

Distribution of Two Distinct Genotypes of Citrus Greening Organism in the Ryukyu Islands of Japan

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Abstract

The Asian type “*Candidatus Liberibacter asiaticus*” (Las, citrus greening organism) is severely damaging citrus production in Asia including Japan. Our previous study suggested that the bacteriophage-type DNA polymerase region (DNA pol) would be useful for molecular differentiation in different Southeast Asian Las isolates. Moreover, Las isolates originated from most of the Southeast Asian regions harbor the DNA pol gene, whereas Japanese ones lack this region. These preliminary findings lead us to a hypothesis that all Japanese isolates lack DNA pol. To try this hypothesis, we collected citrus leaf samples infected with Las throughout the Ryukyu Islands, and examined them by a duplex PCR that could simultaneously amplify two DNA fragments of DNA pol and *nusG-rp/K* operon of Las. The duplex PCR was applied to the collection of 65 Las isolates. Both DNA pol and *nusG-rp/K* operon were successfully amplified from nine isolates, whereas only *nusG-rp/K* operon was amplified with the other 56 isolates. These nine isolates with DNA pol originated from the Hateruma, Irabu, Kohama, Miyako, Tarama, and Yonaguni Islands, which are geographically close to Taiwan. The nucleotide sequence of DNA pol of these nine isolates was identical, and was also the same as four Taiwanese isolates reported previously. These results suggest that Japanese Las isolates comprise at least two distinct genotypes, and the genotype that had DNA pol is highly homogeneous.

Discipline: Plant disease (Plant pathology)

Additional key words: citrus greening disease, *Diaphorina citri*, duplex PCR, Huanglongbing

Introduction

Citrus greening (Huanglongbing) is one of the most devastating citrus diseases in many parts of the world (Asia, North America, South America, and Africa). Citrus greening is a threat to the citrus industry⁴. The causal agents of this disease (citrus greening organism) are *Candidatus Liberibacter* spp., which are gram negative, phloem-limited bacteria that cannot be cultured. The pathogens are mainly transmitted by the psyllids *Trioza*

erytrae (*T. erytrae*) in Africa¹ and *Diaphorina citri* (*D. citri*) in Asia, North America and South America⁵. Compared with *D. citri*, *T. erytrae* is found in cooler areas and at higher altitudes. *D. citri* is more widely spread in warmer lowlands in tropical and subtropical areas. Contaminated plant materials used for propagation of nursery plants also transmit these pathogens. Three *Ca. Liberibacter* spp. that act as pathogens have been identified: *Ca. Liberibacter asiaticus* (Las), *Ca. Liberibacter africanus*⁹ and *Ca. Liberibacter americanus*³.

Infected citrus trees show various symptoms such

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as yellowing and blotchy mottling on their leaves and fruit. Infected trees and branches suffer heavy leaf drop followed by out-of-season flushing and blossoming, and in severe cases, dieback. However, none of the symptoms are specific to citrus greening². In Japan, this disease was found for the first time on Iriomote Island in 1988¹². Subsequently, it was found on Okinawa Main Island in 1994¹⁰, Yoron Island in 2002⁷, and Okinoerabu, Tokunoshima and Kikai Islands in 2003¹⁵. The incidence survey in Okinawa Prefecture revealed that Las was found throughout the Okinawa Islands excluding North and South Daito Islands¹³. The transmission vector, *D. citri* was apparently distributed throughout the Ryukyu Islands, and Las positive psyllids were found in most of these islands¹⁴. These observations indicate that the disease is spreading northward gradually in these areas. North of these islands, there is a main island, Kyushu, which has the main citrus production areas of this region, and northbound dispersion of the disease poses a great threat to citrus cultivation there.

