Complete genome sequence of a previously undescribed monopartite begomovirus and betasatellite infecting Malvastrum coromandelianum in Cambodia

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Received: 28 June 2020 / Accepted: 12 January 2021 / Published online: 3 April 2021
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Abstract
A previously undescribed monopartite begomovirus was identified in Kampot province, Cambodia, in Malvastrum coromandelianum plants exhibiting yellow vein symptoms characteristic of begomovirus infections. The apparently full-length viral component was cloned and sequenced following enrichment of circular DNA by rolling-circle amplification and restriction enzyme digestion. The genome of the virus was 2737 nucleotides in length (KP188831) and exhibited an organization like that of other monopartite begomoviruses, sharing the highest nucleotide sequence similarity (87.7% identity) with ageratum yellow vein virus (AM940137). A satellite molecule was amplified from total DNA by PCR amplification, using the betasatellite-specific primer pair β01/β02. The satellite molecule (1346 nt, KP188832) had structural characteristics like those of other betasatellites associated with begomoviruses and shared the highest nucleotide sequence similarity (84.8% identity) with malvastrum yellow vein betasatellite (MN205547). According to the criteria established for species demarcation for classification of begomoviruses (family Geminiviridae) and betasatellites (family Tolecsatellitidae), respectively, the virus isolate from M. coromandelianum in Cambodia is a previously undescribed novel monopartite begomovirus, for which the name “malvastrum yellow vein Cambodia virus” (MaYVCV) is proposed, and the betasatellite is a previously undescribed novel betasatellite, for which the name “malvastrum yellow vein Cambodia betasatellite” (MaYVKHB) is proposed.

Handle Editor: Jesús Navas-Castillo.

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Plant viruses of the family Geminiviridae have a circular, single-stranded DNA genome encapsidated in a twinned icosahedral particle. They are important global pathogens that cause serious yield losses in many crops. The family includes nine genera, namely Becurtovirus, Begomovirus, Capulavirus, Curtovirus, Eragrovirus, Grablovirus, Mastrevirus, Topocuvirus, and Turncurtovirus, based on genome organization, insect vector, and host range [1, 2]. The genus Begomovirus (family Geminiviridae) is the largest group within the family, comprising at least 424 species recognized by the International Committee on Taxonomy of Viruses (ICTV) (http://www.ictvonline.org/virusTaxonomy.asp), and economically, it is the most important, causing yield losses in many crops, including cassava, cotton, and tomato. Begomoviruses are widely distributed in tropical and subtropical regions of the world and are transmitted exclusively by the whitefly Bemisia tabaci (Gennadius). They can have either a monopartite (DNA-A) or bipartite (DNA-A and DNA-B) genome configurations, depending on the number of genome components they possess [3]. The begomoviral DNA-A or DNA-B component is ~2.6-2.8 kb in size, while the monopartite genome is about 2.8 kb in size. The DNA-A and monopartite genomes encode mostly functionally comparable viral proteins: AV1 and AV2 on the virion-sense strand and AC1, AC2, AC3 and AC4 on the complementary strand, for replication, control of gene expression and encapsidation. DNA-B encodes BV1 and BV2 for movement of the virus in plant cells [4]. Many satellite DNA molecules of approximately 1.3 kb in size have been discovered associated with begomoviruses. These satellite DNA molecules are classified as either betasatellites (genus Betasatellite, family Tolecsatellitidae), or alphasatellites (family Alphasatellitidae). Some satellite DNA molecules of approximately 0.7 kb...
in size have been discovered associated with begomoviruses and classified as deltasatellites (genus Deltasatellite, family Tolecsatellitidae) [5, 6]. Betasatellites are involved in the development of disease symptoms, a characteristic that is attributed to their function as a suppressor of host plant gene silencing. Begomovirus-betasatellite complexes cause important diseases of vegetables, fiber crops, and ornamental plants and infect many uncultivated wild plant species [7].

The perennial plant species Malvastrum coromandelianum is native to South America but has been introduced and established in tropical and subtropical regions, including Asia. To date, at least 10 begomoviruses have been found first in M. coromandelianum plants [8–15].

