# RESEARCH ARTICLE

# The ever increasing diversity of begomoviruses infecting non-cultivated hosts: new species from *Sida* spp. and *Leonurus sibiricus*, plus two New World alphasatellites

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#### Keywords

Begomovirus; geminivirus: recombination; plant virology; satellite DNA.

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## Abstract

Begomoviruses (whitefly-transmitted, single-stranded DNA plant viruses) are among the most damaging pathogens causing epidemics in economically important crops worldwide. Besides cultivated plants, many weed and wild hosts act as virus reservoirs where recombination may occur, resulting in new species. The aim of this study was to further characterise the diversity of begomoviruses infecting two major weed genera, Sida and Leonurus. Total DNA was extracted from samples collected in the states of Rio Grande do Sul, Paraná and Mato Grosso do Sul during the years 2009–2011. Viral genomes were enriched by rolling circle amplification (RCA), linearised into unit length genomes using various restriction enzymes, cloned and sequenced. A total of 78 clones were obtained: 37 clones from Sida spp. plants and 41 clones from Leonurus sibiricus plants. Sequence analysis indicated the presence of six bipartite begomovirus species and two alphasatellites. In Sida spp. plants we found Sida micrantha mosaic virus (SiMMV), Euphorbia yellow mosaic virus (EuYMV), and three isolates that represent new species, for which the following names are proposed: Sida chlorotic mottle virus (SiCMoV), Sida bright yellow mosaic virus (SiBYMV) and Sida golden yellow spot virus (SiGYSV), an Old World-like begomovirus. L. sibiricus plants had a lower diversity of begomoviruses compared to Sida spp., with only Tomato yellow spot virus (ToYSV) and EuYMV (for the first time detected infecting plants of the genus *Leonurus*) detected. Two satellite DNA molecules were found: Euphorbia yellow mosaic alphasatellite, for the first time detected infecting plants of the genus Sida, and a new alphasatellite associated with ToYSV in *L. sibiricus*. These results constitute further evidence of the high species diversity of begomoviruses in non-cultivated hosts, particularly Sida spp.

#### Introduction

The family *Geminiviridae* comprises plant viruses with one or two genomic components of circular, single-strand DNA (ssDNA) encapsidated in geminate particles (Brown *et al.*, 2012). The family is divided into seven genera (*Becurtovirus, Begomovirus, Curtovirus, Eragrovirus, Mastrevirus, Topocuvirus* and *Turncurtovirus*) according to the type of insect vector, host range, genome organisation and phylogeny (Brown *et al.*, 2012; Varsani *et al.*, 2014). The genus *Begomovirus* is the largest in the family and includes mono- and bipartite viruses transmitted by *Bemisia tabaci* (Hemiptera: Aleyrodidae) to dicotyledonous plants (Brown *et al.*, 2015).

Based on phylogenetic analysis and genomic features, begomoviruses are broadly divided into two groups: Old World (OW; Europe, Africa, Asia and Oceania) and New World (NW; the Americas) (Rybicki, 1994; Padidam *et al.*, 1999; Paximadis *et al.*, 1999). Begomoviruses in the New World are mostly bipartite (DNA-A and DNA-B), except for *Tomato leaf deformation virus* (ToLDeV), an indigenous NW monopartite virus (Melgarejo et al., 2013). The DNA-A contains genes involved in replication, encapsidation of viral progeny and suppression of host defenses (Rojas et al., 2005; Hanley-Bowdoin et al., 2013), and the DNA-B contains genes required for intraand intercellular movement in the plant, host range determination and suppression of host defenses (Rojas et al., 2005; Mahajan et al., 2011; Hanley-Bowdoin et al., 2013; Brustolini et al., 2015). The majority of the begomoviruses that occur in the OW are monopartite, with a genomic organisation similar to the DNA-A of bipartite viruses (Padidam et al., 1996; Mansoor et al., 2003) and the presence of an additional open reading frame (ORF) which partially overlaps the cp gene, named v2 in monopartite viruses or av2 in bipartite viruses. The V2/AV2 protein is involved in viral movement and gene silencing suppression (Rybicki, 1994; Padidam et al., 1996; Glick et al., 2008). Begomoviruses in the OW are generally associated with satellite DNA molecules (Zhou, 2013).

Brazil is a begomovirus diversity hotspot, with reports of their detection dating back to the 1950s (Costa & Bennett, 1950; Costa, 1955). Begomoviruses are limiting factors for common bean and tomato production (Faria *et al.*, 2000; Zerbini *et al.*, 2005), and a large number of new species of tomato-infecting begomoviruses has been identified in the country (Ribeiro *et al.*, 2003; Fernandes *et al.*, 2006; Calegario *et al.*, 2007; Ribeiro *et al.*, 2007; Castillo-Urquiza *et al.*, 2008; Fernandes *et al.*, 2008; Albuquerque *et al.*, 2012). The advent of techniques for the unbiased amplification of circular DNA genomes (specially rolling circle amplification, RCA (Inoue-Nagata *et al.*, 2004), created new possibilities for the discovery of novel begomoviruses, and also of divergent ssDNA viruses (Krenz *et al.*, 2012; Loconsole *et al.*, 2012; Basso *et al.*, 2015).

Non-cultivated species of the families Asteraceae, Caparaceae, Euphorbiaceae, Fabaceae, Labiatae, Malvaceae, Solanaceae and Sterculiaceae have been reported as hosts of many begomoviruses in Brazil and in several other countries in the Americas (Frischmuth et al., 1997; Roye et al., 1997; Faria and Maxwell, 1999; Fernandes et al., 1999; Idris et al., 2003; Jovel et al., 2004; Assunção et al., 2006; Amarakoon et al., 2008; Castillo-Urquiza et al., 2008; Barbosa et al., 2009; Silva et al., 2011; Silva et al., 2012; Tavares et al., 2012; Barreto et al., 2013). There is evidence that some of these begomoviruses from noncultivated hosts can be transmitted to cultivated species by the insect vector and by grafting (Arnaud et al., 2007; Castillo-Urquiza et al., 2007; Cotrim et al., 2007; Silva et al., 2010; Barreto et al., 2013; Rocha et al., 2013; Ramos-Sobrinho et al., 2014), highlighting the need to investigate these plants as reservoirs of viral diversity and as a source of new viruses which may cause diseases in crops.

