



# A novel haplotype of ‘*Candidatus Liberibacter solanacearum*’ found in Apiaceae and Polygonaceae family plants

Minna Haapalainen · Satu Latvala ·  
Annika Wickström · Jinhui Wang · Minna Pirhonen ·  
Anne I. Nissinen

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**Abstract** A previously unknown haplotype of the plant pathogen ‘*Candidatus Liberibacter solanacearum*’ (Lso) was found in cultivated carrots and parsnips in eastern Finland. That same haplotype was found in western Finland, over 300 km away, in the family Polygonaceae, the species *Fallopia convolvulus* (wild buckwheat) and *Persicaria lapathifolia* (pale persicaria) growing as weeds within carrot and parsnip fields. The infected plants, both apiaceous and polygonaceous, showed symptoms of foliar discoloration. This is the first report of Lso bacteria in plants of the family Polygonaceae. The finding that the polygonaceous plants infected with a previously unknown haplotype of Lso were growing among the apiaceous plants infected with Lso haplotype C suggests that these two haplotypes might be transmitted by different vectors. Phylogenetic analyses showed that

the new haplotype, called haplotype H, is distinct from the previously characterized haplotypes and appears to have diverged early from their common ancestor. Multi-locus sequence analysis revealed four different sequence types (strains) within the haplotype H. These findings suggest that the haplotype H is likely to be endemic in northern Europe and that the genetic diversity within the Lso species is higher than previously assumed.

**Keywords** ‘*Candidatus Liberibacter solanacearum*’ · Haplotype · Phylogenetics · Polygonaceae · Parsnip · Carrot disease

## Introduction

‘*Candidatus Liberibacter solanacearum*’ (Lso; Liefting et al. 2009) is a Gram negative plant-pathogenic bacterium transmitted by psyllids. When feeding on the phloem sap of the host plant, the psyllids harbouring Lso deliver the bacteria into the phloem sieve cells, where the bacteria may then multiply and colonize the phloem systemically. Because of the dependency on a vector in accessing a plant, the natural host range of Lso is determined by the host plant range of the vector. Different psyllid species feeding on different plants and occurring in different geographical regions have been found to harbour diverse haplotypes of Lso. Thus far, eight distinct haplotypes have been found and named A, B, C, D, E, F, G, and U. The haplotypes A, B and F are associated with zebra chip disease of potato in America - the haplotype A also in New Zealand - and are transmitted

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M. Haapalainen (✉) · A. Wickström · J. Wang ·  
M. Pirhonen  
Department of Agricultural Sciences, University of Helsinki, P. O.  
Box 27, FI-00014 Helsinki, Finland  
e-mail: minna.haapalainen@helsinki.fi

S. Latvala · A. I. Nissinen  
Natural Resources Institute Finland (Luke), Natural Resources,  
Tietotie, FI-31600 Jokioinen, Finland

*Present Address:*  
J. Wang  
College of Plant Protection, Hebei Agricultural University, Leka  
South Street 2596, Baoding 071001 Hebei, China

by the tomato/potato psyllid *Bactericera cockerelli* (Hansen et al. 2008; Nelson et al. 2011; Swisher Grimm and Garczynski 2019). The haplotypes C, D and E have been found in Europe, D and E also in North Africa and D in the eastern Mediterranean area. These haplotypes are associated with diseases in apiaceous plants, including carrot and celery (Hajri et al. 2017; Mawassi et al. 2018; Munyaneza et al. 2010; Nelson et al. 2013; Tahzima et al. 2014; Teresani et al. 2014). In northern Europe, Lso haplotype C is transmitted by the carrot psyllid *Trioza apicalis* (Nissinen et al. 2014), whereas in the Mediterranean region the haplotypes D and E are transmitted by the psyllid *Bactericera trigonica* (Antolinez et al. 2017; Mawassi et al. 2018). In addition to *T. apicalis*, Lso haplotype C was found in *Trioza anthrisci* and its host plant *Anthriscus sylvestris*. Haplotype U was found in *Trioza urticae* and its nettle host plant (*Urtica dioica*) in Finland (Haapalainen et al. 2018). Recently, another haplotype of Lso was discovered in samples of a wild solanaceous plant *Solanum umbelliferum* from California, U.S., and called haplotype G (Mauck et al. 2019).