Assessment of genetic diversity provides a framework for understanding the taxonomy, population structure and dynamics of phyto bacteria. It also provides a key for devising sensitive, specific and rapid methods for detecting the pathogen, diagnosing plant disease, and managing disease risk¹¹. Methods to distinguish Las isolates are fundamental for ecologic and epidemiologic studies. We reported genetic diversity of Las in Southeast Asia previously¹⁶. The degree of genetic diversity was different in each gene region of Las, and the most diverse gene region was the bacteriophage-type DNA polymerase region (DNA pol). The Ryukyu Islands of Japan are located near Taiwan, and the closest island, Yonaguni, is approximately 100 km apart from Taiwan. This close geographical situation has made us postulate that genotypes of Las in the Ryukyu Islands are similar to those of Taiwan.

In this paper, we determined the Las genotypic distribution in the Ryukyu Islands of Japan using the duplex PCR developed in this study. Contrary to the initial assumption, the results showed that there are two distinct genotypes in the population structure of Las in the Ryukyu Islands of Japan. One genotype occurs also in Taiwan and another genotype is unique to Japan. The results also indicated that one of the two genotypes may constitute an absolutely homogeneous population related to Taiwanese isolates. We also discuss the relationship between the genotypic composition and geographical origin.

Materials and methods

1. Sample collection and preparation

Leaf samples were collected from infected citrus trees

in different orchards throughout the Ryukyu Islands of Japan (Table 1). Total DNA was extracted from about 0.2 g of the leaf midrib from the infected citrus tree using a DNeasy Plant Mini kit (Qiagen, Valencia, USA) according to the manufacturer's instructions. The extracted DNA was suspended in 20–200 µl of TE buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8.0).

2. Duplex polymerase chain reaction

Genotypes of Las isolates were determined by the duplex PCR based on DNA pol and *nusG-rplK* operon. DNA pol was chosen as a variable region to differentiate isolates, while *nusG-rplK* operon was selected as a conserved region among isolates to verify the presence of Las. One primer set of DNA pol was previously reported as GODNPFW1 (5'-TCCTGAGAATTACACACAAAC-3') and GODNPRV1 (5'-TCTAAGTCTATCCTGTAACCC-3')¹⁶. The other primer set of *nusG-rplK* operon was previously reported as MHO353 (5'-GTGTCTCTGATGGTCCGTTTGCTTCTTTTA-3') and MHO354 (5'-GAACCTTCCACCATACGCATAGCCCTTCA-3')⁸. Genotyping by the duplex PCR using both of these primer sets was assessed by the following amplicons: 988 bp and 627 bp for DNA pol and *nusG-rplK* operon, respectively. The PCR reactions were performed using an iCycler thermal cycler (Bio-Rad Laboratories Hercules, CA, USA) in 20 µl reaction mixture volumes containing 2 µl of DNA template, 1 µM of each primer, 200 µM of dNTP mixture, 1 × PCR buffer, and 0.5 units of *Ex Taq* DNA polymerase Hot Start Version (TaKaRa, Japan). The thermal cycling conditions were as follows: initial denaturing at 94°C for 2 min; 40 cycles of denaturing at 94°C for 30 s, annealing at 50°C for 30 s, extension at 72°C for 1 min; and a final extension step at 72°C for 5 min. The amplified PCR products were separated by electrophoresis in 1.5% agarose gels. Gels were stained with 0.5 µg/ml ethidium bromide, and visualized under a UV transilluminator.

3. Sequence analysis of bacteriophage-type DNA polymerase region

Among the analyzed isolates in this study, nine isolates that possessed the DNA pol were selected, and the sequence of the DNA pol and flanking regions were determined as described previously¹⁶ and compared among isolates. Sequence data were assembled using BioEdit version 5.0.9⁶.

Results

Duplex PCR results would show that the isolates with DNA pol produce two DNA bands of DNA pol and *nusG-rplK* operon after agarose gel electrophoresis, whereas the