## Materials and methods

During the years 2009 to 2011, leaf samples from Sida spp. (Malvaceae) and Leonurus sibiricus (Lamiaceae) plants displaying symptoms of yellow mosaic and leaf distortion and/or infestation by B. tabaci were collected in the states of Rio Grande do Sul (n = 27), Paraná (n = 33) and Mato Grosso do Sul (n=10). Total DNA was extracted from press-dried samples as described by Doyle & Doyle (1987). Full-length viral circular genomes were enriched by rolling-circle amplification (RCA) as described by Inoue-Nagata et al. (2004). Unit length genomes were excised with ApaI, BamHI, ClaI, EcoRI, HindIII, KpnI, SacI, SalI or SpeI and ligated into the pBLUESCRIPT-KS+ (pKS+) plasmid vector (Stratagene, San Diego, CA, USA), previously cleaved with the same enzyme. Viral inserts were sequenced commercially (Macrogen Inc., Seoul, South Korea) by primer walking. All genome sequences were organised to begin at the nicking site in the invariant nonanucleotide at the origin of replication (5'-TAATATT//AC-3').

Pairwise sequence comparisons were performed using Sequence Demarcation Tool (SDT) v.1.2 (Muhire et al., 2014) using the MUSCLE alignment option (Edgar, 2004). Multiple sequence alignments were obtained using the MUSCLE algorithm implemented in MEGA6 (Tamura et al., 2013). Phylogenetic analyses were performed with the sequences of the closest begomoviruses determined by BLASTn comparison of the clones generated in this study and the sequences deposited in Genbank, plus some begomovirus sequences of the New and Old World. Phylogenetic trees were constructed using Bayesian inference performed with MrBayes v. 3.0b4 (Ronquist & Huelsenbeck, 2003), with the nucleotide substitution model selected by MrModeltest v. 2.2 (Nylander, 2004) in the Akaike Information Criterion (AIC). The analysis was run for 10 million generations, excluding the first 2 000 000 generations as burn-in. The trees were visualised in FigTree v.1.3.1 (tree.bio.ed.ac.uk/software/figtree/). Recombination analysis was performed with Recombination Detection Program (RDP) v.4.5.1 (Martin et al., 2010) using default settings and a Bonferroni-corrected P-value cutoff of 0.05. The same data set used for the phylogenetic analysis was used for recombination analysis. Only those recombination events detected by more than four of the seven tests implemented in RDP were considered to be reliable.

## **Results and discussion**

A total of 70 samples (43 *Sida* spp. samples, 27 *L. sibiricus* samples) were collected and 61 were preliminarily positive for the presence of a begomovirus, based on the

detection of an approximately 2600-bp band after digestion of the RCA products with restriction enzymes (data not shown). A total of 78 clones were obtained: 37 clones from *Sida* spp. samples and 41 clones from *L. sibiricus* samples (Table 1; these include DNA-A, DNA-B and satellite DNA clones). BLASTn analysis and pairwise sequence comparisons indicated the presence of six begomovirus species and two alphasatellites (Table 1; Figs. S1 and S4).

Pairwise sequence comparisons of cloned genome sequences with those deposited in GenBank indicated that *Sida micrantha mosaic virus* (SiMMV) was the predominant begomovirus infecting *Sida* spp. plants. Out of 15 DNA-A clones obtained from this host, 10 correspond to SiMMV isolates, as well as 19 out of the 21 DNA-B clones (Table 1). The DNA-A sequences share >96.3% nucleotide (nt) identity amongst themselves and 96.3% to 97.2% identity with SiMMV (accession number FN436003), and the DNA-B sequences share >88.0% nt identity amongst themselves and 89.1–93.3% identity with SiMMV (FN436004) (Fig. S1). Bayesian phylogenetic trees based on either the DNA-A or DNA-B placed these isolates in a monophyletic branch together with SiMMV (Fig. 1).

Two DNA-A sequences (BR:Trm531.2:10 and BR:Caa691:10) obtained from *Sida* spp. samples #531 and #691 displayed 97.0% and 97.2% nt sequence identity to *Euphorbia yellow mosaic virus* (EuYMV; FJ619507), respectively (Table 1; Fig. S1). This virus has already been found infecting *Sida santaremnensis* in Minas Gerais (Tavares *et al.*, 2012).

The Sida sample #531 actually had a mixed infection. A virus representing a new species (BR:Trm531.1:10) based on the criteria of <91% nt sequence identity for the DNA-A, recently updated by the Geminiviridae Study Group of the ICTV (Brown et al., 2015), was cloned from this sample, for which the name Sida chlorotic mottle virus (SiCMoV) is proposed. Pairwise sequence comparisons of the DNA-A sequence (2601 nt) with those deposited in GenBank indicated a maximum nt sequence identity of 81.5% with Tomato dwarf leaf virus (ToDfLV, JN564749) (Fig. S1). A DNA-B was detected in the sample but has not yet been cloned. SiCMoV is placed in a monophyletic branch with ToDfLV, Tomato chino La Paz virus (ToChLPV), Tomato leaf deformation virus (ToLDeV), Tomato golden mosaic virus (TGMV), Sida mosaic Bolivia virus 1 (SiM-BoV1) and Abutilon mosaic Bolivia virus (AbMBoV; Fig. 1A). This close relationship is consistent with the pairwise sequence identity analysis and with previously reported data (Márquez-Martín et al., 2011; Medina & Lambertini, 2012; Melgarejo et al., 2013). Interestingly, although no recombination events were detected for this virus, it clustered with viruses of the EuYMV group (including BR:Trm531.2:10 obtained from the same sample) in a cp nt sequence tree, but with SiMBoV1, ToDfLV and *Tomato yellow spot virus* (ToYSV) in a *rep* nt sequence tree (Fig. S2), suggesting a recombinant origin. Mixed infections by different begomoviruses are common in non-cultivated hosts (García-Andrés *et al.*, 2006; Alabi *et al.*, 2008; Monde *et al.*, 2010), facilitating recombination events among distantly related begomoviruses which may contribute to the frequent emergence of new species.

