The impact of post acquisition period on detection of \textit{Candidatus} Liberibacter solanacearum in tomato potato psyllid

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\textbf{Abstract} The impact of post acquisition period (PAP) on detection of \textit{Candidatus} Liberibacter solanacearum (Lso) in adult tomato potato psyllid (TPP; \textit{Bactericera cockerelli}) was investigated using qPCR assays on whole insects. Lso in adults was initially tested immediately after acquisition access periods (AAP) of 15, 24, 48, 72, 96 h or 1 week on Lso-infected tomato leaves. These results were compared with those from adults with 15 or 24 h AAP followed by 24-, 48- or 72-h PAP on Lso-free tomato leaves. From 15–20 individuals of each treatment tested, 5–15\% were positive for Lso immediately after AAP of 15 or 24 h. That increased to 84\% and 100\% with 24- and 48-h PAPs respectively, but decreased to 25–47\% after 72 h. Longer AAP (48 h–1 week) without PAP did not exceed 55\% acquisition success. The phloem ingestion and possible acquisition of Lso during 15 and 24 h plant access periods are discussed.

\textbf{Keywords} tomato/potato psyllid, \textit{Candidatus} Liberibacter solanacearum, post acquisition period, phloem ingestion, detection.

\section*{INTRODUCTION}
Zebra chip is an economically important disease of potato (\textit{Solanum tuberosum} L.) in the United States, Mexico, Central America and New Zealand (Munyaneza et al. 2007; Munyaneza 2012). This disease is caused by \textit{Candidatus} Liberibacter solanacearum (Lso), also known as \textit{Candidatus} Liberibacter psyllaurous (Hansen et al. 2008). Lso has been shown to severely disrupt carbohydrate flow in potato plants, leading to Zebra chip symptoms in the tubers (Buchman et al. 2011). Lso is a phloem-limited, Gram-negative, unculturable bacteria that was initially found in solanaceous crops (Liefting et al. 2008, 2009) and is transmitted by tomato potato psyllid (TPP) \textit{Bactericera cockerelli} (Šulc) (Hemiptera: Triozidae) (Munyaneza et al. 2007; Hansen et al. 2008). This bacterium was found later in carrot plants in Finland, Sweden and Norway, where it was vectored by \textit{Trioza apicalis} Forster (Munyaneza 2012). Recent reports from the Canary Islands and Spain have found Lso in carrot and celery crops where it is vectored by \textit{B. trigonica} (Alfaro-Fernandez et al. 2012 a, b). TPP was first reported in New Zealand in 2006 and has now established throughout the North and South Island. Lso disease symptoms
The primary aim of this study was to test the effect of different post acquisition periods on the detection of Lso in TPP using qPCR, based on the assumption that Lso multiplies in TPP after acquisition. The efficiency of feeding of TPP during 15 h and 24 h plant access periods was also measured.

**MATERIALS AND METHODS**

**Insects and plants**

Lso-positive and -negative TPP colonies were maintained in separate glasshouses. The Lso-negative TPP colony was reared on clean (Lso-free) capsicum plants (*Capsicum annuum* L. ‘Giant Bell’) and the Lso-positive TPP colony was reared on tomato plants (*Solanum lycopersicum* L. ‘Moneymaker’) showing Lso symptoms and testing positive for Lso in regular qPCR tests. Both colonies were maintained in 50 × 50 × 50 cm steel frame cages covered with insect net at 24±1°C and under natural light. Insect samples from both colonies were tested monthly by qPCR to confirm presence or absence of Lso. Adults from the Lso-negative TPP colony were used in the experiments. Tomato plants (‘Moneymaker’) were infected with Lso by exposing groups of six clean plants to 50–60 adult TPP from the Lso-positive colony for a week in a 50 × 50 × 70 cm steel frame cage covered with fine net. Those plants were then sprayed with Confidor® at 1 g/litre to kill all the psyllids and left in a separate room in the glasshouse until symptoms typical of Lso appeared. The presence of Lso in symptomatic plants was confirmed by qPCR tests. Lso-free tomato plants (‘Moneymaker’) were grown from seeds in individual pots (10 × 10 × 11.5 cm) with potting mix in an insect- and pathogen-free glasshouse.

