification efficiency and Nt (Number of amplicon molecules at fluorescent threshold). Mostly for every Q-PCR run the peak become raised at 24 to 28 cycles. The conventional PCR run the peak become raised a 16 to 24 cycles. Where as for Nested PCR the peak raised between 27 to 31 cycles, For the second set of nested PCR the peak raised between 25 to 27 cycles. The melting peak increased at 79°C for all the dilutions of the conventional PCR product band in gel electrophoresis which initiate the peak at 0.002 dF/dT and end between 0.006 DT to 0.008 DT. The melting peak increased at 72 to 81 °C for all the dilutions of the Nested PCR first set of primers product band in gel electrophoresis which initiate the maximum peak was initiated at 0.008 and end at 0.0013 DF/DT. The melting peak increased at 79°C for all the dilutions of the Nested PCR second set of primers product band in gel electrophoresis which initiate the maximum peak was initiated at 0.005 and end at 0.0014 DF/DT. The r^2 (Regression coefficient) Es (ES slope derived estimate of amplification efficiency and Nt (Number of amplicon molecules at fluorescent threshold) was 0.997108, 88.56 % and 15637388142.32 obtained for first set of nested PCR, where as 0.991053469, 91.21 % and 5197936724.43 obtained

for second set of nested PCR. Ultimately the amplicon molecules amplification efficiency were high for the second set of Nested PCR. (Table 6 and 7; Plate A, B, C, D, E and F).

4. DISCUSSION

4.1 Huanglongbing disease survey

In the field, strange signs including erratic blotching called mottling and chlorosis (Baranwal, 2004) have been seen. Our findings suggested that Tamil Nadu was home to many CLAs isolates. The symptoms of acid lime and mandarin orange species are comparable to those described by Garnier and Bove in 1993 [43, 44, 32 and 25].

4.2 Detection of CLAs in mandarin orange and acid lime

The identification of *Ca. Liberibacter* spp. from an HLB sample always should be based on analysis of the 16S rDNA locus. Primer set OI1/OI2c amplifies the signature sequence for '*Ca. Liberibacter asiaticus*' and should be used for the species determination. Our data corroborate the results of previous studies (Xialong). A band of 703 bp was obtained from all infected plants. No amplification was observed from water or healthy citrus plants. The band of 703 bp was sequenced and compared with data from GenBank. The results indicated that the sequence was closely related to the record of *Candidatus Liberibacter asiaticus*, since they were a part of the ribosomal protein gene (*rplJ*). It can be concluded that the causal agent of citrus Huanglongbing (greening) disease. (45).

4.3 Nested PCR, q-PCR conventional PCR and their sensitivity

Accordingly, the primer sensitivity was inversely correlated with the template concentration, the detection system sensitivity order was nested PCR > qPCR > conventional PCR, meaning that the nested PCR was 10 times more sensitive than that of qPCR and 100 times more sensitive than of conventional PCR. The amplification efficiency (AE) of qPCR for second set of nested PCR was therefore estimated based on the equation $AE = [10^{-1/\text{slope}} - 1]$ based on the equation given by Rutledge *et al.*, 2003. Ahmad [30] indicated that the efficiency of amplification affected the sensitivity of qPCR. In this study, the efficiency of amplification of second set of nested PCR purified DNA from qPCR was 91.21%. The results showed that the efficiency of

amplification was not a key factor for the **sensitivity** of the assay. (ahmad 2009). **The mathematics of PCR dictate that amplification efficiency and Nt are independent entities.** In reality Nt is determined solely by the fluorescent threshold (Ft), and as such its value is independent of the parameters impacting PCR amplification. Indeed, this interrelationship between Nt and Ft has important practical implications, based on the principle that Ft does not directly reflect the number of amplicon molecules, but rather DNA mass at fluorescent threshold (Mt). This in turn dictates that Mt could be used to predict Nt for any amplicon of known size, if it is assumed that amplicon size and base composition do not significantly impact DNA fluorescence. The calculated **results** in excel sheet were supported the above mentioned equations which was given Rutledge et al., 2003 (40).

CONCLUSION

The disease prevalence revealed that 189 samples out of 210 samples showed HLB disease incidence of CLAs, accounting for 90% of the disease incidence in mandarin oranges throughout the entire Tamil Nadu, and 202 samples showed positive out of 216 samples, accounting for 93.52 percent of acid lime samples in Tamil Nadu. With the exception of Nilgiris, Dindigul, and Thirunelveli, all field samples were positive, but the incidence varied across each district. Research has shown unequivocally that the PCR approach may be used to detect HLB illness, citrus greening, and to validate the presence of the pathogen CLAs. The survey has provided a thorough insight of the highest incidence of HLB occurrence in Tamil Nadu's citrus-growing regions. The low concentration and asymmetrical distribution of CLAs in citrus presented challenges for other detection methods, but the Nested PCR and Q-PCR based assay proved to be the most effective method for the pathogen's detection. Excellent HLB diagnosis by Nested PCR was required for identifying infected plants and developing citrus nurseries free of the illness. Using Nested PCR for HLB detection should make epidemiological research easier and contribute to HLB control. The citrus belts of Tamil Nadu can benefit greatly from these discoveries in terms of HLB diagnosis, as well as the provision of HLB free seedlings to registered nurseries for mass multiplication. The data imply that the nested PCR primer pairs were able to identify Las in a variety of symptomatic tissues and geographical locations. The nested PCR primer pairs are suitable for numerous citrus cultivars and geographical regions as a consequence. The nested PCR primers will be a very valuable addition to the current Las detection techniques.