In October 2014, three leaf samples, Ca-1, Ca-2 and Ca-3, were collected from three different M. coromandelianum plants exhibiting yellow vein symptoms in Kampot Province, Cambodia (Fig. 1). Total DNA was extracted using an Easy-Pure Plant DNA Kit (TransGen Biotech, Beijing, China). To detect the suspected begomovirus, PCR amplification was carried out using degenerate primers to amplify the coat protein gene (cp) (AV494/CoPR) [16, 17] with an expected protein gene. PCR amplification was conducted using Pure Plant DNA Kit (TransGen Biotech, Beijing, China). To detections of RCA products with the BamHI restriction enzyme nuclease (Fermentas, GlenBurnie, MD, USA). The digestion of RCA products with the BamHI restriction enzyme yielded a ~2.7-kbp DNA fragment. The fragments (n = 3) were gel-purified and ligated into the plasmid vector pGEM-3Z (Promega Co., Madison, WI, USA), which had been digested with BamHI, introduced into Escherichia coli DH5α by transformation, and sequenced (Invitrogen Co., Shanghai, China). Total DNA was used as template for PCR amplification with primers for DNA-B components, betasatellites, and alphasatellites (PBLlv2040/PCRc1 [18], β01/β02 [19], and DNA101/DNA102 [20], respectively).

The amplicons of each sample confirmed the presence of a begomovirus in the symptomatic M. coromandelianum plants based on the closest matches of the sequences in a BLASTn search of the GenBank database with ageratum yellow vein virus [AM940137], with 95.61% nt sequence identity and 100% coverage.

The complete genome of the putative begomovirus was amplified from one sample by rolling-circle amplification (RCA) (TempliPhi kit; GE Healthcare, Buckinghamshire, UK), followed by digestion with BamHI and HindIII endonucleases (Fermentas, GlenBurnie, MD, USA). The digestion of RCA products with the BamHI restriction enzyme

Fig. 1 Malvastrum coromandelianum plant exhibiting yellow vein symptoms

Fig. 2 Phylogenetic tree showing the relationships of the complete genome sequence of MaYVCV to other begomoviral sequences (A) and of MaYVKHB to closely related betasatellites (B). The tree was constructed using the maximum-likelihood method implemented in MEGA 6.0. The bootstrap (>50%) consensus tree was inferred from 1000 iterations. AYVV, ageratum yellow vein virus; BGMV, bean golden mosaic virus; MaBYMV, malvastrum bright yellow mosaic virus; MaLCHV, malvastrum leaf curl Philippines virus; MaLCuV, malvastrum leaf curl virus; MaYMV, malvastrum yellow mosaic Jamaica virus; MaYMV, malvastrum yellow mosaic virus; MaYVCV, malvastrum yellow vein Cambodia virus; MaYVHeV, malvastrum yellow mosaic Helshire virus; MaYVHV, malvastrum yellow vein Honghe virus; MaYVLahV, malvastrum yellow vein Lahore virus; MaYVV, malvastrum yellow vein virus; MaYYnV, malvastrum yellow vein Yunnan virus; MiYLCV, mimosa yellow leaf curl virus; SiYVVV, sida yellow vein Vietnam virus; StLCuV, stachytarpheta leaf curl virus; ThLCPuV, tobacco leaf curl Pusa virus; ToLCaV, tomato leaf curl Java virus; ToLCMiV, tomato leaf curl Mindanao virus; ToLCLV, tomato leaf curl Laos virus; ToLCPatV, tomato leaf curl Patna virus; ALCuCMB, ageratum leaf curl Cameroon betasatellite; AYVB, ageratum yellow vein betasatellite; AYVCNB, ageratum yellow vein China betasatellite; AYVINB, ageratum yellow vein India betasatellite; CLCuBaB1, cotton leaf curl Bangalore betasatellite 1; CLCuMuB, cotton leaf curl Multan betasatellite; EpYVMB, eupatorium yellow vein mosaic betasatellite; EpYVB, eupatorium yellow vein betasatellite; KeLuB, kenaf leaf curl betasatellite; LaYVB, Lindernia anagallis yellow vein betasatellite; MaLCuGub, malvastrum leaf curl Guangdong betasatellite; MaYVB, malvastrum yellow vein betasatellite; MaYVKB, malvastrum yellow vein Cambodia betasatellite; MaYVVnB1, malvastrum yellow vein Yunnan betasatellite 1; MaYVVaB2, malvastrum yellow vein Yunnan betasatellite 2; MaLCuV, malvastrum leaf curl betasatellite; OLCuB, okra leaf curl betasatellite; PaLCuCNB, papaya leaf curl China betasatellite; ToLCB, tomato leaf curl betasatellite; ToLCBDDB, tomato leaf curl Bangladesh betasatellite.
Begomovirus and betasatellite in *Malvastrum coromandelianum*

**Old World begomoviruses**

**New World begomoviruses**
were 99.6%-100% identical to one another. The sequence of Ca-1 was chosen as the representative genome sequence of this begomovirus and deposited in the GenBank database under the accession number KP188831. The sequence had the typical genome organization of a monopartite Old World begomovirus, containing six predicted open reading frames (ORFs).