Lso is an unculturable bacterium, and thus the different strains of this species have only been defined by genetic markers. The different haplotypes, which can be seen as groups of closely related genotypes, are defined by the sequence of 16S rRNA and the sequence of 50S ribosomal protein *rplJ/rpL* gene region (Nelson et al. 2011, 2013). To distinguish the different strains within a haplotype, a multi-locus sequence typing (MLST) scheme based on seven protein-coding genes was developed (Haapalainen et al. 2018). These genetic markers can be used for the identification of the haplotype and strain of the Lso found in a plant or insect sample and also for characterizing previously unknown strains.

Since the first discovery of Lso in symptomatic carrots in Finland (Munyaneza et al. 2010), plant samples have been collected from carrot fields every year and the presence of Lso tested by PCR (Haapalainen et al. 2017, 2018). Recently, Lso haplotype C was also detected in parsnips grown beside carrots in Tavastia Proper region (Haapalainen et al. 2018). To assess the frequency of Lso infections in the parsnips grown close to the carrot fields where symptomatic plants had been observed, a survey on parsnip fields was carried out in the autumn 2018. The genotype of the bacterium was determined for selected Lso positive plant samples. However, for some of the samples the marker DNA sequences obtained did not match with any of the known Lso haplotypes.

In this study, we characterized these novel Lso genotypes and determined their phylogenetic relationship to the genotypes previously described.

## Materials and methods

### Plant samples

In autumn 2018, samples of carrots (*Daucus carota ssp. sativus*) and parsnips (*Pastinaca sativa*) showing symptoms of foliar discoloration were collected for Lso testing from different regions of Finland. Carrot samples were collected from ten fields in South Savo, six fields in Satakunta and two fields in Southwest Finland. Parsnip samples were collected from three fields in South Savo and from five fields in Satakunta, at locations close to carrot fields. In addition, samples of weeds with foliar discoloration were collected from one carrot field and one parsnip field in Satakunta. Single Lso positive carrot samples from South Savo and weed samples from Tavastia Proper that had been collected in the earlier surveys in 2013 and 2016 and given unfamiliar *rplJ/rpL* PCR products were included in the sequence analyses.

### DNA extraction, amplification and sequencing

For DNA extraction, 100 mg samples were cut from the petioles of the plants collected and these samples were homogenized in FastPrep tubes with lysing matrix A (MP Biomedicals, Santa Ana, CA, USA), using the FastPrep device at speed 5 for 40 s. DNA extraction was subsequently performed using the DNeasy Plant Mini kit protocol (Qiagen, Hilden, Germany), with DNA elution in 100 µl of nuclease-free water. For Lso detection, either end-point PCR with 16S primers OA2/Lsc2 (Haapalainen et al. 2017; Liefting et al. 2009) or real-time PCR with fluorescent probe (Li et al. 2006, 2009) was used as previously described (Haapalainen et al. 2017, 2018). For analysing the Lso genotype in the positive samples, end-point PCR was run with the ribosomal *rplJ/rpL* gene region primers rp01F(CL514-F)/rp01R(CL514-R) (Liefting et al. 2009) and with the primers amplifying fragments of the seven protein-coding genes - *adk*, *atpA*, *fbpA*, *ftsZ*, *glyA*, *groEL* and *gvrB* - included in the multi-locus sequence typing (MLST) scheme (Haapalainen et al. 2018). For a sample found to contain a novel haplotype of Lso also the 16S–23S intergenic spacer region was amplified with primers

LpFrag4-1611F and LpFrag4-480R (Hansen et al. 2008). The PCR products were purified using QIAquick PCR purification kit (Qiagen) according to the manufacturer's instructions. Capillary sequencing was performed at the Jokioinen laboratory of the Natural Resources Institute Finland and at the FIMM sequencing service of the University of Helsinki, both using an automated sequencer (Applied Biosystems, Waltham, MA, USA).