A virus corresponding to a second new species was cloned from the Sida spp. sample #720 (BR:Tac720:10). Nucleotide sequence identity between the common regions (CR) of the DNA-A and DNA-B was 94.7%, and the two components have identical iterons (TGGGG), indicating that they constitute a cognate pair. Both the DNA-A (2692 nt) and the DNA-B (2656 nt) show the highest nt sequence identity with SiMMV (86% and 75.5%, respectively; Fig. S1). The name Sida bright yellow mosaic virus (SiBYMV) is proposed for this new species. Analysis with the RDP4 program detected one strongly supported recombination event in the DNA-A, with SiMMV (FN436003) and an unknown virus as the putative parents (Table 2). Phylogenetic reconstruction based on the DNA-A placed this isolate in a monophyletic branch with SiMMV isolates, occupying a basal position in the clade (Fig. 1A). The recombination event has strong phylogenetic support: the rep nt sequence tree places BR:Tac720:10 in a monophyletic branch (99% posterior probability) with the SiMMV isolates, while the *cp* nt sequence tree places the isolate in a monophyletic branch (99% posterior probability) with Abutilon mosaic Brazil virus (AbMBV, JF694480; Fig. S2). Phylogenetic analysis based on the DNA-B placed this isolate in a monophyletic branch with SiMMV, Tomato rugose mosaic virus (ToRMV, AF291706) and Tomato severe rugose virus (ToSRV, KC004086), occupying a basal position in the clade (Fig. 1B). One recombination event was detected, with BR:Tol1075:11 and Bean dwarf mosaic virus (BDMV, M88180) as the putative parents (Table 2). The recombination event has good phylogenetic support: BR:Tac720:10 groups with different begomoviruses in the mp and nsp nt sequence trees (Fig. S3). Recombination is a common event among geminiviruses (Padidam et al., 1999; Lefeuvre et al., 2009) and contributes greatly to their evolutionary potential and local adaptation (Harrison & Robinson, 1999; Padidam et al., 1999; Berrie et al., 2001; Monci et al., 2002). A number of natural begomovirus recombinants have been responsible for severe diseases and great economic losses in cassava in East Africa (Zhou et al., 1997; Pita et al., 2001), tomatoes in Spain (Monci et al., 2002; García-Andrés et al., 2006; García-Andrés et al., 2007a,b), and cotton and okra in Pakistan (Zhou et al., 1998; Idris and Brown, 2002; Briddon et al., 2014).