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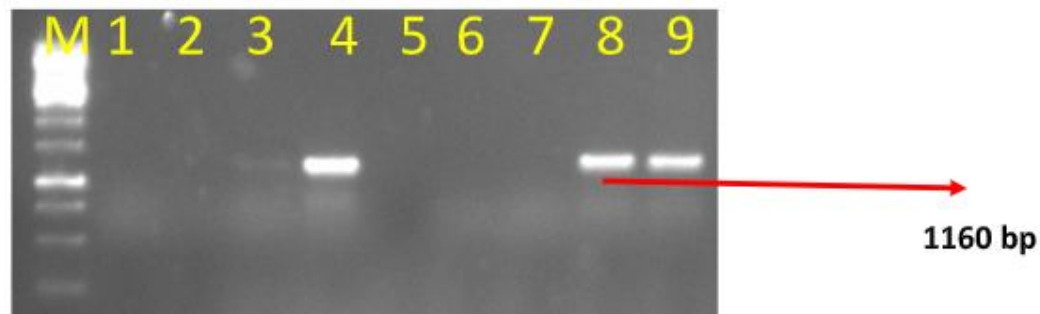


Fig. A. Gel picture of OI1 and OI2C Primers PCR amplified Mandarin orange midrib samples showing 1160 base pairs similar to *Candida Liberibacter asiaticus*; M -1 kb Ladder; Lane 1,2,3- Negative control; 4- Kanalkadu (Dindigul) mandarin orange sample ; Lane 5,6,7 -Negative control; Lane 8- Thandikudi (Dindigul); Lane 9 - Sankarankoil (Thirunelveli) acid lime sample.



Fig. B. Gel picture of A2 and J5 Primers (*rpiKAJL-rpoBC*) gene PCR amplified mandarin orange mid-rib and acid lime midrib samples showing 703 base pairs similar to *Candidatus Liberibacter asiaticus*; M -1 kb Ladder; Lane 1- Kanalkadu (mandarin orange); Lane 2- Negative control; Lane 3- Thandikudi (mandarin orange); Lane 4,5 -Negative control; 6- Sankarankoil (acid lime)

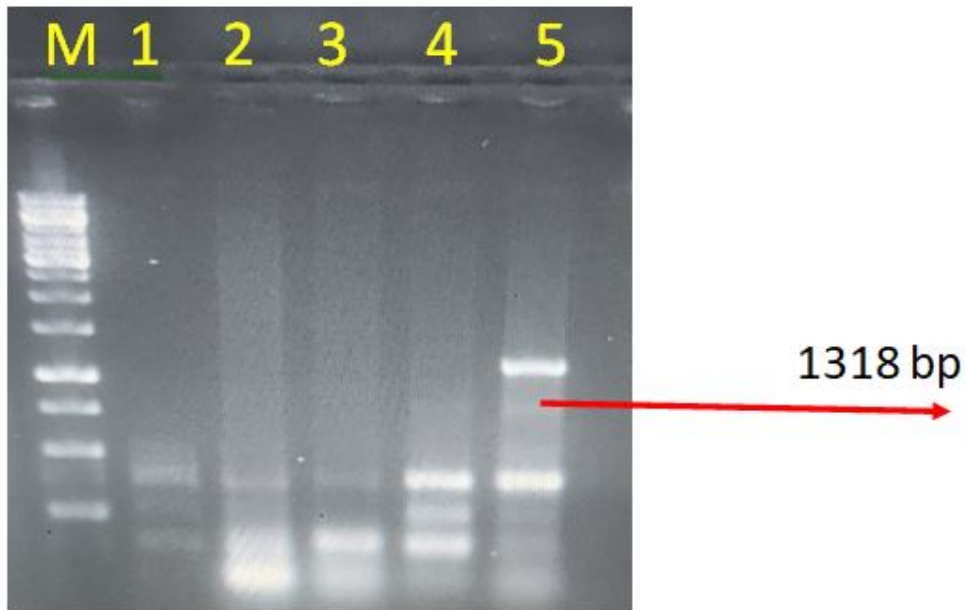


Fig. C. Nested PCR amplification with first set of primers (F1 and B1) at annealing temp- 58°C showing 1318 bp amplification. M- 1Kb Ladder; La 1,2,3,4 – Negative control ; Lane 5 - Mandarin sample (kanalkadu , Dindigul district)