### Phylogenetic analysis

After removing the primer sequences and trimming the 5' ends, the sequences of the *rplJ/rplL* gene region were aligned by Clustal W and the alignment was then corrected based on the translated amino acid sequences of RplJ and RplL. A maximum parsimony (MP) phylogenetic tree was constructed in MEGA7 (Kumar et al. 2016), treating the gaps as the fifth character state, and bootstrapping with 1000 replications. The 16S rRNA gene sequences and the 16S–23S intergenic spacer regions were also aligned by Clustal W in MEGA7 and the small nuclear polymorphisms searched. The seven MLST gene fragments were aligned after primer trimming and then concatenated, forming a supermatrix, and used for pair-wise comparisons of the different sequence types as previously described (Haapalainen et al. 2018). To include the shorter haplotype G sequences in the analysis, all the gene fragments were trimmed at the 5' end accordingly. Due to the lack of published sequence material, the haplotypes E and F could not be included in the MLST analyses. Phylogenetic distance matrixes were calculated using the substitution models TN93 (Tamura and Nei 1993), F84 (Felsenstein 1984) and K80 (Kimura 1980), and clustering was performed by unweighted-pair group method with arithmetic means (UPGMA). The maximum-likelihood tree was constructed using RAxML (Randomized Axelerated Maximum Likelihood) program v8.2.11 (Stamatakis et al. 2005) and applying GTR + G model to each partition (locus alignment). A majority rule (50%) consensus tree was computed out of 1000 bootstrap replicates.

## Results and discussion

A novel haplotype of Lso in carrots and parsnips in eastern Finland

A previously unknown haplotype of the plant pathogen '*Candidatus* Liberibacter solanacearum' was found in

symptomatic cultivated carrots and parsnips in South Savo, eastern Finland. Two carrot plants infected with this haplotype, from now on called haplotype H, were collected as samples in the same locality within South Savo in 2013 and 2016 (Table 1; Fig. 1). In 2018, carrots infected with Lso at low bacterial titres (Ct values from 36 to 38) were found in seven of the ten fields sampled in South Savo and for two of the fields the Lso haplotype could be confirmed and was found to be C. In all the carrot fields sampled in Satakunta and Southwest Finland regions the symptomatic carrots were infected with Lso haplotype C. In the previous study in 2013 and 2014 carrot samples had been collected for Lso tests in Tavastia Proper, Ostrobothnia and South Savo regions in addition to Satakunta and Southwest Finland, and plants infected with Lso haplotype C had been found (Haapalainen et al. 2017). Thus, haplotype C is the major haplotype of Lso occurring in carrots in Finland and haplotype H is rare. In 2018, a survey on the occurrence of Lso in parsnips was carried out. Parsnips infected with Lso were found in all the three fields sampled in South Savo, and for two of these fields the Lso haplotype was identified as H. In the first field, four out of five samples were Lso positive and in the second field, 7.2 km distant from the first one, all five were Lso positive. In the third field three out of six samples were Lso positive, but due to the very low bacterial titre (real-time PCR Ct value >37) the haplotype could not be resolved. In contrast, the symptomatic parsnips collected in Tavastia Proper in 2016 (Haapalainen et al. 2018) and in Satakunta region in 2018 were infected with haplotype C. Thus, in these apiaceous crop plants the haplotype H has thus far only been detected within one sampling area located in eastern Finland.

The carrots infected with Lso haplotype H showed strong foliar discolouration, similar to the symptom previously associated with a high titre of Lso haplotype C (Haapalainen et al. 2017; Nissinen et al. 2014). No proliferation of the shoot was observed, and the PCR test for phytoplasma with primers P1/fU5 and P7/rU3 (Crosslin et al. 2006) was negative. However, it is possible that other biotic or abiotic factors could have contributed to the discolouration symptom. In those Lso-infected parsnips that showed some symptoms, the symptoms were mild and included red discolouration of the petioles and slightly wrinkled leaves with blotchy bronze discolouration. Two of the plants (18-11P2 and 18-11P3) had brown discolouration in the root tip. No significant correlation was observed