Based on SDT analysis, the pairwise nt sequence identity values ranged from 69.9 to 87.7% in comparisons of the viral component from this study to those of 21 other closely related begomoviruses available in the GenBank database. The viral component shared the highest nt sequence similarity (87.7% identity) with ageratum yellow vein virus (AYVV, AM940137) and Malvastrum leaf curl Philippines virus (MaLCPHV, KC577540) (87.5% identity). Based on the species demarcation threshold for begomoviruses (91% nt sequence identity) [15], this virus isolated from M. coromandelianum in Cambodia is a previously undescribed begomovirus, for which the name "malvastrum yellow vein Cambodia virus" (MaYVCV) is proposed.

Phylogenetic analysis carried out using MEGA [23] (maximum likelihood with bootstrap values >50% and 1000 iterations) between MaYVCV and 21 other begomoviral DNA-A sequences showed that it clustered with AYVV (AM940137), stachytarpheta leaf curl virus (StaLCuV, AJ564743), MaLCPHV (KC577540), and tomato leaf curl Mindanao virus (ToLCMiV, EU487046) to form a unique clade (Fig. 2A).

Recombination analysis was carried out with default settings in Recombination Detection Program (RDP) 4.0, using the GENECOV, Max Chi, RDP, Bootscan, Chimaera, 3Seq, and SiSan algorithms [24], but no recombination events were detected in MaYVCV.

The M. coromandelianum samples were tested by PCR for the presence of DNA-B, betasatellites, and alphasatellites, using the primer pairs PBLlv2040/PCRc1 [18], β01/β02 [19], and DNA101/DNA102 [20], respectively. An amplicon of ~1.4 kb was obtained for samples from three diseased plants using β01/β02, but no amplicon was obtained with PBLlv2040/PCRc1 or DNA101/DNA102, suggesting the presence of an associated betasatellite, but no alphasatellite or DNA-B component.

The betasatellite was 1,346 nt in length (GenBank accession no. KP188832) and had the typical structure of a betasatellite, with a single ORF (βC1) located on the complementary-sense strand, an A-rich region, and a satellite conserved region (SCR) containing a predicted stem-loop structure with the nonanucleotide sequence TAATATTAC [7]. Based on SDT analysis, the pairwise nucleotide sequence identity between the betasatellite and betasatellites available in GenBank was 59.1-84.8%. It shared the highest nt sequence similarity (84.8% identity) with malvastrum yellow vein betasatellite from China (MaYVB, MN205547). Phylogenetic analysis indicated that it clustered with MaYVB (MN205547) to form a unique clade (Fig. 2B). Thus, according to the species demarcation threshold for betasatellites (91% nt sequence identity) [25], this is a previously undescribed betasatellite, for which the name "malvastrum yellow vein Cambodia betasatellite" (MaYVKHB) is proposed.

Alphasatellites infecting malvaceous species have been reported throughout Asia, and also in Burkina Faso, Cameroon, Egypt, Kenya, and Mali [26], and deltasatellites infecting M. coromandelianum plants have been reported in Cuba [5]. However, these DNA satellites were not found in the analyzed samples.

Wild plant hosts of viruses are known to be important reservoirs of begomoviruses and are expected to contribute to viral evolution and to the spread of viruses in cultivated crops. The identification and molecular characterization of viral genomes in wild plants and weeds contributes to our understanding of the genetic diversity, ecology, and evolution of begomoviruses. For example, malvastrum leaf curl virus has been identified infecting M. coromandelianum and was also detected in the cultivated fruit crop plant papaya [27], suggesting the wild host may serve as an important reservoir for the virus. Additional research is required to determine the extent of MaYVCV spread between M. coromandelianum and papaya and other cultivated plant species in Cambodia.
Acknowledgements This study was funded by the Scientific and Technological Assistance Project for Developing Countries, China (KY201402015), the National Natural Science Foundation of China (31871937), Guangdong Basic and Applied Basic Research Foundation (2019A1515012150), the Science and Technology Program of Guangzhou, China (201904010173), and the Special Fund for Scientific Innovation Strategy-Construction of High-Level Academy of Agriculture Science (R2019PY-JX005).

Declarations

Conflict of interest The authors declare that they have no conflict of interest.

Ethical approval This research did not use human participants or other animals.

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