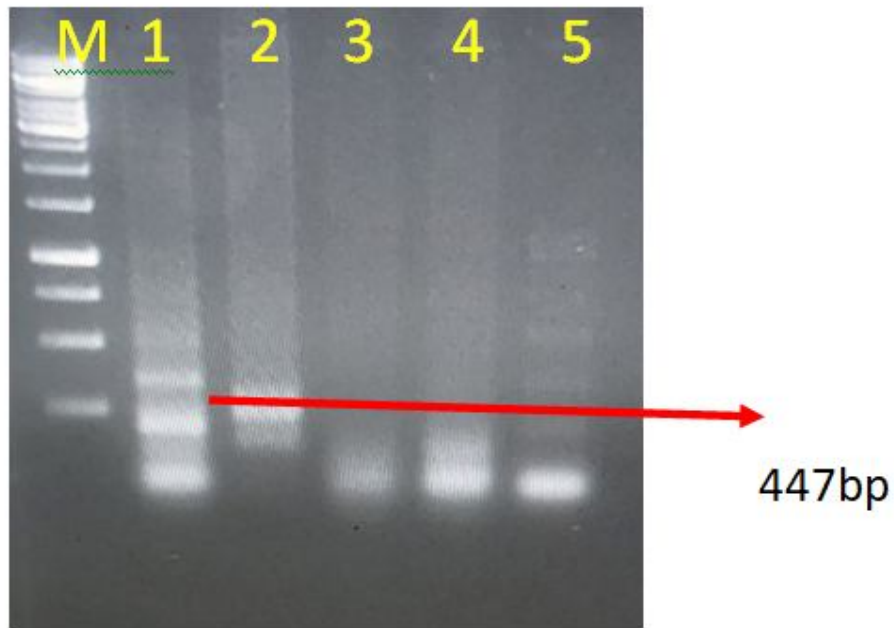


Fig. D. Nested PCR amplification with second set of primers (F3 and B3) at annealing temp- 62°C showing 443 bp amplification. M- 1Kb Ladder. Lane 1 – Mandarin sample (kanalkadu , Dindigul district) nested PCR first set PCR product as a DNA template amplified with second set of nested PCR primers; Lane 2,3,4,5 – Negative control.

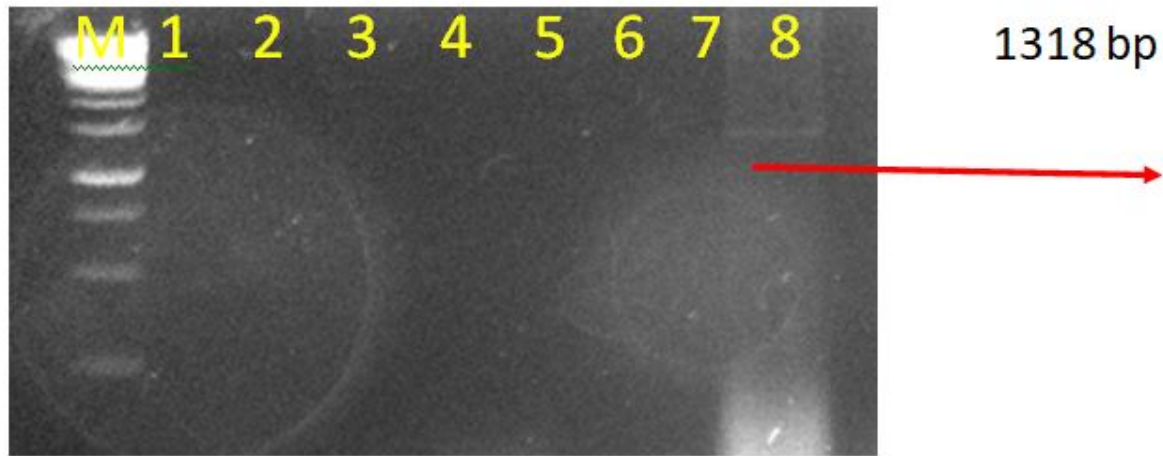
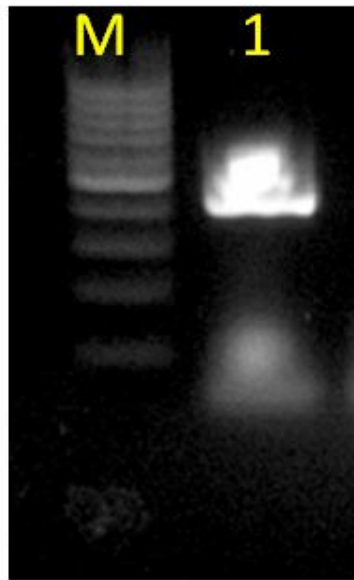


Fig. E. Nested PCR amplification with first set of primers (F1 and B1) at annealing temp- 58°C showing 1318 bp amplification. M- 1Kb Ladder; La 1,2,3,4,5,6,7 – Negative control ; Lane 8 - Acid lime sample (Sanarankoil , Thirunelveli district)



447bp

Fig. F . Nested PCR amplification with second set of primers (F3 and B3) at annealing temp- 62°C showing 443 bp amplification. M- 100bp Ladder Lane 1 – acid lime sample (Sankarankoil , Thirunelveli district) nested PCR first set PCR product as a DNA template amplified with second set nested PCR primers.