|                   | Samuling                |                               |                  |                 |                    | Enzyme <sup>a</sup> |         |           |                 | JenBank       |
|-------------------|-------------------------|-------------------------------|------------------|-----------------|--------------------|---------------------|---------|-----------|-----------------|---------------|
| Sample Code       | Date                    | Location                      | Geographical Coo | rdinates        | Host               | DNA-A               | DNA-B   | Satellite | Isolate Name    | Access Number |
| Sida micrantha    | mosaic virus ( <u>5</u> | SiMMV)                        |                  |                 |                    |                     |         |           |                 |               |
| CF48              | 03/19/09                | Santo Angelo, RS              | S28 22' 54.70''  | W54 18' 17.24'' | Sida spp.          | EcoRI               |         |           | BR:Saa48:09     | KX348158      |
| CF65              | 03/20/09                | São Miguel das Missões, RS    | S28 29' 35.59''  | W54 33' 37.15"  | Sida spp.          | Spel                |         |           | BR:Smm65:09     | KX348159      |
| CF69              | 03/20/09                | Santo Antonio das Missões, RS | S28 29' 42.70''  | W55 25' 18.00'' | Sida spp.          |                     | Spel    |           | BR: Sam 69: 09  | KX348195      |
| CF115             | 03/20/09                | Panambi, RS                   | S28 18' 24.45"   | W53 29' 21.38"  | Sida spp.          | EcoRI               | Spel    |           | BR: Pan 115:09  | KX348160 (A)  |
|                   |                         |                               |                  |                 |                    |                     |         |           |                 | KX348192 (B)  |
| CF547             | 03/24/10                | São Miguel das Missões, RS    | S28 29' 35.59''  | W54 33' 37.15'' | Sida spp.          |                     | Spel    |           | BR:Smm547:10    | KX348196      |
| CF556             | 03/25/10                | Cruz Alta, RS                 | S28 36' 10.89''  | W53 39' 25.34"  | Sida spp.          |                     | Sall    |           | BR:Cra556:10    | KX348194      |
| CF662             | 03/25/10                | Realeza, PR                   | S25 40' 45.00''  | W53 33' 09.00'' | Sida spp.          | EcoRI               | Spel    |           | BR:Rea662:10    | KX348156 (A)  |
|                   |                         |                               |                  |                 |                    |                     |         |           |                 | KX348186 (B)  |
| CF679             | 06/08/10                | Marechal Candido Rondon, PR   | S24 30' 59.50''  | W54 04' 37.20'' | Sida spp.          |                     | Apal    |           | BR:Mcr679:10    | KX348197      |
| CF698             | 06/09/10                | Dourados, MS                  | S22 17' 58.00''  | W54 49' 14.10'' | Sida spp.          |                     | HindIII |           | BR:Dou698:10    | KX348191      |
| CF704             | 06/09/10                | Laguna Carapã, MS             | S22 25' 35.60''  | W55 21' 30.00'' | Sida spp.          |                     | Sall    |           | BR:Lac704:10    | KX348193      |
| CF732             | 06/10/10                | Umuarama, PR                  | S23 50' 12.30''  | W53 17' 23.60'' | Sida spp.          | EcoRI               |         |           | BR:Umu732:10    | KX348164      |
| CF755             | 06/10/10                | Marialva, PR                  | S23 30' 05.00''  | W51 47' 08.00'' | Sida spp.          |                     | Sall    |           | BR:Mar755:10    | KX348198      |
| CF799             | 08/11/10                | São Domingos, PR              | S24 00' 51.00''  | W51 30' 37.00'' | Sida spp.          |                     | Apal    |           | BR:Sad799:10    | KX348190      |
| CF822             | 08/24/10                | Chapada, RS                   | S28 01' 08.80''  | W53 05' 52.31"  | Sida spp.          | Spel                | Spel    |           | BR:Cha822:10    | KX348161 (A)  |
|                   |                         |                               |                  |                 |                    |                     |         |           |                 | KX348199 (B)  |
| CF832             | 08/25/10                | Santo Angelo, RS              | S28 22' 54.70''  | W54 18' 17.23'' | Sida spp.          |                     | Spel    |           | BR:Saa832:10    | KX348202      |
| CF876             | 10/06/10                | São Miguel das Missões, RS    | S28 23' 59.00''  | W54 39' 52.00'' | Sida spp.          | Sacl                | Spel    |           | BR:Smm876:10    | KX348155 (A)  |
|                   |                         |                               |                  |                 |                    |                     |         |           |                 | KX348200 (B)  |
| CF895             | 1 0/0 7/1 0             | Tapatuba, RS                  | S29 03' 57.00''  | W54 43' 53.00'' | Sida spp.          |                     | Sall    |           | BR:Tap895:10    | KX348189      |
| CF926             | 03/15/11                | Chapada, RS                   | S28 01' 08.80''  | W53 05' 52.31"  | Sida spp.          | Spel                | Spel    |           | BR:Cha926:11    | KX348162 (A)  |
|                   |                         |                               |                  |                 |                    |                     |         |           |                 | KX348201 (B)  |
| CF949             | 03/16/11                | São Miguel Das Missões, RS    | S28 29' 35.59''  | W54 33' 37.15"  | Sida spp.          |                     | Sall    |           | BR:Smm949:11    | KX348203      |
| CF1116            | 06/08/11                | Tacuru, RS                    | S23 38' 17.60''  | W54 58' 17.70'' | Sida spp.          |                     | Apal    |           | BR:Tac1116:11   | KX348188      |
| CF1120            | 06/09/11                | Guaíra, PR                    | S24 04' 57.00''  | W54 10' 04.00'' | Sida spp.          |                     | Apal    |           | BR:Gua1120:11   | KX348204      |
| CF1121            | 06/09/11                | Umuarama, PR                  | S23 50' 12.30''  | W53 17' 23.60'' | Sida spp.          | Spel                |         |           | BR:Umu1121:11   | KX348157      |
| CF1128            | 06/09/11                | Janiópolis, PR                | S24 08' 02.00''  | W52 47' 22.00'' | Sida spp.          |                     | BamHI   |           | BR:Jan1128:11   | KX348187      |
| CF1142            | 06/09/11                | Marialva, PR                  | S23 30' 09.40''  | W5147'42.90''   | Sida spp.          | EcoRI               |         |           | BR: Mar 1142:11 | KX348163      |
| Euphorbia yellov  | v mosaic virus          | s (EuYMV)                     |                  |                 |                    |                     |         |           |                 |               |
| CF531             | 03/24/10                | Três de Maio, RS              | S27 45' 39.44''  | W54 15' 42.87'' | Sida spp.          | HindIII             |         |           | BR:Trm531.2:10  | KX348180      |
| CF691             | 06/08/11                | Caarapo, MS                   | S22 26' 42.50''  | W54 49' 49.50'' | Sida spp.          | Sall                |         |           | BR:Caa691:10    | KX348181      |
| CF713             | 06/09/10                | Aral Moreira, MS              | S22 46' 19.70''  | W55 24' 43.90'' | Leonurus sibiricus | Apal                | Kpnl    |           | BR:Arm713:10    | KX348182 (A)  |
|                   |                         |                               |                  |                 |                    |                     |         |           |                 | KX348224 (B)  |
| sida chiorotic m  | ottle virus (Sic        | LIMOV)                        | :                |                 |                    |                     |         |           |                 |               |
| CF531             | 03/24/10                | Três de Maio, RS              | S27 45' 39.44''  | W54 15' 42.87'' | Sida spp.          | Apal                |         |           | BR:Trm531.1:10  | KX348183      |
| Sida bright yellc | w mosaic viru           | us (SiBYMV)                   |                  |                 |                    |                     |         |           |                 |               |
| CF720             | 06/09/10                | Tacuru, MS                    | S23 38′ 17.60′′  | W54 58' 17.70'' | Sida spp.          | Spel                | Clal    |           | BR:Tac720:10    | KX348184 (A)  |
|                   |                         |                               |                  |                 |                    |                     |         |           |                 | KX348225 (B)  |