Table 1. Isolates of *Candidatus Liberibacter asiaticus* used in this study

	Isolate	Code	Location	Year of collection
1) Kikai Island	Kikai-130	Hm1	Oasato, Kikai, Kagoshima	2006
	Kikai-145	Hm2	Oasato, Kikai, Kagoshima	2006
	Kikai-147	Hm3	Oasato, Kikai, Kagoshima	2006
	Kikai-269	Hm4	Oasato, Kikai, Kagoshima	2007
	Kikai-301	Hm5	Oasato, Kikai, Kagoshima	2007
	Kikai-318	Hm6	Oasato, Kikai, Kagoshima	2007
	Kikai-323	Hm7	Oasato, Kikai, Kagoshima	2007
2) Tokunoshima Island	Toku-225	Hm8	Kinen, Isen, Tokunoshima, Kagoshima	2006
	Toku-228	Hm9	Kinen, Isen, Tokunoshima, Kagoshima	2006
	Toku-229	Hm10	Kinen, Isen, Tokunoshima, Kagoshima	2006
	Toku-230	Hm11	Kinen, Isen, Tokunoshima, Kagoshima	2006
	Toku-231	Hm12	Kinen, Isen, Tokunoshima, Kagoshima	2006
	Toku-232	Hm13	Kinen, Isen, Tokunoshima, Kagoshima	2006
	Toku-233	Hm14	Kinen, Isen, Tokunoshima, Kagoshima	2006
	Toku-234	Hm15	Nishi-metegu, Isen, Tokunoshima, Kagoshima	2006
	Toku-235	Hm16	Higashi-metegu, Isen, Tokunoshima, Kagoshima	2006
	Toku-236	Hm17	Higashi-metegu, Isen, Tokunoshima, Kagoshima	2006
	Toku-237	Hm18	Higashi-metegu, Isen, Tokunoshima, Kagoshima	2006
	Toku-238	Hm19	Higashi-metegu, Isen, Tokunoshima, Kagoshima	2006
	Toku-239	Hm20	Higashi-metegu, Isen, Tokunoshima, Kagoshima	2006
	Toku-240	Hm21	Saben, Isen, Tokunoshima, Kagoshima	2006
	Toku-241	Hm22	Saben, Isen, Tokunoshima, Kagoshima	2006
	Toku-244	Hm23	Saben, Isen, Tokunoshima, Kagoshima	2006
	3) Yoron Island	Yoron-57	H1	Yoron, Kagoshima
Yoron-83		H2	Yoron, Kagoshima	2002
Yoron-121		H3	Yoron, Kagoshima	2002
Yoron-127		H4	Yoron, Kagoshima	2002
4) Okinawa Main Island	KIN-1	Iw2	Kin, Okinawa	1994
	Honto-4	Iw5	Okinawa	2005
	KIN-3	Ns1	Kin, Okinawa	2007
	Ishi-2	Ns2	Ishikawa, Uruma, Okinawa	2007
	Nago-Nc-1	K14	Nago, Okinawa	2007
	Nago-4	K15	Nago, Okinawa	2007
	MotobuB-1	K16	Motobu, Okinawa	2007
	Kin2-1	K17	Kin, Okinawa	2007
	Nakijin-5	K18	Nakijin, Okinawa	2007

Table 1. (continued)

	Isolate	Code	Location	Year of collection
	HigashiA-3	K19	Higashi, Okinawa	2007
	OgimiA-3	K20	Ogimi, Okinawa	2007
	Uruma1-1	K21	Gushikawa, Uruma, Okinawa	2007
	UrumaKA-5	K22	Katuren, Uruma, Okinawa	2007
	A-17	K23	Okinawa, Okinawa	2007
	A2-12	K24	Okinawa, Okinawa	2007
	B-8	K25	Okinawa, Okinawa	2007
	A-11	K26	Tomigusuku, Okinawa	2007
	C-3	K27	Itoman, Okinawa	2007
	A-3	K28	Naha, Okinawa	2007
	Hae-5	K29	Haebaru, Okinawa	2007
	KO-7	K30	Kochinda, Yaese, Okinawa	2007
5) Miyako Island	Miyako-13	Iw4	Miyakojima, Okinawa	2007
	S-2-4	K1	Shimoji, Miyakojima, Okinawa	2007
	H-3	K3	Hirara, Miyakojima, Okinawa	2007
	U-4	K4	Ueno, Miyakojima, Okinawa	2007
6) Irabu Island	I-1	K2	Irabu, Miyakojima, Okinawa	2007
7) Ishigaki Island	Ishi-1	Iw3	Ishigaki, Okinawa	2005
	Ishigaki-16	Iw6	Ishigaki, Okinawa	2007
	Hirakubo-5	K5	Hirakubo, Ishigaki, Okinawa	2007
	Hirano-4	K6	Hirano, Ishigaki, Okinawa	2007
	Kawahara-2	K7	Kawahara, Ishigaki, Okinawa	2007
	Hirae-1	K8	Hirae, Ishigaki, Okinawa	2007
8) Hateruma Island	Hateruma-1	K9	Hateruma, Taketomi, Okinawa	2007
9) Kohama Island	Kohama-4	K10	Kohama, Taketomi, Okinawa	2007
10) Yonaguni Island	Higawa-1	K11	Higawa, Yonaguni, Okinawa	2007
11) Tarama Island	Tarama-12	K12	Tarama, Okinawa	2007
12) Iheya Island	Iheya-2	K13	Iheya, Okinawa	2007
13) Iriomote Island	OK-901	Iw1	Iriomote, Taketomi, Okinawa	1988