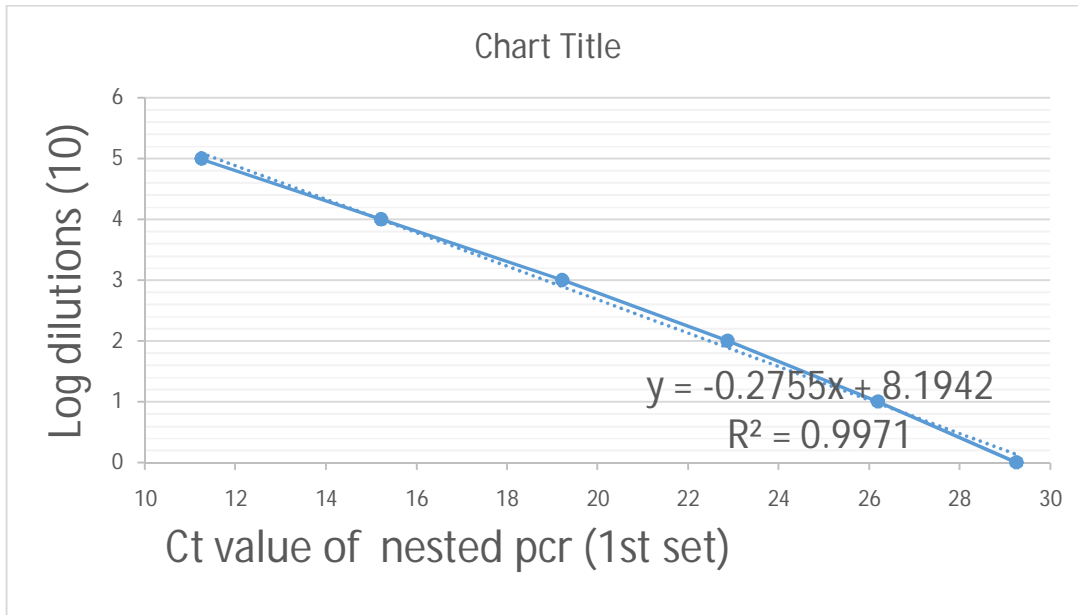


Fig. G. Standard Curve constructed through the Log dilutions 0 to 5 with the average Ct Values of Q-PCR formed through first set of nested PCR primers (F1 and B). The linear regression value obtained through the standard curve is 0.9971.

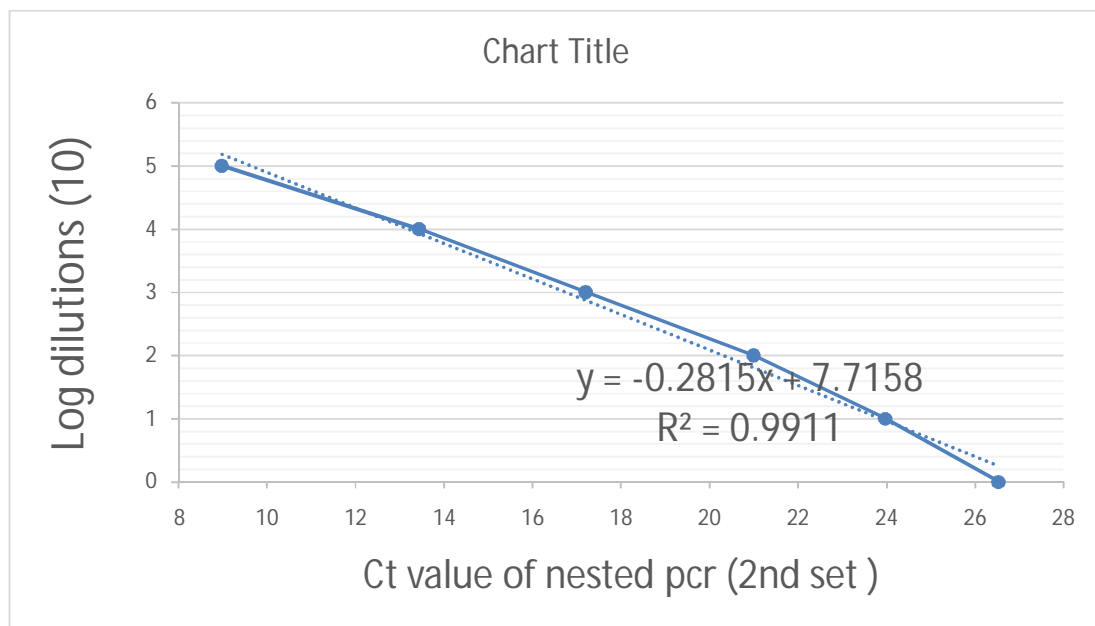


Fig. H. Standard Curve constructed through the Log dilutions 0 to 5 with the average Ct Values of Q-PCR formed though second set of nested PCR primers (and B). The linear regression value obtained through the standard curve is 0.9911.