**Acquisition access periods**

In the first experiment, TPP adults from the Lso-free colony were exposed to excised symptomatic leaves of Lso-positive tomato plants for AAPs of 15, 24, 48, 72 or 96 h or 1 week. Each excised leaf from a Lso-infected tomato plant was placed into a vertically-held Petri dish (85 × 15 mm), with the stem extending through a foam plug fitted into an opening cut in the base of both leaves.
the lid and the dish. The end of the stem was submerged in a 50 × 10 mm tube of water held under the Petri dish and the tube was plugged with the foam. Twenty TPP were left on each leaf for each of the assigned AAPs (a total of 120 TPP on six excised leaves) after which the insects were individually collected into 0.6 ml tubes and stored dry at -20°C for subsequent qPCR testing for Lso. Two samples from each leaf were also collected for qPCR tests to confirm the presence of Lso in the leaves. A logistic regression model was used to test the relationship between AAP and the number of Lso-positive TPP. The model was fitted using the GENMOD procedure in SAS, allowing for over-dispersion of the binomial (SAS Institute Inc. 2008).

**Post-acquisition period**

The impact of three different post-acquisition periods (PAPs) on detecting Lso in TPP was tested in the second experiment. Twenty TPP adults were placed on each of six Lso-symptomatic leaves excised from Lso-positive tomato plants, set up as described for the first experiment. Sixty TPP were removed after a 15-h AAP and immediately released onto a Lso-free tomato plant housed in a cage (50 × 50 × 50 cm) covered with fine net. The remaining sixty TPP on Lso-positive leaves were removed after a 24-h AAP and immediately released onto another Lso-free tomato plant in a cage. Groups of 15–20 TPP were then removed from each plant after 24, 48 and 72 h of PAPs and individually stored in 0.6 ml tubes at -20°C for subsequent qPCR testing for Lso.

Both experiments were carried out at 24±1°C and 16:8 h (light:dark) periods.

**EPG monitoring to measure real feeding time**

The duration of actual phloem ingestion by tethered TPP adults was measured for 15-h and 24-h plant access periods in a follow-up EPG test. The aim of this test was to estimate the actual periods TPP spent ingesting phloem and potentially acquiring Lso from infected leaves during the AAPs tested in the second experiment. The stylet penetration activities of 20 TPP adults (Lso-free) on shoots of Lso-infected tomato plants were recorded for 24 h. To prepare for the EPG tests, adults were collected from the Lso-free colony into 1.5 ml tubes (4–5 adults/tube) and the insects were immobilised under CO₂ for 2–3 s before wiring. Individual TPP were attached to a 2 cm long, 18.5 μm diameter gold wire, with a drop of silver conductive paint (n-butyl acetate solvent; Ladd Research Industries, Williston, VT). The other end of the gold wire was attached to a 3 cm long copper wire connected to the head of a 3 mm diameter copper nail, which fitted tightly into the input probe of the EPG monitor. Wired insects were left for 15 min to recover before being placed on the abaxial surface of Lso-infected tomato leaves. The plant electrode (a thick copper wire) was inserted into the water-filled tube into which the leaf stem had been placed. The stylet penetration of each insect was monitored for 24 h on a Giga-4 DC-EPG system (WF Tjallingii, Wageningen, The Netherlands) under fluorescent lights for a 16 h photoperiod with the photo phase of 6 am – 10 pm at 24±1°C. Four insects were monitored on four channels of the EPG monitor simultaneously, and a total of 20 insects was monitored. Data output was digitized using DI-720 analogue to digital board (DATAQ instruments). Stylet penetration data were acquired, stored and measured using WinDaq Pro+ software (DATAQ instruments). The duration between the start of the recording and the first phloem ingestion of individual TPP was measured. The total phloem ingestion period of each TPP was also measured for the first 15 h of the recording and then the full recording period (24 h). The efficiency of feeding by individual TPP (calculated by dividing the total duration of phloem ingestion by the recording period) for the first 15 h and the last 9 h of the total 24 h recording period were compared using a paired-t test of the null hypothesis that the average of the differences between two efficiencies is 0.

**Detection of Lso**

Individual plant samples were prepared for testing by grinding with a Qiagen Tissue Lyser and the DNA extracted with a QIAxtractor®
system according to the manufacturer’s protocol (QIAGen, 2010). Individual TPP were processed according to the Zygem protocol outlined in Beard & Scott (2013). The qPCR analysis to detect Lso was performed according to the method of Beard & Scott (2013), with a modification being a transfer to the Roche 480 LightCycler® Platform (Roche Diagnostics) using Roche Sybr Green I Master Mix. This detection platform has a larger sample number capacity, allowing both the Lso-exposed and control samples to be run in unison. An observed Ct shift meant samples were run for 45 cycles instead of 40 (see Puketapu et al. (2013) for a more detailed description of detection methods). Samples that amplify in the last five cycles of a qPCR programme are at the limits of detection for this test and detection standards ranged from 1 x 10⁹ gene copies to 10 copies. However, only 2 µl of the total 50 µl DNA extraction of each psyllid was used, therefore, it was possible that positive individuals with a very low Lso titre would not be detected before cycle 40. It was therefore decided that in the case where one or more of the three replicates tested positive after cycle 40 and the product showed the correct melt curve, the sample was counted as positive when interpreting the data.