isolates without DNA pol produce only one DNA band of *nusG-rplK* operon. We named these two genotypes as genotype I and II, respectively. Both genotype I and II were observed among 65 isolates analyzed in this study (Figs. 1 & 2). For example, the result of Fig. 2E showed that the isolates of Hateruma-1, Kohama-4, Higawa-1, and Tarama-12 were classified as genotype I, while the isolate of Iheya-2 as genotype II. Nine isolates belonged to geno-

type I, and these isolates were collected from Hateruma, Irabu, Kohama, Miyako, Tarama, and Yonaguni Islands, which are relatively close to Taiwan (Fig. 1, 2D & 2E). The other 56 isolates belonged to genotype II, which were collected from Iheya, Iriomote, Ishigaki, Kikai, Okinawa Main, Tokunoshima, and Yoron Islands (Figs. 1 & 2A to 2C). The relationship between the year of collection and genotype could not be observed. For example, in spite of

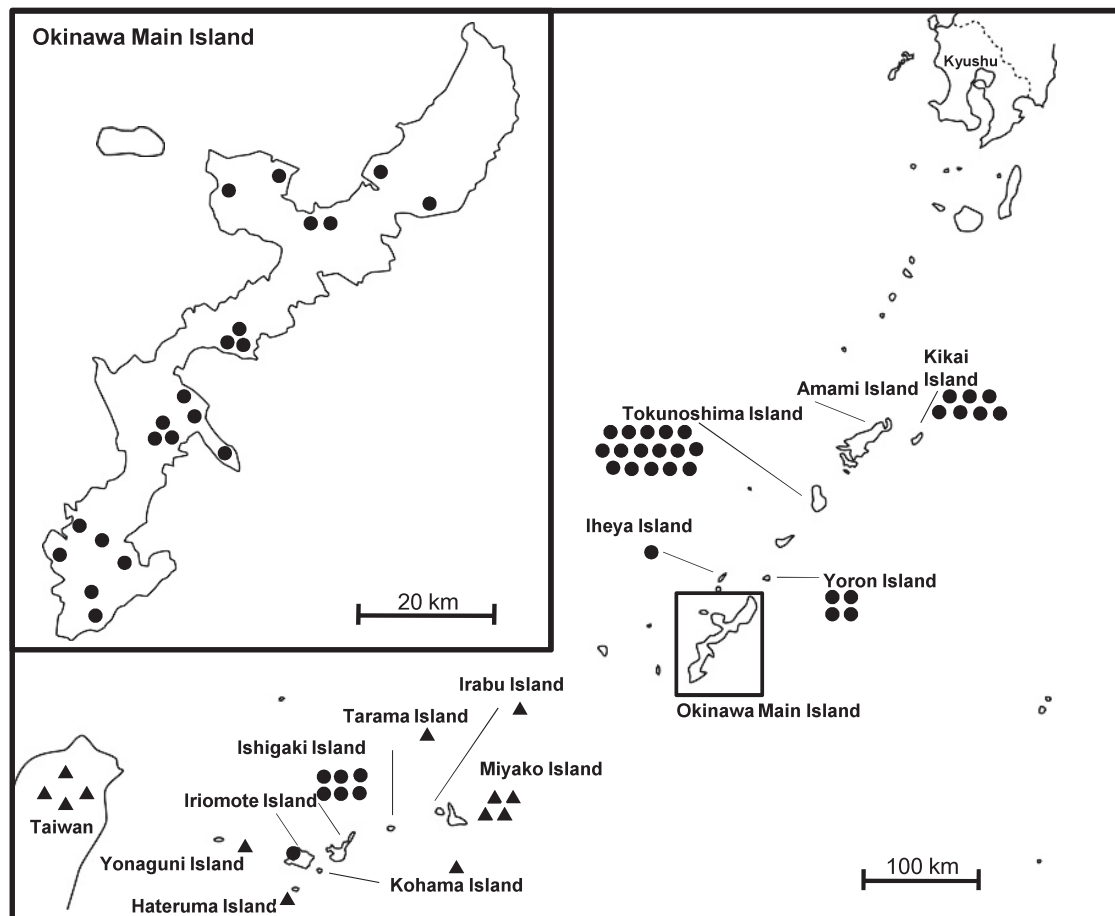


Fig. 1. Map of the Ryukyu Islands

An enlarged map of Okinawa Main Island is shown in the inset. The black triangles and circles show genotype I and II, respectively. Four Taiwanese isolates of genotype I are also shown.

the difference in year of collection, the isolates collected from Kikai and Okinawa Main Islands constituted the same genotype II.

The nine isolates of genotype I were sequenced around DNA pol. All the isolates sequenced in this study were 3,609 nucleotides long, and were identical among isolates. Moreover, sequence alignment was carried out using the sequences obtained in this study and the previously reported sequences of the Southeast Asian isolates¹⁶. Interestingly, the nine Japanese isolates sequenced in this study were identical with four Taiwanese isolates (TW2, TW3, TW5, and TW6, data not shown).