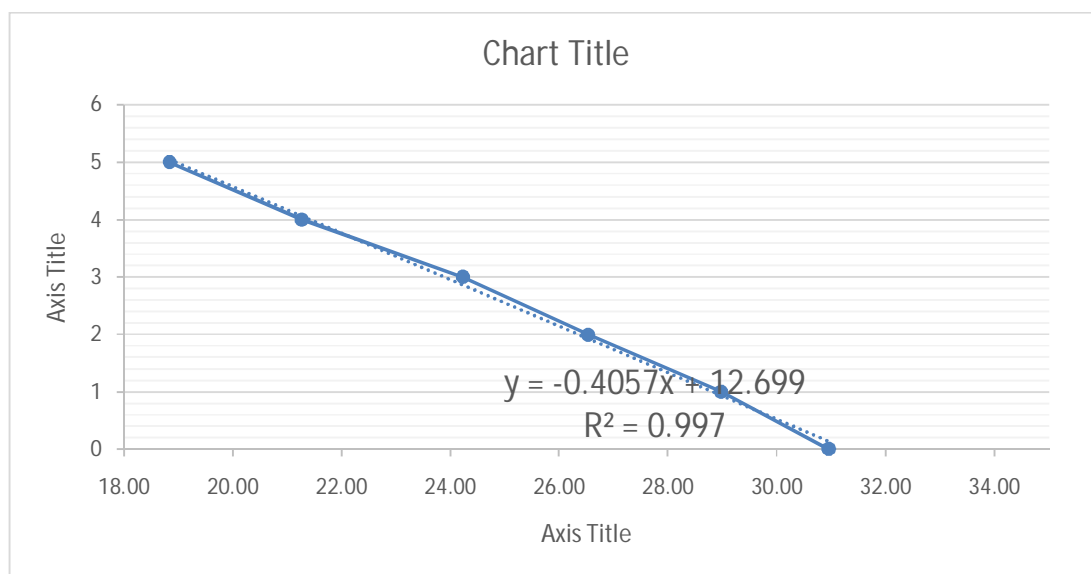


Fig. I. Standard Curve constructed through the Log dilutions 0 to 5 with the average Ct Values of Q-PCR formed through Conventional primers (OI1 and OI2). The linear regression value obtained through the standard curve is 0.9911.

Table.1 Survey and Detection of CLas bacterium in mandarin orange plants at various locations in Tamil Nadu, PDI (%) represent the mean of three replicates, ± represent standard deviation, The treatment means are compared using Duncan's Multiple Range Test (DMRT). In a column, mean followed by a common letter (s) are not significantly different ($p=0.05$)

S.NO	Locations	District	Latitude(°N)	Longitude(°E)	Altitude(metres)	Number of trees evaluated per field	Number of trees infected	PDI(%)	HLB infection
1.	Yercaud	sSalem	11.7748	78.2097	1623	24	22	88.66±2.345 ^{bc} (70.389)	+VE
2.	Kolli hills	Namakka l	11.2485	78.3387	1400	22	20	86.54±2.292 ^{bc} (68.525)	+VE
3.	Conoor	Nilgiris	11.3502	76.7961	1850	23	20	90.00± 3.740 ^{bc} (71.995)	+VE
4.	Katteri	Nilgiris	12.4382	78.1764	1850	22	20	88.90±2.142 ^{bc} (70.737)	+VE
5.	Ooty	Nilgiris	11.4102	76.6950	2240	20	16	84.00±1.455 ^c (66.435)	+VE
6.	Kanalkadu	Dindigul	10.3022	77.6911	1292	25	25	100.00±0.00 ^a (87.134)	+VE
7.	patlankadu	Dindigul	10.3093	77.6435	1325	25	22	91.66±0.465 ^b (73.839)	+VE
8.	thandikudi	Dindigul	10.3093	77.6435	1500	25	23	92.00±1.457 ^b (73.784)	+VE
9.	Thadiyankudisai	Dindigul	10.2368	77.7088	1098	24	21	90.00±1.670 ^{bc} (71.823)	+VE
	CD(0.05)							4.409	
	SE(m)							1.428	
	SE(d)							2.019	
	CV							3.502	

S.NO	Locations	District	Latitude (°N)	Longitude (°E)	Altitude (metres)	Number of Trees evaluated per field	Number of trees infected	PDI(%)	HLB infection
1.	Pollachi	Coimbatore	10.6609	77.0048	293	28	26	92.85±2.461 ^{bc} (74.655)	+VE
2.	Sankarankoil	Thirunelveli	9.168902	77.5413	154	28	28	100±0.000 ^a (87.134)	+VE
3.	Kolli hills	Namakkal	11.2485	78.3387	1400	24	21	94.00 ± 2.311 ^{bc}	+VE

Table.2 Survey and Detection of CLas bacterium in acid lime plants at Various locations in Tamil Nadu, PDI(%) represent the mean of three replicates, ± represent standard deviation, The treatment means are compared using Duncan's Multiple Range Test (DMRT). In a column, means followed by a common letter (s) are not significantly different ($p=0.05$).