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|                             | Samuling        |                                  |                         |                      |                  | Enzyme <sup>a</sup> |              |           |                 | GenBank                   |
|-----------------------------|-----------------|----------------------------------|-------------------------|----------------------|------------------|---------------------|--------------|-----------|-----------------|---------------------------|
| Sample Code                 | Date            | Location                         | Geographical Cool       | rdinates             | Host             | DNA-A               | DNA-B        | Satellite | Isolate Name    | Access Number             |
| CF677                       | 06/08/10        | Marechal Candido Rondon, PR      | S24 30' 59.50"          | W54 04' 37.20''      | L. sibiricus     | Apal                |              |           | BR:Mcr677:10    | KX348170                  |
| Sida golden yeli            | low spot virus  | (SiGY SV)                        |                         |                      |                  |                     |              |           |                 |                           |
| CF889                       | 06/10/10        | São Borja, RS                    | S28 57′ 30.00″          | W55 32' 30.00''      | <i>Sida</i> spp. | Apal                |              |           | BR:Sab889:10    | KX348185                  |
| Tomato yellow :             | spot virus (ToY | (SV)                             |                         |                      |                  |                     |              |           |                 |                           |
| CF724                       | 06/10/10        | Guaíra, PR                       | S24 08' 35.30"          | W54 14' 53.20''      | Sida spp.        |                     | Apal         |           | BR:Gua724:10    | KX348226                  |
| CF661                       | 06/07/10        | Realeza, PR                      | S25 40' 45.00"          | W53 33' 09.00''      | L. sibiricus     | Spel                | Kpnl         |           | BR:Rea661:10    | KX348169 (A) KX348214 (B) |
| CF673                       | 06/08/10        | Toledo, PR                       | S24 46' 45.00"          | W53 40' 41.00''      | L. sibiricus     | Apal                | Apal         |           | BR:Tol673:10    | KX348174 (A) KX348223 (B) |
| CF697                       | 06/09/10        | Dourados, MS                     | S22 17' 58.00"          | W54 49' 14.10''      | L. sibiricus     | Apal                | HindIII      |           | BR:Dou697:10    | KX348171 (A) KX348205 (B) |
| CF703                       | 06/09/10        | Laguna Carapã, MS                | S22 25' 35.60"          | W55 21' 30.00"       | L. sibiricus     | Apal                |              |           | BR:Lac703:10    | KX348177                  |
| CF735                       | 06/10/10        | Janiópolis, PR                   | S24 08' 02.00"          | W52 47' 22.00"       | L. sibiricus     |                     | Apal         |           | BR:Jan735:10    | KX348206                  |
| CF739                       | 06/10/10        | Engenheiro Beltrão, PR           | S23 42' 12.60"          | W52 08' 54.60''      | L. sibiricus     | Sacl                | HindIII      |           | BR:Egb739:10    | KX348168 (A) KX348207 (B) |
| CF775                       | 08/10/10        | Cascavel, PR                     | S24 52' 06.00"          | W53 20' 28.00''      | L. sibiricus     | Apal                | Apal         |           | BR:Cas775:10    | KX348172 (A) KX348222 (B) |
| CF776                       | 08/10/10        | Cafelândia, PR                   | S24 39' 21.30"          | W53 13' 07.40''      | L. sibiricus     | Apal                | Apal         |           | BR:Caf776:10    | KX348175 (A) KX348221 (B) |
| CF779                       | 08/10/10        | Ubiratã, PR                      | S24 33' 07.00"          | W53 00' 34.00''      | L. sibiricus     |                     | HindIII      |           | BR:Ubi779:10    | KX348208                  |
| CF784                       | 08/10/10        | Campo Mourão, PR                 | S23 57' 14.90"          | W52 20' 59.30''      | L. sibiricus     | Sacl                | Kpnl         |           | BR:Cam784:10    | KX348167 (A) KX348209 (B) |
| CF793                       | 08/11/10        | Londrina, PR                     | S23 26' 20.30"          | W51 08' 17.90''      | L. sibiricus     |                     | Apal         |           | BR:Lon793:10    | KX348207                  |
| CF796                       | 08/11/10        | São Domingos, PR                 | S24 00' 51.00"          | W51 30' 37.00''      | L. sibiricus     |                     | Kpnl         |           | BR:Sad796:10    | KX348219                  |
| CF802                       | 08/11/10        | Ivaiporã, PR                     | S24 18' 37.00"          | W51 43' 24.00"       | L. sibiricus     |                     | Apal         |           | BR:lva802:10    | KX348218                  |
| CF1024                      | 04/26/11        | Francisco Beltrão, PR            | S26 00' 09.00"          | W52 56' 36.00''      | L. sibiricus     |                     | Apal         |           | BR:Frb1024:11   | KX348210                  |
| CF1058                      | 06/06/11        | Pato Branco, PR                  | S26 11' 47.44"          | W52 49' 23.87"       | L. sibiricus     | Apal                |              |           | BR:Pab1058:11   | KX348179                  |
| CF1067                      | 06/06/11        | Ampere, PR                       | S25 57' 06.90"          | W53 24' 25.00"       | L. sibiricus     | Apal                |              |           | BR:Amp1067:11   | KX348173                  |
| CF1075                      | 06/07/11        | Toledo, PR                       | S24 46' 46.00"          | W53 40' 41.00''      | L. sibiricus     |                     | Apal         |           | BR:Tol1075:11   | KX348211                  |
| CF1077                      | 06/07/11        | Nova Mercedes, PR                | S24 30' 54.70"          | W54 07' 0.22"        | L. sibiricus     |                     | Apal         |           | BR:Nom1077:11   | KX348217                  |
| CF1083                      | 06/07/11        | Guaíra, PR                       | S24 14' 04.00"          | W54 11' 57.00"       | L. sibiricus     | Spel                | Apal         |           | BR:Gua1083:11   | KX348166 (A) KX348216 (B) |
| CF1 095                     | 06/08/11        | Dourados, MS                     | S22 17' 58.00"          | W54 49' 14.10''      | L. sibiricus     | Apal                | Apal         |           | BR:Dou1095:11   | KX348165 (A) KX348212 (B) |
| CF1111                      | 06/08/11        | Aral Moreira, MS                 | S22 46' 19.70"          | W55 24' 43.90"       | L. sibiricus     |                     | HindIII      |           | BR:Arm1111:11   | KX348213                  |
| CF1130                      | 06/09/11        | Araruna, PR                      | S24 03' 50.00"          | W52 33' 52.00''      | L. sibiricus     | Apal                |              |           | BR:Ara1130:11   | KX348176                  |
| CF1135                      | 06/09/11        | Sertanópolis, PR                 | S23 50' 25.00"          | W52 18' 19.00''      | L. sibiricus     | Apal                | Apal         |           | BR:Ser1135:11   | KX348178 (A) KX348215 (B) |
| Euphorbia yello             | w mosaic alph   | nasatellite                      |                         |                      |                  |                     |              |           |                 |                           |
| CF18                        | 03/19/09        | Chapada, RS                      | S28 01' 08.80"          | W53 05' 52.31'       | <i>Sida</i> spp. |                     |              | EcoRI     | BR:Cha18:09     | KX348227                  |
| Leonurus yellow             | v spot alphasa  | tellite                          |                         |                      |                  |                     |              |           |                 |                           |
| CF1 095                     | 06/08/11        | Dourados, MS                     | S22 17' 58.00"          | W54 49' 14.10''      | L. sibiricus     |                     |              | EcoRI     | BR:Dou1095.1:11 | KX348228                  |
|                             |                 |                                  |                         |                      |                  |                     |              |           | BR:Dou1095.2:11 | KX348229                  |
|                             |                 |                                  |                         |                      |                  |                     |              |           | BR:Dou1095.3:11 | KX348230                  |
|                             |                 |                                  |                         |                      |                  |                     |              |           | BR:Dou1095.4:11 | KX348231                  |
|                             |                 |                                  |                         |                      |                  |                     |              |           | BR:Dou1095.5:11 | KX348232                  |
| <sup>a</sup> Enzyme used fi | or releasing g  | enome-length DNA components afte | er rolling-circle ampli | fication and subsedu | ent cloning into | the plasmic         | d vector pKS |           |                 |                           |