Discussion

Genetic diversity in bacteria can be assessed by examining specific restriction enzyme sites, repetitive elements, amplicons produced by random primers, or genome sequences¹¹. Our previous study suggests that the polymorphic DNA pol may be useful for molecular differentiation among the Southeast Asian Las isolates¹⁶.

In this study, we used a duplex PCR of variable DNA pol and conserved *nusG-rpIK* operon to differentiate among Las isolates of the Ryukyu Islands, and examined the genotypic distribution. Using this duplex PCR, the Las isolates collected from the Ryukyu Islands were clearly differentiated to two distinct genotypes. The genotype I in this study includes the group A, B and C that had been previously reported¹⁶, as well as the India-Poona isolate⁹. The genotype II consists of the Japanese isolates that lack DNA pol¹⁶. The method developed in this study could differentiate clearly among Las, especially genotype II.

Genotypic composition of Las isolates was different on each island. Especially, many Las isolates were collected from Okinawa Main and Tokunoshima Islands. These islands harbored only genotype II which is unique to Japan. These islands are the northern boundary of occurrence of greening from Southeast Asia. Dominance of unique isolates in the northern boundary region suggests that invasion of various foreign Las from southern Asian countries to these areas is not very frequent¹⁶. The Miyako-Yaemama area excluding Ishigaki and Iriomote