								(76.989)	
4.	Thadiyankudisai	Dindigul	10.2368	77.7088	1098	20	18	96.42 ±2.483 ^{ab} (79.498)	+VE
5.	Ayyampalayam	Dindigul	10.2271°	77.7482	281	25	25	100±0.000 ^a (87.134)	+VE
6.	Conoor	Nilgiris	11.350208	76.7961	1850	20	18	90.00±2.381 ^{cd} (71.642)	+VE
7.	Kallar	Nilgiris	11.3360°	76.8617°	372	25	25	100±0.000 ^a (87.134)	+VE
8.	Ooty	Nilgiris	11.4102	76.6950	2240	24	21	87.50±2.567 ^d (69.394)	+VE
9.	Yercaud	sSalem	11.7748	78.2097	1623	22	20	96.90±2.025 ^{ab} (80.295)	+VE
	C.D.(0.05)							3.580	
	SE(m)							1.184	
	SE(d)							1.674	
	C.V.							2.152	

Table.3 Mandarin orange Clas DNA sample at various dilutions and each sample is having four replicates and dilutions is at $1, 10^1, 10^2, 10^3, 10^4$ and 10^5 , The Nested PCR first set of primers (F1,B1 primers PCR amplified sample at annealing temperature 58°C and run through Q-PCR F and R primers and Ct value and SD was noted.

	No	Rep#1	Rep#2	Rep#3	Rep#4	Av. Ct	SD
10^5	10000000	11.1276	11.0152	11.5772	11.24	11.24	0.242811752
10^4	1000000	14.9156	15.6766	15.0678	15.22	15.22	0.328789578
10^3	100000	18.84344	19.61256	19.03572	19.42028	19.228	0.351053645
10^2	10000	21.96864	23.11284	23.57052	22.884	22.884	0.673686719
10^1	1000	25.676	26.462	26.462	26.2	26.2	0.370523953
1	100	28.3822	29.26	29.5526	29.8452	29.26	0.632088243

Table.4 Mandarin orange Clas DNA sample at various dilutions and each sample is having four replicates and dilutions is at $1, 10^1, 10^2, 10^3, 10^4$ and 10^5 , The Nested PCR second set of primers (F3,B3 primers PCR amplified sample at annealing temperature 62°C and run through Q-PCR F and R primers and Ct value and SD was noted.

	No	Rep#1	Rep#2	Rep#3	Rep#4	Av. Ct	SD
10^5	10000000	8.89218	8.80236	9.25146	8.982	8.982	0.194033377
10^4	1000000	13.1712	13.8432	13.3056	13.44	13.440	0.290337183
10^3	100000	16.8658	17.5542	17.0379	17.3821	17.210	0.314210174
10^2	10000	20.16768	21.21808	21.63824	21.008	21.008	0.618458774
10^1	1000	23.5004	24.2198	24.2198	23.98	23.980	0.339128412
1	100	25.7438	26.54	26.8054	27.0708	26.540	0.573329527

Table.5 Mandarin orange Clas DNA sample at various dilutions and each sample is having four replicates and dilutions is at $1, 10^1, 10^2, 10^3, 10^4$ and 10^5 , The conventional PCR primers (OI1 and OI2C primers PCR amplified sample at annealing temperature 60.5°C and run through Q-PCR F and R primers and Ct value and SD was noted.

	No	Rep#1	Rep#2	Rep#3	Rep#4	Av. Ct	SD
10^5	10000000	18.6516	18.4632	19.4052	18.84	18.84	0.406990516
10^4	1000000	20.8544	21.9184	21.0672	21.28	21.28	0.45970054
10^3	100000	23.7552	24.7248	23.9976	24.4824	24.24	0.442559826
10^2	10000	25.4784	26.8054	27.3362	26.54	26.54	0.781316445
10^1	1000	28.4004	29.2698	29.2698	28.98	28.98	0.40983909
1	100	30.033625	30.9625	31.272125	31.58175	30.9625	0.668866446

Table. 6 The r^2 (Regression coefficient) E_s (ES slope derived estimate of amplification efficiency and N_t (Number of amplicon molecules at fluorescent threshold) were derived through Log factor and Average ct value derived through Q-PCR of first set nested PCR diluted PCR sample ($1, 10^1, 10^2, 10^3, 10^4$ and 10^5 dilutions. The amplification efficiency is 88.56% for first set of nested PCR primers,

Dilutions:	No	Log (No)	nested pcr primers 1
10 ⁵	10000000	7.000	11.240
10 ⁴	1000000	6.000	15.220
10 ³	100000	5.000	19.228
10 ²	10000	4.000	22.884
10 ¹	1000	3.000	26.200
1	100	2.000	29.260
		r2:	0.997108345
		Eff:	0.885614852 (88.56%)
		Nt:	15637388142.32

Table.7 The r2 (Regression coefficient) Es (ES slope derived estimate of amplification efficiency and Nt (Number of amplicon molecules at fluorescent threshold) were derived through Log factor and Average ct value derived through Q-PCR of second set nested PCR diluted PCR sample (1, 10¹, 10², 10³, 10⁴ and 10⁵ dilutions. The amplification efficiency is 91.21 % for second set of nested PCR primers.