Ann Appl Biol **170** (2017) 204–218 © 2017 Association of Applied Biologists

208



Figure 1 Bayesian phylogenetic trees based on the complete DNA-A (A) and DNA-B (B) nucleotide sequences of the begomoviruses obtained from *Sida* spp. (red) and *Leonurus sibiricus* plants (green) in this study plus begomoviruses from the New World and Old World (see Table S1 for full names and GenBank access numbers). BR:Vic25.4:11 and BR:Vic26:11 (blue) are unpublished sequences determined in our laboratory (Xavier, 2015). The OW begomoviruses *African cassava mosaic virus* (ACMV), *East African cassava mosaic virus* (EACMV), *Tomato yellow leaf curl virus* (TYLCV) and *Watermelon chlorotic stunt virus* (WmCSV) were used as outgroups. Nodes with posterior probability values between 0.50 and 0.89 are indicated by empty circles and those with values equal to or greater than 0.90 are indicated by filled circles.

Ann Appl Biol **170** (2017) 204–218 © 2017 Association of Applied Biologists



Figure 1 Continued.

Ann Appl Biol **170** (2017) 204–218 © 2017 Association of Applied Biologists

### Diversity of begomovirus and satellites in Sida and Leonurus

Table 2 Recombination events detected in the Sida bright yellow mosaic virus (SiBYMV-BR:Tac720:10) and Sida golden yellow spot virus (SiGYSV -BR:Sab889:10) genomes, based on a data set including the nucleotide sequences of the begomoviruses described in this study and additional begomoviruses from the New World (NW) and Old World (OW)

|       |                             | Recombination Breakpoints <sup>b</sup> |      | Parents           |                    |                     |                          |
|-------|-----------------------------|--|------|-------------------|--------------------|---------------------|--------------------------|
| Event | Recombinant                 | Begin                                  | End  | Minor             | Major              | Method <sup>c</sup> | P-value <sup>d</sup>     |
|       | Sigysv <sup>a</sup> (DNA-A) |  |      |                   |                    |                     |                          |
| 1     | BR:Sab889:10                | 180                                    | 1178 | Unknown           | SiMBoV1 (HM585441) | <b>R</b> GBMCS3     | $5.066 \times 10^{-38}$  |
| 2     | BR:Sab889:10                | 1448                                   | 1889 | SiYLCV (KC706539) | Unknown            | RMC3                | $2.524 \times 10^{-7}$   |
|       | Sibymv (DNA-A)              |  |      |                   |                    |                     |                          |
| 3     | BR:Tac720:10                | 193                                    | 1906 | Unknown           | SiMMV (FN436003)   | RGBMCS <b>3</b>     | $7.891 \times 10^{-28}$  |
|       | Sibymv (DNA-B)              |  |      |                   |                    |                     |                          |
| 4     | BR:Tac720:10                | 2562                                   | 587  | BR:Tol1075:11     | BDMV (M88180)      | RBMCS               | 1.725 × 10 <sup>-6</sup> |

<sup>a</sup>See Table S1 for the complete data set and full virus names.

<sup>b</sup>Recombination breakpoint coordinates are according to the first nucleotide after the cleavage site at the origin of replication, increasing clockwise.

<sup>c</sup>Recombination events and their putative parental viruses were identified using the Rdp (R), Geneconv (G), Boostcan (B), Maxichi (M), Chimaera (C), Siscan (S) and 3Seg (3) modules in RDP4.

<sup>d</sup>The reported *P*-values are for the programs indicated in bold in the 'Method' column and are the lowest *P*-values calculated for the region in question.

A DNA-A component (BR:Sab889:10) was cloned from the *Sida* spp. sample #889 (Table 1; a DNA-B was detected in sample #889 but has not yet been cloned). Pairwise comparisons indicated the highest nt sequence identities of 72.1% with *Tomato mottle leaf curl virus* (ToMoLCV, JF803249) and 73.8% with two unpublished sequences obtained from *Sida acuta* samples in our laboratory (BR:Vic25.4:11 and BR:Vic26.1:11; Xavier, 2015). Thus, BR:Sab889:10 constitutes the third new virus species detected in the *Sida* spp. samples, for which the name Sida golden yellow spot virus (SiGYSV) is proposed.

