Deinbollia mosaic virus: a novel begomovirus infecting the sapindaceous weed Deinbollia borbonica in Kenya and Tanzania

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ABSTRACT Four isolates of a bipartite begomovirus from naturally infected Deinbollia borbonica plants exhibiting yellow mosaic symptoms in Kenya and Tanzania were molecularly characterised. The DNA-A was most closely related to that of tomato leaf curl Mayotte virus (AM701764; 82%), while the DNA-B shared the highest nucleotide sequence identity with that of East African cassava mosaic virus (AJ704953) at 65%. Based on the current ICTV species demarcation criterion for the genus Begomovirus (≥91% sequence identity for the complete DNA-A), we report the full-length genome sequence of this novel bipartite begomovirus. The results reveal additional diversity and reservoir hosts of begomoviruses in East Africa.

Begomoviruses (family Geminiviridae) are an extremely successful group of emerging viruses infecting cultivated (crop), and non-cultivated (weed) plants from different botanical families [1]. The high degree of genetic variability and species diversity exhibited by begomoviruses in Africa threatens the cultivation of economically important crops such as cassava, tomato and beans [2]. Weed-infecting begomoviruses are transmitted by the polyphagous Bemisia tabaci (Gennadius) (Aleyrodidae) [3] and may act as progenitors of crop-infecting begomoviruses [4] or increase the genetic diversity of the latter by recombination [5]. Despite the critical role that weed-infecting begomoviruses play in the epidemiology of crop diseases, they remain under-studied in Africa.

Deinbollia borbonica Scheff. (Sapindaceae) is a common perennial tropical shrub whose geographical distribution spreads from the coastal belt of Somalia to northern Mozambique. In East Africa, it grows as a weed within mixed cropping farming systems where crops such as cassava, tomato, and beans are grown. Despite D. borbonica existing in Africa for over a century (https://plants.jstor.org/stable/10.5555/al.ap.specimen.m0108980), it was not until in 2012 that a begomovirus naturally infecting D. borbonica was identified in northeastern Tanzania by Ndunguru et al. (unpublished; GenBank accession no. KT799138). Here, we report the full-length genome sequence of this novel bipartite begomovirus infecting D. borbonica in Kenya and Tanzania.

Infected leaf samples showing yellow mosaic symptoms (Supplementary Fig. S1) were collected from D. borbonica in Kenya (sample K1; GPS coordinates 03.31669S, 39.96528E; altitude 12 masl; May 2014) and Tanzania (samples T1, T2, and T3; GPS coordinates 05.10219S, 38.47172E; altitude 202 masl; March 2015). Total DNA was extracted using a ZR Plant/Seed DNA MiniPrep Kit (Zymo Research Corp.). The identity of the plant species was confirmed by PCR amplification and Sanger sequencing of the chloroplast rbcL gene [6, 7]. The virus
(A)

(B)
The genome was enriched by rolling-circle amplification (RCA) using a TempliPhi 100 RCA Kit (GE Healthcare), and sequenced using an Illumina MiSeq system at BecA-ILRI Hub (Nairobi, Kenya). De novo assembly using CLC Genomics Workbench version 7.0.4 (CLC Bio; QIAGEN) generated two consensus contigs identified as DNA-A and DNA-B by BLASTn analysis (http://www.ncbi.nlm.nih.gov/BLAST). The genome components were amplified using abutting primers (DNA-Af, 5'-TTGGGCTCAAGTITTT GACG-3'; DNA-Ar, 5'-TACCGTCAAACCTTGAGGC-3') and DNA-Bf, 5'-TGTCGTATGCGTGCTTTTGG-3'; DNA-Br, 5'-AATAGCCTCAAAAGCAGCGC-3') using Phusion High-Fidelity DNA Polymerase (Thermo Fisher Scientific). The blunt-ended PCR products were cloned into PUC18 and sequenced by the Sanger method using overlapping PCR primers (Supplementary Table S1).

The genome organization of the virus was identical to that of Old World bipartite begomoviruses (Supplementary Table S2). The DNA-A and DNA-B contained a 158-nucleotide (nt)-long common region (CR) with >97% nucleotide sequence identity. Additionally, a unique putative iteron sequence (GAGGGCA), appearing twice as perfect repeats and once as an inverted repeat sequence, was identified, indicating that the cloned DNA-A and DNA-B from each plant sample constituted a cognate pair. The CR also possessed the nonanucleotide sequence (5'-TAATATTAC-3') typical of begomoviruses [8]. Recombination analysis using RDP4 revealed no evidence of significant recombination events.

Pairwise nucleotide sequence identities calculated using Sequence Demarcation Tool (SDT v1.2) [9] revealed that the isolates were >98% identical to each other and <91% identical to currently reported begomoviruses (Supplementary Fig. S2). According to the recently updated ICTV begomovirus species demarcation criteria [10], the isolates qualify to belong to a novel begomovirus species, tentatively named “Deinbollia mosaic virus” (based on the host plant and disease symptom). Phylogenetic reconstruction based on the full-length nucleotide sequences of the DNA-A (Fig. 1A) and DNA-B (Fig. 1B) using GTR + G + I as the best-fit model of evolution in MEGA 6 software [11], clustered DMV isolates together with African begomoviruses. The DNA-A sequence was most similar (82%) to that of tomato leaf curl Mayotte virus (ToLCYT; AM701764), while the DNA-B sequence shared 65% nucleotide sequence identity with that of East African cassava mosaic virus (EACMV: AJ704953). The results reveal additional diversity and reservoir hosts of begomoviruses in East Africa.

**GenBank accession numbers**

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**Compliance with ethical standards**

**Conflict of interest** The authors declare no conflicts of interest.

**Research involving human participants and/or animals** No part of this study was performed with human participants or animals by any of the authors.

**Informed consent** Informed consent was obtained from individual farmers for the samples used in this study.

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