**Table 1** Plant samples infected with ‘*Candidatus Liberibacter solanacearum*’ haplotype H

Sample ID	Plant species	Sampling year	Sampling region	Real-time PCR Ct value for Lso <sup>a</sup>	<i>ftsZ</i> site 252	<i>gyrB</i> site 155	Haplotype H sequence type <sup>c</sup>
13-356	<i>Daucus carota</i>	2013	South Savo	22.22	T	G	1
16-P8	<i>Daucus carota</i>	2016	South Savo	19.70	T	A	2
16-UT1	<i>Persicaria lapathifolia</i>	2016	Tavastia Proper	29.23	T	A	2
16-KT3	<i>Fallopia convolvulus</i>	2016	Tavastia Proper	23.15	T	A	2
18-T1	<i>Persicaria lapathifolia</i>	2018	Satakunta	27.02	T	G	1
18-T2	<i>Persicaria lapathifolia</i>	2018	Satakunta	21.92	G	G	3
18-11P1	<i>Pastinaca sativa</i>	2018	South Savo	22.59	T	G	1
18-11P7	<i>Pastinaca sativa</i>	2018	South Savo	21.58	G	A	4
18-11P2	<i>Pastinaca sativa</i>	2018	South Savo	22.42	T	nd <sup>b</sup>	nd
18-11P3	<i>Pastinaca sativa</i>	2018	South Savo	21.43	T	nd	nd
18-11P4	<i>Pastinaca sativa</i>	2018	South Savo	33.76	T	nd	nd
18-11P8	<i>Pastinaca sativa</i>	2018	South Savo	32.62	G	nd	nd

<sup>a</sup> Average of the three PCR results of the plant petiole sample

<sup>b</sup> nd, not determined

<sup>c</sup> Based on the combination of the small nuclear polymorphisms



**Fig. 1** Regions in Finland where plants infected with ‘*Candidatus Liberibacter solanacearum*’ (Lso) were found. Lso haplotype C (diamonds) was detected in carrot and parsnip samples from Satakunta, Tavastia Proper, southwest Finland, south Savo and in the previous study (Haapalainen et al. 2017) also in carrots from South Ostrobothnia. Haplotype H was detected in carrots and parsnips in south Savo (solid circle) and in samples of polygonaceous plants from Satakunta and Tavastia Proper (open circles)

between the root/shoot weight ratio and the Lso titre in parsnips in this study: Spearman’s rank correlation coefficient was  $-0.193$  with a  $p$  value  $0.199$ . Previously, carrots infected with Lso haplotype C were shown to have a reduced root weight in a greenhouse experiment (Nissinen et al. 2014), whereas for the samples collected from fields the effect was not statistically significant, due to the differences between carrot varieties and growth conditions (Haapalainen et al. 2017).



**Fig. 2** Foliar discoloration symptoms in *Persicaria lapathifolia* plant infected with ‘*Candidatus Liberibacter solanacearum*’ haplotype H

**Table 2** Polymorphic sites in the 50S ribosomal protein *rplJ/rplL* gene region, the 16S rRNA gene and the 16S–23S intergenic spacer region of ‘*Candidatus Liberibacter solanacearum*’

DNA region	Nucleotide position <sup>a</sup>	Haplotype of ‘ <i>Candidatus Liberibacter solanacearum</i> ’ <sup>b</sup>								
		U	A	D	E	C	B	F	G	H
50S ribosomal proteins <i>rplJ/rplL</i> gene region (637–640 nt)	29	T	T	T	T	T	T	G	G	T
	54	G	G	G	G	C	G	G	G	G
	90	G	G	G	G	G	G	G	G	A
	93	A	A	G	A	A	A	A	A	A
	111	C	C	C	C	T	C	C	C	C
	140	G	G	G	G	G	C	G	G	G
	145	A	A	A	A	A	A	A	A	G
	150	A	A	A	A	A	A	A	A	G
	160	T	C	T	T	C	C	C	C	C
	162	G	G	G	G	T	T	T	T	T
	166	G	G	G	G	G	G	A	A	G
	168	A	A	A	A	A	A	G	G	A
	171	A	A	G	A	A	A	A	A	A
	179	A	A	A	A	A	A	A	A	C
	183	G	G	G	G	G	T	T	T	G
	193	G	G	G	A	G	G	G	G	G
	220	C	C	A/C	C	C	C	C	C	C
	251–252	AA	A-	AA	AA/A-	AA	A-	–	–	A-
	257	G	G	G	G	G	A	G	G	G
	267	–	–	–	–	–	–	–	AA	–
	323	T	T	C	C	T	T	T	T	T
	383	C	T	C	C	C	C	C	C	C
	393–395	–	–	–	–	GCT	–	GTT	GTT	GCT
	397	T	T	T	T	T	C	C	C	C
	410	T	T	T	T	T	T/C	T	T	T
	415	T	C	C	C	C	C	C	C	C
	432	G	G	G	G	T	G	G	G	G
	438	T	G	G	G	G	G	G	G	G
	464	G	T	G	G	G	G	G	G	G
	470	A	A	A	A	G	A	A	A	A
	482	T	T	T	T	T	T	C	T	T
	512	A	A	A	A	A	A	A	A	T
	518–519	AA	GA	GA	GA	AA	AA	AG	AG	AG
	526	A	A	A	A	A	G	A	A	A
	545	C	C	T/C	C	C	C	C	C	T
	551	G	G	G	G	G	G	G	G	A
	554	T	T	T	T	T	T	T	T	C
	584	G	G	G	G	G	A	A	A	G
	587	T	C	C	C	C	C	C	C	C
	599	A	G	A	A	A	A	A	A	A
614	A	A	A	A	A	A	G	G	G	
620	G	G	G	G	G	A	A	A	A	
629	A	A	A	A	A	A	A	A	G	

