Complete genome analysis of a novel recombinant isolate of pepper veinal mottle virus from mainland China

Songbai Zhang1,3, Zibing Zhao3, Limin Zheng1, Jian Liu2, Jing Peng1, Fei Yan4, Fan Li5, Yan Xie6, Zhaobang Cheng7, Xuguo Zhou1, Deyong Zhang1,2* and Yong Liu1,2*

Abstract

Background: Pepper veinal mottle virus (PVMV) was well established in Africa, and also reported infecting pepper (Capsicum annuum L) in Taiwan and India. However, there is not available of PVMV in mainland China. Here, the first complete genome sequence of PVMV isolated from pepper in mainland China was reported.

Finding: The complete genomic sequence of isolate PVMV-HN isolated from pepper in mainland China is reported in this study. The genome of PVMV-HN is 9793 nucleotides (nt) excluding the poly (A) tail, shares 98-99 % nucleotide sequence identity with those two PVMV isolates from Ghana and Taiwan. Recombinant analysis showed that PVMV-HN probably represents a novel recombinant of PVMV. The phylogenetic relationship of PVMV-HN isolate to other PVMV isolates and other potyviruses based on genome or polyprotein sequence level and CP gene level, was also analyzed in this study.

Conclusion: The current study will help to understand phylogenetic relationship of isolate PVMV-HN.

Background

Five Potyvirus spp. have been reported to infect pepper crops (Capsicus spp.) [1], of these, Pepper veinal mottle virus (PVMV), has been considered to be a major constraint to pepper production for yield and fruit quality reduction [2]. The virion of PVMV is flexuous filaments, enveloped a single molecule of linear, positive-sense, single-stranded ribonucleic acid (ssRNA), about 9.7-10 Kb in size, which has a poly (A) tract at the 3' end, and its 5' end covalently linked to the virus-encoded VPg protein [3, 4]. PVMV could form viral inclusion bodies and “pinwheels” in the cytoplasm of infected cells, which similar with other potyviruses [5], however, its serologically unrelated to other several pepper potyviruses, such as Tobacco etch virus (TEV) [6], Potato virus Y (PVY) [7] and Chili veinal mottle virus (ChiVMV) [8], and so on are extensively infecting pepper crops in China.

The first complete genome RNA sequence of PVMV was originally isolated from Capsicus frutescens in Ghana [9], and subsequently the other complete genome RNA sequence isolated from Taiwan have been deposited in GenBank (No. FJ617225). This virus is well documented of infected pepper crops in several countries in Africa [10, 11], and the phylogenetic relationship among different isolates with CP gene level has been extensively studied [9]. In Asia, this virus was reported infecting pepper crops in India and Taiwan [12]. However, as our best knowledge, there is no complete genome RNA sequence of PVMV infecting pepper crops in mainland China so far. In this study, the full genome sequence of a PVMV isolated from pepper (Capsicum annuum) in Hunan province, China was determined and compared to those of various PVMV isolates and other potyviruses.

* Correspondence: dyzhang78@163.com; haosi1iu@163.com
1 Key Laboratory of Pest Management of Horticultural Crop of Hunan Province, Hunan Plant Protection Institute, Hunan Academy of Agricultural Science, Changsha 410125, China
Full list of author information is available at the end of the article

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**Viral material and sequence analysis**

The viral symptoms of the pepper sample display mottle, yellowing and malformation on leaf, this pepper plant was sampled in a survey of pepper diseases in Hunan province, China in 2014. Total RNA infected pepper sample was extracted by an RNeasy mini kit (QIAGEN, Germany). Full-length cDNA was synthesized by using an oligo(dT)18 primer and Transcript® All-in-One First-Strand cDNA Synthesis SuperMix for PCR Kit (TransGen Biotech, Beijing, China) according to manufacturer’s introduction. The complete genome sequence of the isolate PVMV-HN was amplified from eight overlapping fragments using specific primers (Additional file 1: Table S1), and the 5’- and 3’-terminal ends of the genome was reconfirmed by 5’-RACE and 3’-RACE, respectively, as described previously [9]. PCR products were cloned into pEASY-T5 Vector (TransGen, Beijing, China) and sequenced by Sangon Biotechnology Co., Ltd. (Shanghai, China) using an ABI3730 automated DNA sequencer (Applied Biosystems, USA). Sequences were assembled using DNAMAN version 8 (Lynnon, Quebec, Canada). Sequences analysis and comparison of PVMV-HN to the reference sequences were performed using DNAStar 7.01 package (DNASTAR, Madison, USA). The complete RNA genome or polyprotein sequences and CP sequences of other PVMV isolates and other Potyviruses were downloaded from the National Center for Biotechnology Information (NCBI) database (http://www.ncbi.nlm.nih.gov/). The information of all used sequences was listed in Additional file 1: Table S2. Multiple nucleotide sequence alignments were performed with CLUST W, possible recombination events were evaluated by Recombination Detection Program (RDP3) [13] and a phylogenetic tree was reconstructed by the maximum-likelihood method using MEGA5 [14].