Dilutions	No	Log (No)	nested pcr primers 2
10 ⁵	10000000	7.000	8.982
10 ⁴	1000000	6.000	13.440
10 ³	100000	5.000	17.210
10 ²	10000	4.000	21.008
10 ¹	1000	3.000	23.980
1	100	2.000	26.540
		r2:	0.991053469
		Eff:	0.912189811 (91.21%)
		Nt:	5197936724.43

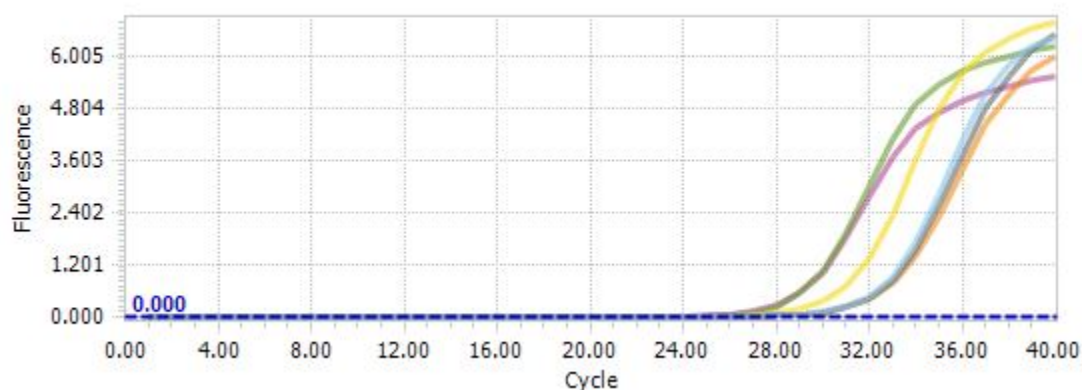


Plate. A. Amplification curve shown raised curves at 27th to 31 cycle for mandarin orange samples and 28th cycle for 10⁵ dilution and curves became stationary at 4.824 to 6005 fluorescence level for

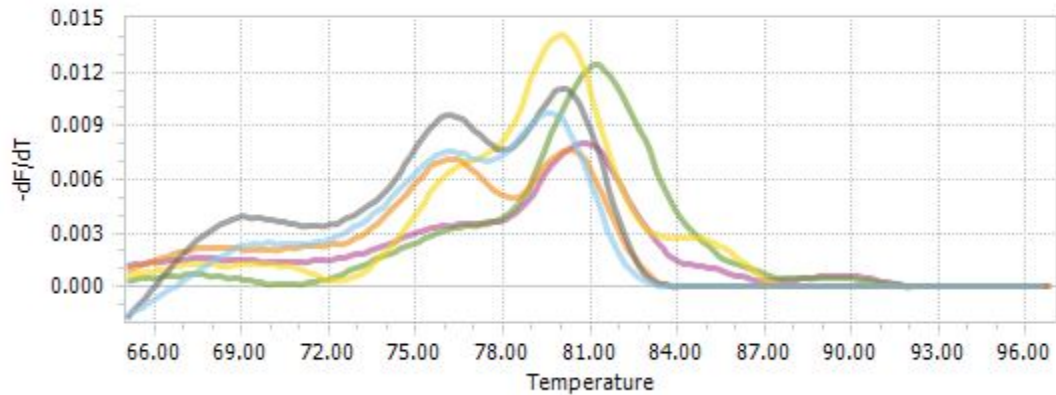


Plate. B . Melting peaks of first set nested PCR amplified PCR product run through Q-PCR primers shown the change in fluorescence level (df/dt) at 0.001 to 0.002 with the increase of peak at temperature 79.0°C (yellow) for mandarin orange sample at Various dilution peaks (1, 10¹, 10², 10³, 10⁴ and 10⁵ (sky blue ,violet, yellow, pink, blue red), the yelow peak at 10⁵ dilution showed the minimum ct value11.24 and 0.013 is the maximum peak at a temperature of 80 °C.

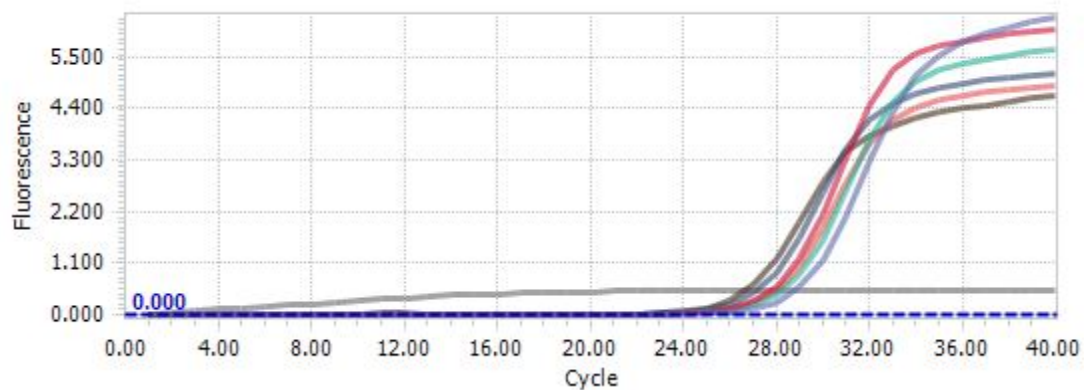


Plate. C. Amplification curve shown raised curves at 24th cycle for mandarin orange (black) and 27th cycle for mandarin orange (light blue) 10⁵ dilution and curves became stationary at 3.300 to 4.400