RESULTS

All the leaves used for Lso acquisition were confirmed to be positive for Lso by qPCR. Results revealed that a single TPP can acquire Lso within 15 h of being placed on an infected plant, but not more than 15% of the insects were detected as positive for Lso when tested immediately after a 15- or 24-h AAP (Figure 1). The proportion of Lso-positive insects increased to 55% when insects were tested after a 48-h AAP or longer. The results of the logistic model showed evidence of a relationship (P=0.0377) between the percentage TPP testing positive for Lso and the length of the AAP. However, the fitted curve showed a trend of increasing numbers of Lso-positive TPP with increasing AAP.

The PAP on Lso-free tomato plants increased the proportion of Lso-positive TPP in groups with 15- or 24-h AAP, compared with those without PAP (Figure 2). All TPP with a 15-h AAP followed by a 48-h PAP were found to be positive for Lso. TPP that had a 24-h AAP followed by a 24-h PAP were 84.2% positive for Lso. The proportion of TPP with Lso was less in TPP groups with a 72-h PAP than the groups with 24-h or 48-h PAP.

![Figure 1](image_url)

**Figure 1** The observed values and fitted line showing the relationship between the percentage tomato potato psyllids testing positive by qPCR for *Candidatus Liberibacter solanacearum* (Lso) immediately after exposure to Lso-infected plants and the time spent on the plants (acquisition access period).
The phloem ingestion periods of tethered TPP measured in the follow-up EPG test provided an estimate of the duration of feeding of the TPP given 15-h and 24-h AAPs in the other experiments. The EPG waveforms were characterised as shown in Butler et al. (2012). The recordings started with non-probing baseline (np) (Figure 3). The initial electrical contact of stylet with plant tissues (probe) appeared with the highest amplitude waveform, which continued with a mixture of waveforms with different frequencies, representing saliva secretion while probing in the vicinity of epidermal and parenchyma cells (C waveform) and sometime xylem ingestion (G waveform). The initial contact with phloem tissues was indicated by D waveform, which was usually followed by phloem salivation (E1 waveform) and then phloem ingestion (E2 waveform). E2 frequently changed to E1 during the phloem phase. Two of 20 TPP monitored

Figure 2 The percentage of tomato potato psyllids testing positive for *Candidatus* Liberibacter solanacearum (Lso) by qPCR after 15 or 24 h of exposure to Lso-infected plants (acquisition access period, AAP) followed by 0, 24, 48 or 72 h of post acquisition periods on healthy plants.

Figure 3 An overview of electrical penetration graph recordings of the stylet penetration behaviour of tomato potato psyllids on tomato leaves. The upper plot shows non-probing baseline (NP), saliva secretion while probing in the vicinity of epidermal and parenchyma cells (C), ingestion from xylem tissues (G) and initial contact with phloem tissues (D). Expanded phloem salivation (E1) waveforms and phloem ingestion (E2) waveforms are shown in the lower plot.
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on EPG did not show phloem ingestion during the 24 h of recording so they were not included in the graph (Figure 4). Thirteen TPP started ingesting phloem during the first 15 h and 18 TPP showed phloem ingestion during 24 h. Variability in phloem ingestion was noticed between individual TPP. The mean duration between the start of the recordings and the first phloem ingestion was 11.19±1.6 h, with a minimum duration of 107 min. The mean duration of phloem ingestion measured within the first 15 h and 24 h of recordings was 5.96±1.08 and 9.01±1.05 h respectively. T-test results indicated that the average efficiency of feeding in the last 9 h was significantly higher (P=0.031) than that in the first 15 h.

DISCUSSION

The results demonstrated a trend of detectable Lso infections in TPP increasing with longer AAPs. It is suggested that PAPs had an impact on qPCR results as more psyllids tested positive for Lso after a PAP than when tested immediately after their AAP. Variability in qPCR results from TPP tested after acquisition may also be influenced by the Lso titre at the acquisition site and the length of time required for accumulation or multiplication of Lso in insect bodies. It has been shown previously that Lso acquisition occurs during phloem ingestion on an infected plant (Sandanayaka et al. 2012).

In contrast to Beard et al. (2013), samples returning only 1/3 positives were considered to be positive in the present analysis, instead of 2/3. This is because the original method of Beard et al. (2013) had a detection limit of 5 bacterial genome copies, but transfer to the Roche 480 Light Cycler platform meant that the limit of detection was raised to 10 bacterial genome copies per extraction. Thus, a positive on the Roche 480 Light Cycler platform meant that a much greater number of copies of the bacterial genome was present. It is recognised that PCR runs above 40 cycles can produce artefacts or non target amplifications giving false positives. However, in the present study the addition of a melt curve assessment of the qPCR product was used to confirm that the right target was being amplified. As a single bacterium can multiply to a larger, more readily detectable, titre given enough time, it was felt that the 1/3 positive approach allowed for dilution factor effects associated with the large DNA extraction volume.