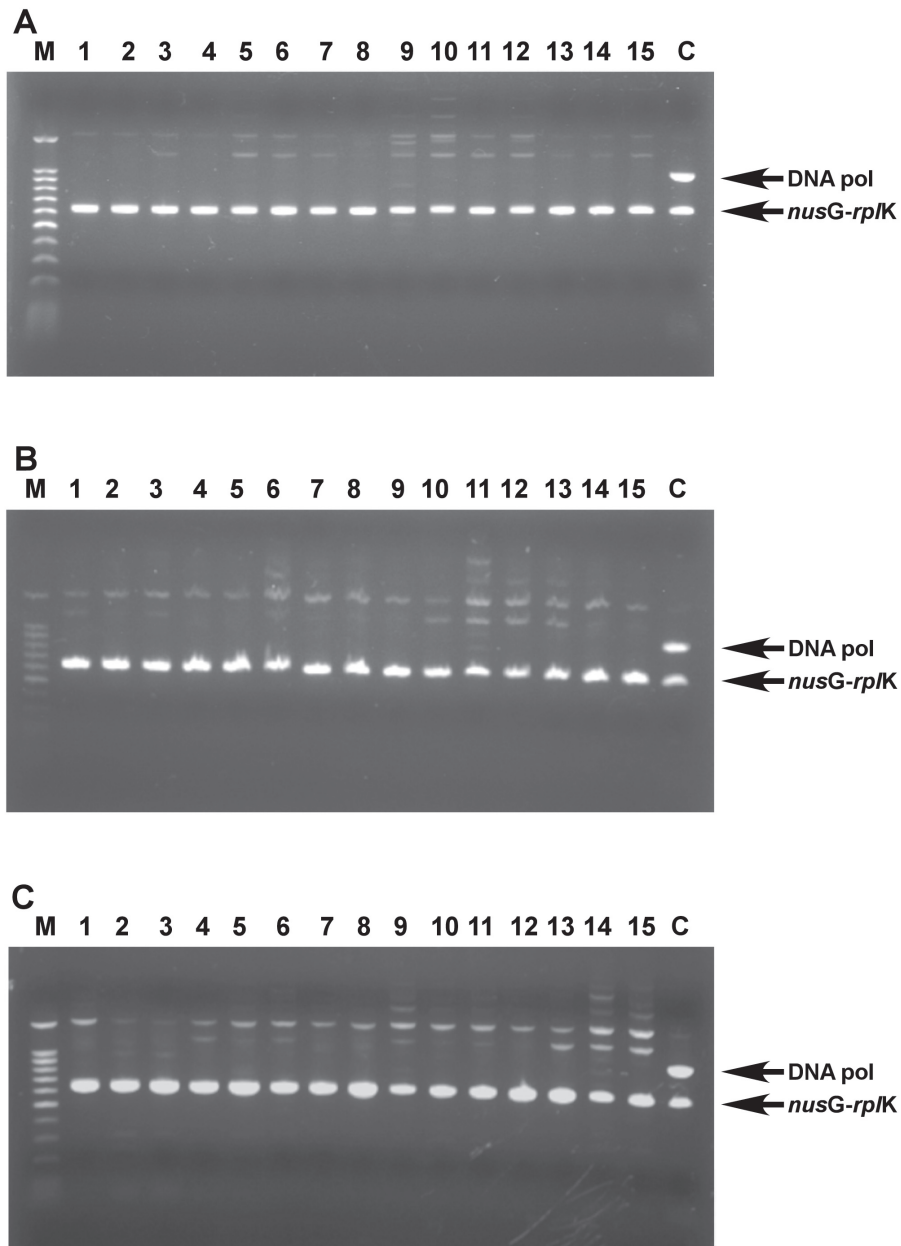


Fig. 2. Results of genotypes of Las isolates determined by the two primer pairs of the bacteriophage-type DNA polymerase region (*DNA pol*) and *nusG-rpIK* operon