**Table 2** (continued)

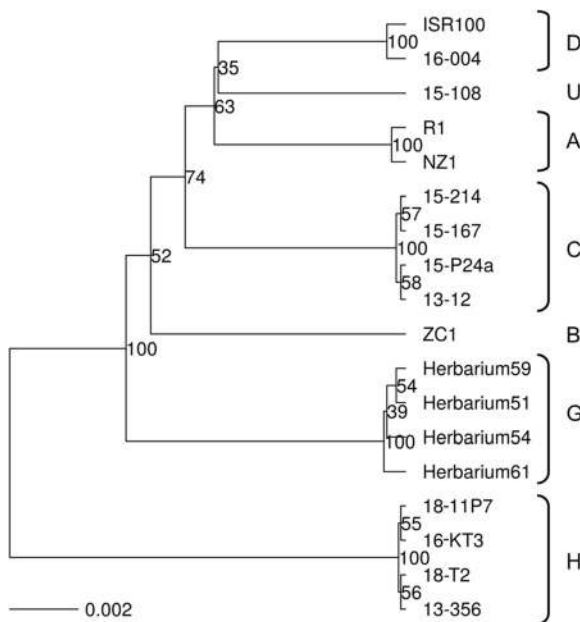
DNA region	Nucleotide position <sup>a</sup>	Haplotype of ' <i>Candidatus</i> Liberibacter solanacearum' <sup>b</sup>								
		U	A	D	E	C	B	F	G	H
16S internal fragment (1168 nt)	27	A	A	A	G	A	A	A	A	A
	28	C	C	T	C	C	C	C	C	C
	63	A	A	A	G	A	A	G	G	G
	100	A	A	A	A	A	A	A	A	G
	124	T	T	T	T	T	G	T	T	T
	271	A	A	A	A	A	A	C	C	A
	412	C	C	C	C	C	C	C	T	C
	436	G	G	G	G	G	G	A	A	G
	493	T	T	T	T	T	C	C	C	C
	597	G	G	G	G	G	G	G	A/G	G
	833	A	A	A	A	A	A	A	A	G
	871	C	C	C	T	C	C	C	C	C
	951	G	A	G	A	G	A	G	G	A
	985	G	G	A	G	G	G	G	G	G
1031	G	G	A	G	G	G	G	G	G	
1116	T	C	T/C	nd <sup>c</sup>	C	C	C	nd	T	
16S–23S intergenic spacer region	39	G	A	A	nd	A/G	A	nd	A	A
	72	C	C/T	C	nd	C	C	nd	C	C
	92	A	A	G	nd	A	A	nd	A	A
	98	C	C	T	nd	C	C	nd	C	C
	177	A	A	A	nd	A	A	nd	G	A
	199	–	–	–	nd	–	–	nd	T	–
	209–210	G -	AT	G -	nd	GA	GA	nd	GA	AA
	219–220	T -	–	–	nd	T -	T - /TT	nd	–	–
	273	T	T	T	nd	C	T	nd	T	T
	296	G	G	G	nd	G	A	nd	G	G
	368	T	T	T	nd	T	T	nd	A	T
	372	T	T	T	nd	T	T	nd	T	C
	408	C	C	C	nd	C	T	nd	C	C
	434	G	G	A	nd	G	G	nd	G	G
	558	C	C	C	nd	C	C	nd	C	T
	571	G	G	G	nd	G	A	nd	G	G
	592	T	T	T	nd	T	T	nd	T	–
	613	C	C	C	nd	C	T	nd	C	C
	627	C	T	T	nd	T	T	nd	T	T
	647	A	A	A	nd	A	A	nd	A/G	A
	732	C	C	C	nd	C	C	nd	A	C
	806	C	C	C	nd	C	C	nd	G	C
841	A	G	G	nd	G	G	nd	G	G	