EPG recordings revealed that 65% and 90% of tethered TPP ingested from phloem tissues during plant access periods of 15 h and 24 h respectively. It can be assumed that the free TPP in the first and

Figure 4 Real time feeding (phloem ingestion) of 18 tomato potato psyllids during the first 15 h and full electrical penetration graph recording period of 24 h.
the second experiments were as likely to feed on phloem during the same plant access periods (15 h and 24 h) as the tethered insects. Thus, it is likely that the proportions of free TPP that acquired Lso in AAPs of 15 or 24 h would be equal to or greater than those observed in the EPG experiment. The average time taken by a tethered TPP to start phloem ingestion from the beginning of the plant access period was 11.19 h. All of the TPP with 15-h AAP followed by a 48-h PAP tested positive for Lso, indicating that all TPP had acquired Lso, which suggested that the combination of PAP and AAP improved detection of Lso in TPP. The results of qPCR after a 24-h PAP showed that 84.2% of TPP acquired Lso during a 24-h AAP, which correlates to EPG data showing 90% of TPP successfully ingested phloem during a 24-h plant access period. Therefore it is suggested that every single TPP that succeeded in feeding and acquiring Lso after 15- or 24-h AAPs may not have detected as positive for Lso, when tested immediately after AAP. Presumably the success of detection may be impacted by the amount of Lso acquired by the TPP during the AAP and the length of time available for Lso to multiply inside the insect body during any PAP.

Tethered TPP demonstrated that the efficiency of feeding in the first 15 h is lower than that in the last 9 h as the insects may need some adaptation time to a new environment. Although the tethering effect on the insect behaviour has to be considered (Tjallingii 1986), an average TPP spent a considerable proportion of the first 15 h attempting to locate phloem tissues and spent more time feeding during the last 9 h. When phloem tissues were located by the TPP, most of the time was spent on phloem ingestion and most likely acquiring Lso. Therefore it is suggested that a greater number of TPP acquire Lso when given a 24-h AAP than with a 15-h AAP.

The proportions of Lso-positive TPP decreased when TPP had a 72-h PAP, suggesting that longer feeding on Lso-free plants may reduce the Lso concentration in the TPP’s body. This agrees with the findings of Pelz-Stelinski et al. (2010) who reported that the proportion of Las-positive adult D. citri, determined using qPCR, decreased over time when held on healthy plants. Declines in bacteria titre in the insect over time may be the result of initial concentrations of bacteria acquired by the insect, age of the insect or negative effects of the bacterium on the insect host (Purcell 1982).

In the first and the second experiments, Lso donor leaves were picked from the same tomato plant to reduce the possible influence of the physiology of host plant on acquisition. A small number (n=20) of TPP were tested on each Lso donor leaf to avoid competition for feeding, especially during longer plant access periods. Although an increased acquisition rate was expected from TPP after the longer confinement on Lso-infected leaves, the number of TPP that tested positive for Lso changed little among the TPP groups that had 48 h, 72 h, 96 h or 7 days of AAP. Pelz-Stelinski et al. (2010) suggested that longer confinement of D. citri on Las-infected plants may allow time for the bacterial titre to increase in the insect. In the present study excised leaves from an infected plant were used as the source of Lso for acquisition. Although these leaves were kept fresh until 7 days AAP, TPP feeding might have been affected by the physiological changes in those leaves over time. Further investigation using whole plants for acquisition is required to confirm that. In studies carried out by Rashed et al. (2012), the highest percentage of acquisition success of Lso occurred when TPP had access to the whole plant and a higher proportion of TPP acquired Lso from stem tissue compared with leaflets and petioles. The effect of feeding site on acquisition success has been shown to be important in other vector-borne pathogen systems where the pathogen was heterogeneously distributed within the host (Daugherty et al. 2010).

Understanding the nature of the relationship between Lso and TPP is important for developing more effective control measures for zebra chip disease. Presumably a latent period is essential for Lso to multiply and enter the salivary gland of the psyllid before inoculation of another plant by the insect can occur. Munyanzeza (2012) has reported that latent period in TPP is about 2 weeks after an acquisition period of 8–24 h on Lso-infected potato plants. He also stated that the latent period
has been observed to be shorter when psyllids have fed on Lso-infected tomato plants, apparently because of a much higher Lso titre in tomato than potato. This study provides preliminary information on the time that might be required for Lso to multiply in the bodies of TPP.

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