**Sequence properties**

The complete genome of the PVMV-HN isolate comprises 9793 nucleotides (nt), with 194 nt at the 5’ non-translated region (NTR) and 373 nt at the 3’ NTR. Similar as other potyviruses, a large typical open reading frame (ORF) was identified as a polyprotein containing 3074 amino acids (aa) with an AUG start codon and UGA stop codon. PVM-HN shares 99 % nucleotide sequence identity with PVMV isolate of Ghana (DQ645484), and 98 % with PVMV isolate of Taiwan (FJ617225) with genome level, however, only 94 % nucleotide sequence identity with PVMV isolates (GQ918276 and GQ918274) from Mali with polyprotein gene level.

A search for possible recombination event was carried out using RDP3, there were three modules (Maxchi, Chimaera and SiScan) of RDP3 supported possible two recombination events between PVMV-HN with PVMV isolates from Ghana and Taiwan with high confidence \((p < 2.25 \times 10^{-5}, \text{Fig. 1})\). The sites of beginning and ending breakpoint were occurring at 3764 nt and 4146 nt, 4726 nt and 5275 nt, respectively. The potential major and minor parents were the PVMV-p (Ghana isolate) and PVMV-ns1 (Taiwan isolate). The existing two recombination events in PVMV-HN probably hinted it was a novel recombinant.

Phylogenetic analysis of PVMV-HN isolate from China to other potyviruses based on nucleotide sequences of complete genome or polyprotein gene level (Fig. 2a) and CP gene level (Fig. 2b) was built and two unique consensuses trees topology were obtained. The two major clades were distinct existence well supported by bootstrap values associated to the branches of the tree (Fig. 2a). The first clade comprises of PVY, Pepper mottle virus, Potato virus V, and the second one includes PVMV and ChiVMV, shows close relationship of PVMV with ChiVMV, as mentioned by Moury et al. [1]. PVMV group was divided into two subgroups, the Africa subgroup and the East Asia subgroup. It is possible that the diversification of PVMV group was relevant with geography. Phylogenetic analysis of more PVMV isolates from different geographic origins based on nucleotide sequence of CP gene was also built and divided 12 PVMV isolates into two distinct major groups (Fig. 2b), reconfirmed that the diversification of PVMV group was relevant with geography, and PVMV-HN from China clustered in the East Asia subgroup.
Phylogenetic analysis suggested that PVMV-HN likely originate from Taiwan and India as its closest evolution relationship (Fig. 2). Thus the recombination events (Fig. 1) were likely critical for PVMV-HN adaption and epidemic [15]. More sequences of PVMV isolates in China would help to understand phylogenetic relationship of PVMV in mainland China with that in other geography.

This study provided the first complete genome sequence about PVMV from mainland China and revealed it is probably a novel recombinant.
Nucleotide sequence accession number
The complete genome sequence of the isolate of PVMV-HN is permanently available in the GenBank database under the accession number KR002568.

Additional file

Additional file 1: Table S1. Primers used in this study. Table S2. Complete genome sequences quoted in this article. (DOC 64 kb)

Competing interests
The authors declare no competing interests.

Authors’ contributions
SZ, ZZ, LZ and JL carried the experimental work, JP, FY, FL, YX and ZC contributed to sampling and sequence analysis, XZ, DZ and YL contributed to recombinant analysis and phylogenetic tree construction. All authors read and approved the final manuscript.

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Author details
1 Key Laboratory of Pest Management of Horticultural Crop of Hunan Province, Hunan Plant Protection Institute, Hunan Academy of Agricultural Science, Changsha 410125, China. 2 Longping Branch, Graduate College, Central South University, Changsha 410125, China. 3 Shaoyang University, Shaoyang 422000, China. 4 Virology and Biotechnology Institute, Zhejiang Academy of Agricultural Sciences, Hangzhou 310021, China. 5 Key Laboratory of Agricultural Biodiversity for Pest Management of China Education Ministry, Yunnan Agricultural University, Kunming 650201, China. 6 Institute of Biotechnology, Zhejiang University, Hangzhou 310058, China. 7 Institute of Plant Protection, Jiangsu Academy of Agricultural Sciences, Nanjing 201014, China.

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