<sup>a</sup> Nucleotide position in the *rplJ/rplL* sequence starting from GCTAATGATAACAGT (the alanine codon 98 of RplJ (L10)), in the 16S rRNA sequence starting from ATCTACCTTTTCTAC, and in the 16S–23S intergenic spacer region starting from TTTATGTTAAGGGCCC

<sup>b</sup> References for the haplotypes: U (Haapalainen et al. 2018), A, B and C (Nelson et al. 2011), D (Nelson et al. 2013), E (Teresani et al. 2014), F (Swisher Grimm and Garczynski 2019), and G (Mauck et al. 2019)

<sup>c</sup> nd = no data available





**Fig. 4** Phylogenetic tree of the different multi-locus sequence types of ‘*Candidatus Liberibacter solanacearum*’ (Lso), based on unweighted-pair group method with arithmetic means (UPGMA) and on TN93 substitution model. The genomic sequences of ‘*Ca. L. solanacearum*’ used in the analysis were of haplotype A strains NZ1 and R1 (JMTK01000005.1 and GCA\_000756225.1), haplotype B strain ZC1 (CP002371.1) and haplotype D strain ISR100 (GCA\_002918245.2). Samples 13–12, 15-P24a, 15-167 and 15-214 represent haplotype C, sample 16-004 haplotype D and 15-108 haplotype U. Herbarium samples 51, 54, 59 and 61 represent haplotype G. Samples 13-356, 16-KT3, 18-11P7 and 18-T2 represent the novel haplotype H. The Lso haplotypes are indicated by letters on the right

Despite careful inspection, no adult psyllids, nymphs or eggs were detected on the plants infected with Lso haplotype H, and thus the vector(s) transmitting this haplotype is still unknown. However, since all the *Ca. Liberibacter* species and their haplotypes characterized thus far have been shown to be psyllid-transmitted, it is likely that the vector of haplotype H also belongs to the family Psyllidae. All the Lso-positive *T. apicalis* and *T. anthrisci* samples for which the haplotype was determined, contained Lso haplotype C (Haapalainen et al. 2018), and thus there might be another vector transmitting haplotype H. The finding in 2018 that within the same field the parsnips were infected with Lso haplotype C and the *P. lapathifolia* plants with haplotype H suggests that there could have been two different vectors with different host range. The absence of adult psyllids, nymphs or eggs on the infected parsnips suggests that parsnip is not a host plant of the new vector(s). Parsnip