Arrows indicate that PCR amplicons of the *DNA pol* (988 bp, upper) and *nusG-rpIK* operon (627 bp, below). (A) Lane M, 100 bp Ladder (Promega). Lanes 1-7, Kikai-130, 145, 147, 269, 301, 318, and 323, respectively. Lanes 8-15, Toku-225, 228, 229, 230, 231, 232, 233 and 234, respectively. Lane C, TW2 (positive control). (B) Lane M, 100 bp Ladder. Lanes 1-8, Toku-235, 236, 237, 238, 239, 240, 241, and 244, respectively. Lanes 9-12, Yoron-57, 83, 121, and 127, respectively. Lanes 13-15, KIN-1, Honto-4, and KIN-3, respectively. Lane C, TW2. (C) Lane M, 100 bp Ladder. Lanes 1-15, Ishi-2, Nago-Nc-1, Nago-4, MotobuB-1, Kin2-1, Nakijin-5, HigashiA-3, OgimiA-3, Uruma1-1, UrumaKA-5, A-17, A2-12, B-8, A-11, and C-3, respectively. Lane C, TW2. (D) Lane M, 100 bp Ladder. Lanes 1-15, A-3, Hae-5, KO-7, Miyako-13, S-2-4, I-1, H-3, U-4, Ishi-1, Ishigaki-16, Hirakubo-5, Hirano-4, Kawahara-2, Hirae-1, and OK-901, respectively. Lane C, TW2. (E) Lane M, 100 bp Ladder. Lanes 1-5, Hateruma-1, Kohama-4, Higawa-1, Tarama-12, and Iheya-2, respectively. Lane C, TW2.

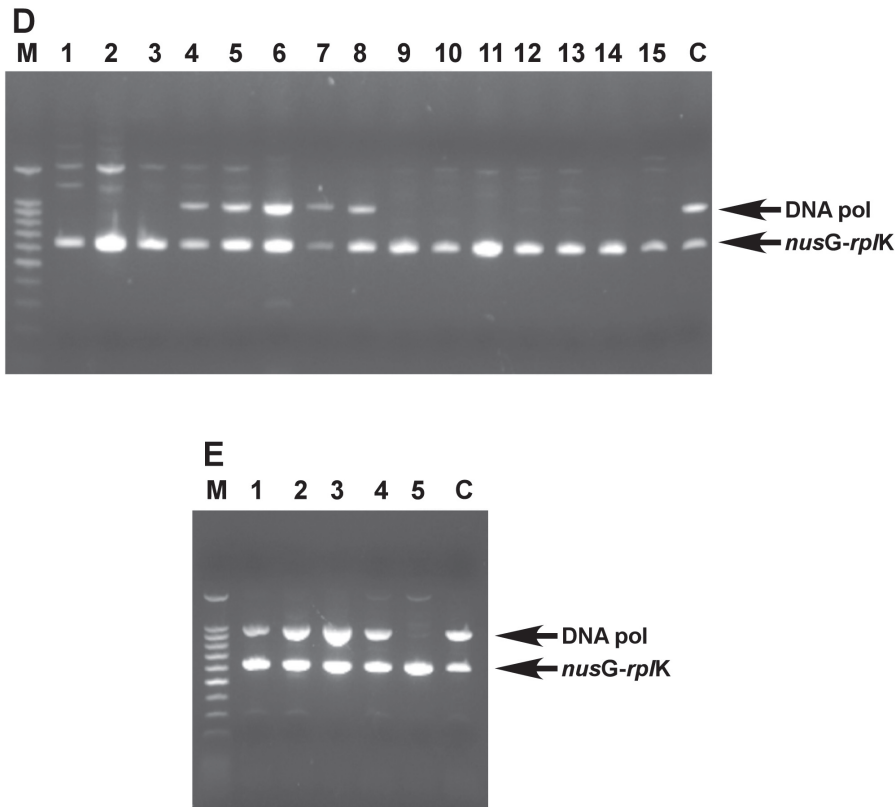


Fig. 2. (continued)

Islands in this survey apparently harbored only genotype I. However, the number of the isolates collected on these islands is insufficient to discuss the detailed within-island population structure. Another research survey would be needed to answer this question in the future.

All the genotype I isolates in the Ryukyu Islands sequenced in this study had identical DNA pol nucleotide sequences, moreover, these isolates were identical to four Taiwanese isolates. These results suggested that a part of the Las population in the Ryukyu Islands may share ancestral history with the Taiwanese population.

Citrus greening diseases are distributed mainly in tropical regions. Greening in Japan had been found first in the southernmost island of Iriomote, which is very close to Taiwan. The distribution of genotype I both in the Miyako-Yaeyama area and Taiwan suggests that some Las isolates may have been introduced from Taiwan to this region. On the other hand, the origin of genotype II, which is unique to Japan, remains unknown.

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