Strikingly, the DNA-A components of BR:Sab889:10, BR:Vic25.4:11 and BR:Vic26.1:11 have a length within the range of OW begomoviruses (2813, 2828 and 2828 nt, respectively) and contain an av2-like gene (which is present only in OW begomoviruses). The region encompassing part of the CR and the av2-like and cp genes of these three components (approximately 1100 nt) has very low similarity to any other begomovirus. The deduced amino acid sequences of their CP and AV2-like proteins were further analysed with BLASTp and with the program Interpro. The analysis performed with Interpro indicated the presence of a domain related to geminivirus CPs, despite the divergence of the cp gene (data not shown). No functional domains were predicted in the AV2-like protein. BLASTp analysis with the CP and AV2 proteins detected only a very low similarity with a highly divergent monopartite geminivirus recently described in China infecting apple trees, named 'apple geminivirus' (AGV) (Liang et al., 2015). The BR:Sab889:10 CP shares 27% amino acid (aa) identity (90% coverage, *E* value  $2e^{-12}$ ) with the AGV CP and the AV2-like protein shares 43% aa identity (83% coverage, *E* value  $1e^{-15}$ ) with the putative V2 protein of AGV.

Analysis with the RDP4 program detected two strongly supported recombination events in BR:Sab889:10, with Sida yellow leaf curl virus (SiYLCV, KC706539), Sida mosaic Bolivia virus 1 (SiMBoV1, HM585441) and two unknown viruses identified as putative parents (Table 2). In the phylogenetic tree, SiGYSV was placed in a cluster with BR:Vic25.4:11, BR:Vic26.1:11 and ToMoLCV (Fig. 1A), which is consistent with the pairwise identity analysis. A cp nt sequence tree was constructed with a data set including four highly divergent geminiviruses (Citrus chlorotic dwarf-associated virus, CCDaV; Euphorbia caput-medusae latent virus, EcmLV; Grapevine red blotch associated virus, GRBaV; apple geminivirus, AGV), one topocuvirus (Tomato pseudo-curly top virus, TPCTV), BR:Vic25.4:11 and BR:Vic26.1:11, and seven NW and OW begomoviruses (Fig. 2). BR:Sab889:10 clusters with BR:Vic25.4:11, BR:Vic26.1:11 and AGV, reflecting the BLASTp analysis in which a similarity was found among the CPs of these four viruses. Studies are in progress in our laboratory to characterise these divergent, OW-like begomovirus species.

The vast majority of the DNA-A clones obtained from *L. sibiricus* samples corresponded to *Tomato yellow spot virus* (ToYSV) isolates (15 out of 16 DNA-A clones and 19 out of 20 DNA-B clones; Table 1). Pairwise sequence comparisons indicated that the DNA-A sequences share >93.4% nt sequence identity with each other and 92.3% to 95.5% nt identity with the sequence of ToYSV (DQ336350), and the DNA-B sequences share >91.9% nt identity amongst themselves and 91.0% to 92.3% nt identity with ToYSV (DQ336351) (Table 1; Fig. S1). The Bayesian phylogenetics trees based on the DNA-A and DNA-B components placed these isolates in clusters with ToYSV (Fig. 1).

The cloned DNA-A and DNA-B (BR:Arm713:10) from *L. sibiricus* sample #713 have 97.2% and 95.3%



**Figure 2** Bayesian phylogenetic tree based on the nucleotide sequences of the *cp* genes of Sida golden yellow spot virus (SiGYSV, in red and selected geminiviruses). The unrooted *cp* tree includes four highly divergent geminiviruses (apple geminivirus, AGV; citrus chlorotic dwarf associated virus, CCDaV; *Euphorbia caput-medusae* latent virus, ECMLV; and grapevine red-blotch associated virus, GRBaV), one topocuvirus (*Tomato pseudo-curly top virus*, TPCTV), the BR:Vic25.4:11 and BR:Vic26:11 sequences (blue) determined in our laboratory, in addition to seven New World and Old World begomoviruses (see Table S1 for full names and GenBank access numbers). Nodes with posterior probability values between 0.60 and 0.89 are indicated by empty circles and those with values equal to or greater than 0.90 are indicated by filled circles.

nt sequence identity with EuYMV (FJ619507 and FJ619508), respectively (Table 1; Fig. S1). EuYMV has been found sporadically in some non-cultivated hosts (Fernandes *et al.*, 2011; Silva *et al.*, 2012; Tavares *et al.*, 2012; Barreto *et al.*, 2013; Rocha *et al.*, 2013), but to our knowledge, this is the first report of EuYMV infecting plants of the genus *Leonurus*.

Alphasatellite DNA molecules were cloned from samples #18 and #1095 (*Sida* spp. and *L. sibiricus*, respectively; Table 1). Isolate BR:Cha18:09 (1338 nt) showed the highest nt sequence identity (93.2%) to *Euphorbia yellow mosaic alphasatellite* (EuYMA, FN436008) (Fig. S4) and a close phylogenetic relationship with this isolate (Fig. 3).

From sample #1095, five alphasatellite clones (BR:Dou1095.1:11 to BR:Dou1095.5:11; all 1367 nt) as well as begomovirus DNA-A and DNA-B components were cloned (Table 1). Pairwise nt sequence comparisons indicated that the DNA-A and DNA-B sequences showed the highest identities with ToYSV (94.6% and 93.0%, respectively; Fig. S1). Alphassatellite sequences shared >99.9% nt identity amongst themselves and 82.3% to

82.4% nt identity with the sequence of EuYMA (Fig. S4). The sequences showed typical features of alphasatellite molecules, containing one ORF (*alpha-Rep*) potentially encoding a Rep protein with 313 amino acids (*data not shown*). The deduced amino acid sequences of the ORF display 86.9% identity with the *Cleome leaf crumple alphasatellite* (ClLCrA) alpha-Rep protein. The sequences also contain an A-rich region located immediately downstream of the ORF (coordinates 1115–1222, with a 57% adenine content) and a predicted hairpin structure containing, within the loop, the nonanucleotide TAG-TATTAC, which is conserved in alphasatellites (Zhou, 2013).