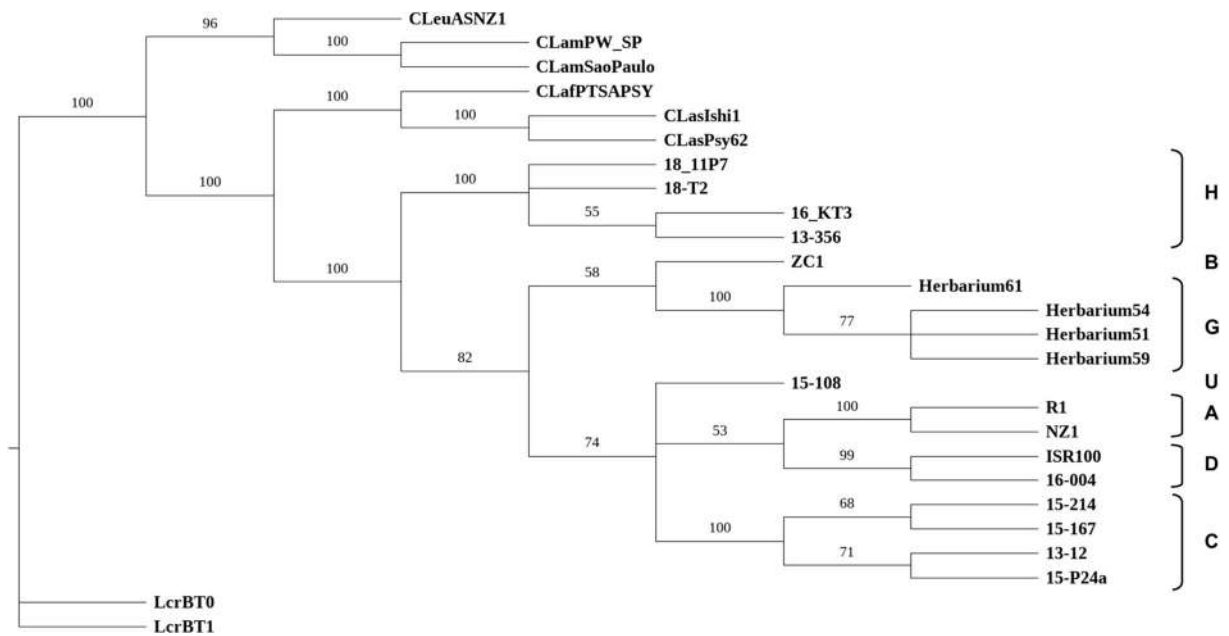
could, however, serve as an alternative host for *T. apicalis* and Lso haplotype C, if cultivated in crop rotation between carrot crops. Since *T. apicalis* showed clear oviposition preference for cultivated and wild carrot and coriander, while few eggs were laid on the other apiaceous plants (Valterová et al. 1997), it is unlikely that the carrot psyllids would prefer parsnip when both carrots and parsnips are cultivated during the same season. Psyllid species that occur in Finland and that could feed on *P. lapathifolia* are *Aphalara borealis*, *A. freiji* (syn. *A. polygona*, Burckhardt and Lauterer 1997) and *A. maculipennis* (Ossiannilsson 1992). However, there is very little knowledge about their host plant range and preferences. According to Ossiannilsson (1992) they are strictly oligophagous on a few species belonging to *Polygonum* family, in which *P. lapathifolia* also was previously classified.

#### Genetic markers of the haplotype H

The 50S *rplJ/rplL* sequence was identical in all the eight haplotype H samples which were sequenced from this region. Previously, it was suggested that in the 50S *rplJ/rplL* sequence an alanine codon (GCT) had been inserted in the haplotype C *rplL* gene (Haapalainen et al. 2018). However, as haplotype H also has an alanine codon at this position and the haplotypes F and G and *Ca. L. asiaticus* have a valine codon (GTT) (Table 2; Supplementary File S1), it now seems more likely that Lso haplotypes A, B, D, E and U have lost one codon from *rplL*. In comparison with haplotype C ST1, haplotype H has six nucleotide differences in the 16S rRNA 1168 nt internal fragment (Table 2; Supplementary File S2) and five nucleotide differences in the 16S–23S intergenic spacer region (Table 2; Supplementary File S3).

Analysis of the seven protein-coding sequences of the multi-locus sequence typing (MLST) scheme revealed one polymorphic site (252: T or G) in the *ftsZ* gene fragment and one polymorphic site (155: G or A) in the *gyrB* gene fragment between the haplotype H samples. The G > T change in *ftsZ* turns an aspartate codon GAT into a tyrosine codon TAT, whereas the nucleotide change at the site 155 in *gyrB* is silent. All the four different combinations of these two SNPs were found amongst the eight sequenced samples, which were accordingly divided into four different sequence types (STs) (Table 1). The other five MLST gene fragments were identical in these eight samples





**Fig. 5** Majority-rule consensus cladogram of ‘*Candidatus Liberibacter solanacearum*’ (Lso) and related (*Candidatus*) *Liberibacter* species constructed with RAxML (Randomized Axelerated Maximum Likelihood) program. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test with 1000 replicates are shown next to the branches. The genomic sequences used in the analysis were strains LcrBT0 and LcrBT1 (CP010522.1 and CP003789.1) for *Liberibacter crescens*, CLeuASNZ1 (PSQJ01000002.1) for ‘*Ca. L. europaeus*’, CLamSaoPaulo and CLamPW\_SP (CP006604.1 and GCA\_000350385.1) for ‘*Ca. L. americanus*’, CLafPTSAPSY