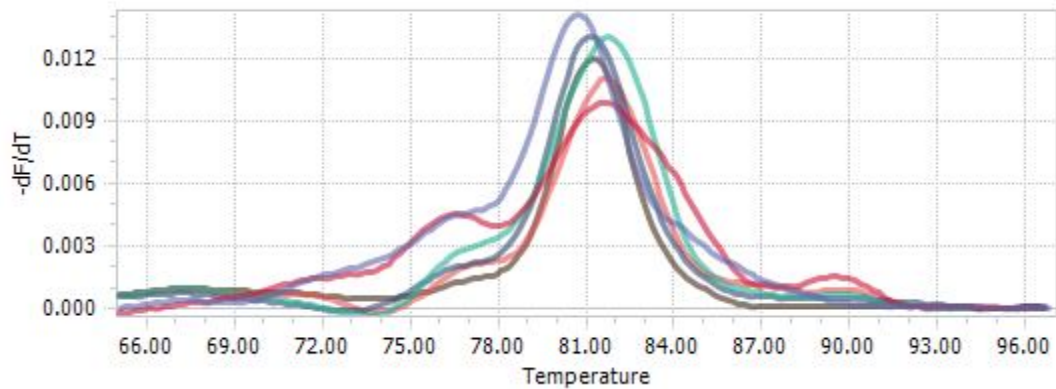


Plate. D . Melting peaks of second set nested PCR amplified PCR product run through Q-PCR primers shown the change in fluorescence level (df/dt) at 0.001 with the increase of peak at temperature 80.0°C (light blue) for mandarin orange sample at Various dilution peaks (1, 10¹, 10², 10³, 10⁴ and 10⁵ (dark blue ,pink, blue, red, sky blue), the light blue peak at 10⁵ dilution showed the minimum ct value 8.982 and 0.013 is the maximum peak at a temperature of 80 °C.

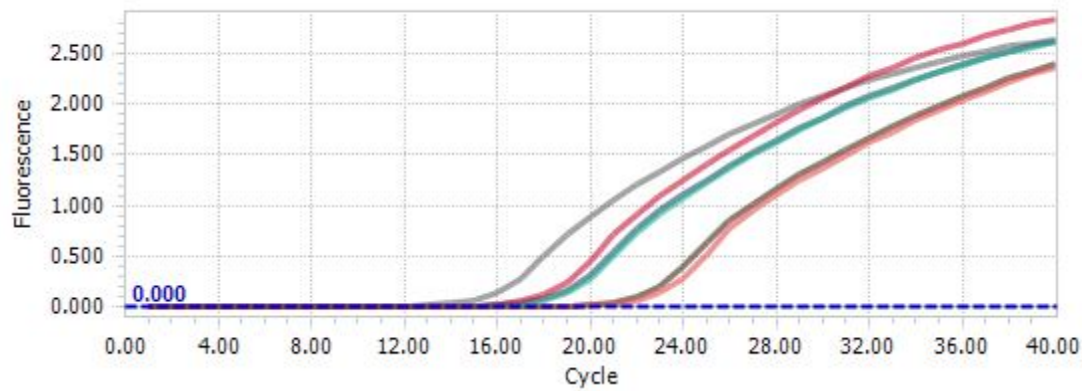


Plate. E Amplification curve shown raised curves at 16th to 21st cycle for mandarin orange samples and 18th cycle for 10⁵ dilution and curves became stationary at 2.000 to 2.400 fluorescence level

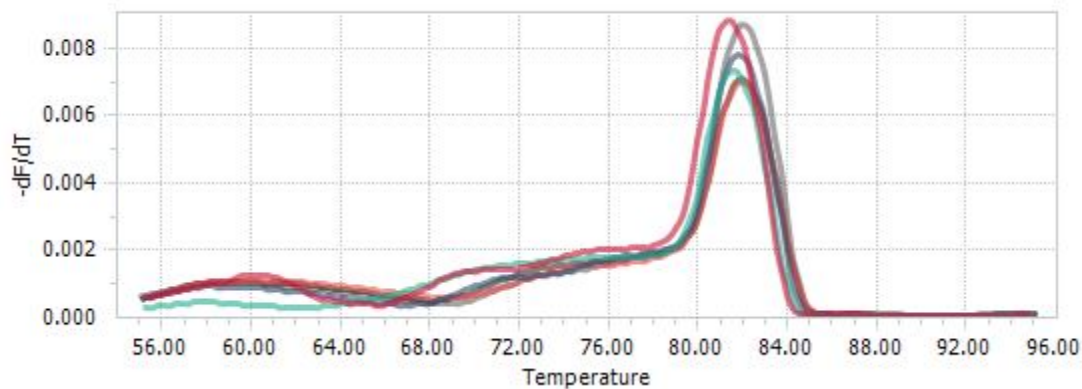


Plate. F. Melting peaks of Conventional PCR amplified PCR product run through Q-PCR primers shown the change in fluorescence level (df/dt) at 0.002 with the increase of temperature at 78.0°C for mandarin orange sample at Various dilution peaks (1, 10¹, 10², 10³, 10⁴ and 10⁵ (Red,skyblue,blue,kight blue and dark red), the dark red peak at 10⁵ dilution showed the minimum ct value 18.84 and 0.008 is the maximum peak at a temperature of 85 °C.

UNDER PEER REVIEW