Phylogenetic analysis showed that the isolates grouped most closely with EuYMA (Fig. 3), consistent with the pairwise identity analysis. According to the proposed demarcation threshold of 83% nt sequence identity for alphasatellites (Mubin *et al.*, 2009, 2012), the five clones from *L. sibiricus* sample #1095 represent a distinct alphasatellite, for which the name Leonurus yellow spot alphasatellite (LeYSA) is proposed.

Diversity of begomovirus and satellites in Sida and Leonurus



Figure 3 Bayesian phylogenetic tree based on the alphasatellites sequences described in this study from *Sida spp.* (red) and *Leonurus sibiricus* plants (green) and the most closely related alphasatellites (see Table S2 for full names and GenBank access numbers). Nodes with posterior probability values between 0.50 and 0.89 are indicated by empty circles and those with values equal to or greater than 0.90 are indicated by filled circles.

Originally thought to be restricted to the OW, alphasatellites have recently been found in association with bipartite begomoviruses in Brazil (state of Mato Grosso do Sul), Cuba and Venezuela (Paprotka *et al.*, 2010; Romay *et al.*, 2010; Jeske *et al.*, 2014). Our results extend the geographical range of alphasatellites in South America, suggesting that these molecules may be widespread in the continent, and also their host range, with the first report of EuYMA infecting plants of the genus *Sida* and the detection of a new alphasatellite, LeYSA, in *L. sibiricus*.

*Sida* spp. are arguably the most abundant natural reservoirs for begomoviruses in several regions of the world (Frischmuth *et al.*, 1997; Hofer *et al.*, 1997; Roye *et al.*, 1997; Echemendía *et al.*, 2004; Jovel *et al.*, 2004; Xiong *et al.*, 2005; Guo & Zhou, 2006; Das *et al.*, 2008; Fiallo-Olivé *et al.*, 2010; Fiallo-Olivé *et al.*, 2012). For example, in a recent survey, Tavares *et al.* (2012) reported the occurrence of nine begomoviruses (including four new species) in 57 *Sida* spp. samples collected in the Brazilian states of Minas Gerais and Alagoas. Here, out of 43 *Sida* spp. samples, we found two previously described begomoviruses, three new species, and also (for the first time) an alphasatellite, further emphasising the tremendous diversity of begomoviruses and associated DNA satellites naturally infecting *Sida* spp.

Ann Appl Biol **170** (2017) 204–218 © 2017 Association of Applied Biologists Conversely, *L. sibiricus* harbours a much lower diversity of begomoviruses, with ToYSV as the causative agent of infection in almost all samples, corroborating with published data (Fernandes *et al.*, 2014). ToYSV was first reported infecting tomato plants in the state of Minas Gerais (Ambrozevicius *et al.*, 2002), and was later reported in bean and soybean plants in northwestern Argentina (Rodríguez-Pardina *et al.*, 2011). *L. sibiricus* is a widely distributed plant in Brazil, and seems to be the main natural reservoir, as well as a potential source of inoculum, of ToYSV to bean, soybean and tomato crops, as previously noted also by Barbosa *et al.* (2012).

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## **Supporting Information**

Additional Supporting Information may be found in the online version of this article:

**Table S1.** Geminivirus sequences used for pairwise sequence comparisons, phylogenetic analysis and detection of recombination events.

**Table S2.** Alphasatellite sequences used for pairwise sequence comparisons, phylogenetic analysis and detection of recombination events.

**Fig. S1.** Pairwise sequence identity matrices of the DNA-A (A) and DNA-B (B), between the begomoviruses obtained from *Sida* spp. (red) and *Leonurus sibiricus* plants (green) in this study and additional begomovirus from the Americas. BR:Vic25.4:11 and BR:Vic26:11 (blue) are unpublished sequences determined in our laboratory (Xavier, 2015).

Fig. S2. Bayesian phylogenetic trees based on the nucleotide sequences of the cp (A) and rep (B) genes of the begomoviruses obtained from Sida spp. (red) and Leonurus sibiricus plants (green) in this study plus begomoviruses from the New World and Old World (see Table S1, Supporting Information, for full names and GenBank access numbers). BR:Vic25.4:11 and BR:Vic26:11 (blue) are unpublished sequences determined in our laboratory (Xavier, 2015). The OW begomoviruses African cassava mosaic virus (ACMV), East African cassava mosaic virus (EACMV), Tomato yellow leaf curl virus (TYLCV) and Watermelon chlorotic stunt virus (WmCSV) were used as outgroups. Nodes with posterior probability values between 0.50 and 0.89 are indicated by empty circles and those with values equal to or greater than 0.90 are indicated by filled circles.

**Fig. S3.** Bayesian phylogenetic trees based on the nucleotide sequences of the *mp* (A) and *nsp* (B) genes of the begomoviruses obtained from *Sida* spp. (red) and *Leonurus sibiricus* plants (green) in this study plus begomoviruses from the New World and Old World (see Table S1 for full names and GenBank access numbers). The OW begomoviruses *African cassava mosaic virus* (ACMV), *East African cassava mosaic virus* (EACMV) and *Watermelon* 

#### Diversity of begomovirus and satellites in Sida and Leonurus

*chlorotic stunt virus* (WmCSV) were used as outgroups. Nodes with posterior probability values between 0.50 and 0.89 are indicated by empty circles and those with values equal to or greater than 0.90 are indicated by filled circles. **Figure S4.** Pairwise sequence identity matrices between the alphasatellites sequences described in this study from *Sida* spp. (red) and *Leonurus sibiricus* plants (green) and the most closely related alphasatellites (see Table S2 for full names and GenBank access numbers).