(CP004021.1) for ‘*Ca. L. africanus*’, CLasIshi1 and CLasPsy62 (AP014595.1 and CP001677.5) for ‘*Ca. L. asiaticus*’, NZ1 and R1 (JMTK01000005.1 and GCA\_000756225.1) for ‘*Ca. L. solanacearum*’ haplotype A, ZC1 (CP002371.1) for ‘*Ca. L. solanacearum*’ haplotype B and ISR100 (GCA\_002918245.2) for ‘*Ca. L. solanacearum*’ haplotype D. Samples 13–12, 15-P24a, 15–167 and 15–214 represent haplotype C, sample 16–004 haplotype D and 15–108 haplotype U. Herbarium samples 51, 54, 59 and 61 represent haplotype G and samples 13–356, 16-KT3, 18-11P7 and 18-T2 represent the novel haplotype H. The Lso haplotypes are indicated by letters on the right

(Supplementary File S4). The finding of four STs (bacterial “strains”) from such a small number of samples suggests that either the different STs have fairly equal frequencies or that the sampling was exceptionally lucky. The different alleles of *ftsZ* and *gyrB* do not seem to be linked to either western or eastern geographical location or to a certain plant species. This kind of pattern suggests that these STs have been present in Finland for a long time. For the Lso positive samples 18-11P2, 18-11P3, 18-11P4 and 18-11P8 only the *ftsZ* marker was sequenced. The results indicated that they also contained Lso haplotype H and that in 18-11P2, 18-11P3 and 18-11P4 the bacteria probably represented either ST1 or ST2 and in 18-11P8 either ST3 or ST4 (Table 1). Interestingly, the samples 18-11P7 and 18-11P8 that had G at the site 252 in *ftsZ* were collected from the same field, whereas the samples from 18-11P1 to 18-11P4 that had the SNP G > T were collected from another field. Thus, it is possible that the different fields had been visited

by vectors belonging to different populations, harbouring different strains of Lso haplotype H.

The *groEL* PCR product of haplotype H showed one base change at the reverse primer *groEL*-1R binding site, where the first nucleotide was C instead of the A found in the other Lso haplotypes. Surprisingly, despite this mismatch, the reverse primer *groEL*-1R was still able to produce the expected 634 bp PCR product together with the forward primer *groEL*-1F. This SNP, a silent mutation in the leucine codon 341, was confirmed by performing a new set of PCR and sequencing reactions with alternative primers *groEL*-F (5'-TAGCAACTATTTCTGCTAACGG-3') and *groEL*-R (5'-CTCTTTAAGCTTATCACGGTCA-3').

#### Phylogeny of the Lso haplotypes

Based on the 50S ribosomal protein gene region *rplJ/rpL* and on the concatenated sequences of the

seven MLST gene fragments, haplotype H appears distinct from the previously characterized haplotypes of Lso, which cluster together in phylogenetic analyses (Figs. 3, 4 and 5). The haplotype B that was seen as an outlier in previous analyses (Haapalainen et al. 2018; Hajri et al. 2017, 2019) appears to cluster together with the recently characterized haplotypes F and G. Separate from them, the haplotypes A, U, D and E form a cluster of closely related haplotypes. The genetic diversity within the haplotypes D and E was recently studied in detail (Hajri et al. 2019). As the haplotype A was originally found west of the mountain divide of North America (Nelson et al. 2011), its close relationship to three haplotypes found in Europe suggests that the ancestor of this cluster was distributed over an area including both north-western America and parts of Eurasia, connected by the Beringian land bridge. Haplotype C is closely related to this cluster. The new outlier is the haplotype H, which seems to represent an early divergent lineage of Lso. The three main branches of the Lso phylogeny are best visualized in the maximum likelihood tree based on the MLST sequences (Fig. 5). Thus, including the newly discovered haplotypes F (Swisher Grimm and Garczynski 2019), G (Mauck et al. 2019) and H in the phylogenetic analyses not only widens the family tree but also sharpens the picture of the relationships between the other haplotypes.

#### DNA sequences

The DNA sequences of Lso haplotype H 16S ribosomal RNA gene, 16S–23S intergenic region, 50S ribosomal protein gene region *rplJ/rplL* and the alleles of the seven MLST gene fragments of *adk*, *atpA*, *fbpA*, *ftsZ*, *glyA*, *groEL* and *gyrB* were deposited at GenBank with accession numbers from MK800158 to MK800169.

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#### Compliance with ethical standards

**Conflict of interest** All the authors declare that they have no conflicts of interest.

**Ethical approval** This article does not contain any studies with human participants or animals performed by any of the authors.

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