A novel putative member of the family *Benyviridae* is associated with soilborne wheat mosaic disease in Brazil

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Soilborne wheat mosaic disease (SBWMD), originally attributed to infections by *Soilborne wheat mosaic virus* (SBWMV) and *Wheat spindle streak mosaic virus* (WSSMV), is one of the most frequent virus diseases and causes economic losses in wheat in southern Brazil. This study aimed to characterize molecularly the viral species associated with wheat plants showing mosaic symptoms in Brazil. Wheat leaves and stems displaying mosaic symptoms were collected from different wheat cultivars in Passo Fundo municipality, Rio Grande do Sul State, southern Brazil. Double-stranded RNA was extracted and submitted to cDNA library synthesis and next-generation sequencing. No sequences of SBWMV and WSSMV were detected but the complete genome sequence of a putative new member of the family *Benyviridae* was determined, for which the name wheat stripe mosaic virus (WhSMV) is proposed. WhSMV has a bipartite genome with RNA 1 and RNA 2 organization similar to that of viruses belonging to *Benyviridae*. WhSMV RNA 1 has a single open reading frame (ORF) encoding a polyprotein with putative viral replicase function. WhSMV RNA 2 has six ORFs encoding the coat protein, the major protein (read-through), triple gene block movement proteins (TGB 1, 2 and 3) and ORF 6 (hypothetical protein). In addition to the genomic organization and nucleotide and amino acid sequence identities, phylogenetic analyses also corroborated that WhSMV is a virus species of the *Benyviridae*. However, isolates of WhSMV formed a clade distinct from members of the genus *Benyivirus*. It was also demonstrated that the plasmodiophorid *Polymyxa graminis* is associated with wheat roots showing SBWMD symptoms and infected by WhSMV.

**Keywords:** *Benyviridae*, mosaic symptoms, next-generation sequencing, *Polymyxa graminis*, *Triticum aestivum*, wheat stripe mosaic virus

**Introduction**

Wheat (*Triticum* spp.) belongs to the family Poaceae and is one of the most cultivated cereals globally. The *Triticum aestivum* species, known as common wheat, is heavily cultivated and responsible for over 80% of world wheat production. In Brazil, wheat is grown during the winter and spring, mainly in the three southern states: Paraná (PR), Santa Catarina (SC) and Rio Grande do Sul (RS). These states together accounted for 85% of the total Brazilian wheat produced during the 2017 season (Companhia Nacional de Abastecimento, 2018).

Viral diseases are important factors impeding the management of wheat crops in Brazil and can cause serious damage and losses. Yellow dwarf disease together with soilborne wheat mosaic disease (SBWMD) are the most common viral diseases and can cause significant losses in wheat fields in southern Brazil (Lau, 2014). Yellow dwarf disease is caused by a number of related viruses, such as *Barley yellow dwarf virus*-PAV (BYDV-PAV, *Luteovirus*, *Luteoviridae*) and *Cereal yellow dwarf virus*-RPV (CYDV-RPV, *Polerovirus*, *Luteoviridae*; Caetano, 1972; Mar et al., 2013); SBWMD is attributed to *Soilborne wheat mosaic virus* (SBWMV, *Furovirus*, *Virgaviroidae*; Caetano et al., 1971, 1978) and *Wheat spindle streak mosaic virus* (WSSMV, *Bymovirus*, *Potyviridae*; Caetano, 1998). In southern Brazil, under ideal conditions such as wet and compacted soil and with temperatures near to 15 °C, SBWMD induces yield reductions to the wheat crop of over 50% (Barbosa et al., 2001). In an assay carried out in an area infested with the plasmodiophorid *Polymyxa graminis* and with a history of SBWMD, Dalbosco et al. (2002b) verified a reduction in grain weight per plant from 7% to 56% in wheat and from 51% to 59% in triticale, depending on the genotype. In the 1000-grain weight, the decrease varied from 4% to 26% in wheat and from 23% to 30% in triticale (Dalbosco et al., 2002b).

Caetano et al. (1971) described that SBWMD had been seen in the southern states of Brazil since the 1970s.

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and that SBWMV was its causal agent. The diagnosis of SBWMV was based on mechanical transmission tests, in vitro physical properties, particle morphology and ultrastructural aspects of affected plant tissues (Caetano et al., 1978). However, these diagnostic methods are subjective and prone to error. Nowadays in Brazil, the detection of the aetiological agent of SBWMD is still based on observation of the symptoms and viral particles. In addition, the use of antisera or primers designed from viral sequences from other countries has only resulted in negative reactions in the diagnostic tests, suggesting that the SBWMV and WSSMV isolates found in Brazil could be divergent from other regions or that a species not yet characterized is associated with SBWMD, especially because there is a lack of an appropriate tool for the characterization of viral species associated with SBWMD. 

Next-generation sequencing (NGS) is an appropriate tool for the characterization of viral species associated with SBWMD, especially because there is a possibility of mixed infections and differences in nucleotide sequences of the viral populations associated with wheat fields in Brazil. A metagenomic approach may also detect the presence of viral species not yet characterized.

The presence of *P. graminis* in the roots of wheat plants with SBWMD symptoms has been reported in Brazil by Caetano et al. (1971), Pierobom et al. (1972) and Dalbosco et al. (2002a). *Polymyxa graminis* is an obligate root parasite of several temperate grasses and represents a significant threat to agriculture due to its ability to transmit viruses of agronomic importance, including SBWMV. *Polymyxa graminis* is responsible for the transmission of at least 14 plant viruses belonging to the *Furovirus*, *Bymovirus*, *Pecluvirus* and *Benyvirus* genera (Adams & Jacquier, 1994).

Currently, the family *Benyviridae* includes four species, *Beet necrotic yellow vein virus* (BNYVV), *Beet soilborne mosaic virus* (BSBMV), *Burdock mottle virus* (BdMV) and *Rice stripe necrosis virus* (RSNV), all of the genus *Benyivirus*, and two tentative species, Mangifera indica latent virus (MILV) and Hubei beny-like virus 1 (HBBLV; Gilmer et al., 2017). The genome of members of the genus *Benyivirus* is composed of at least two polyadenylated positive-sense single-strand (ss) RNA molecules of approximately 7 and 4.6 kb (Bouzoubaa et al., 1986, 1987; Kiguchi et al., 1996; Saito et al., 1996). The better-known members of the genus *Benyivirus* are transmitted by the root-infesting vectors *Polymyxa betae* and *P. graminis* (Tamada et al., 1996; Morales et al., 1999).

The main objective of this study was to characterize molecularly the viral species associated with wheat plants showing SBWMD symptoms and to determine whether the plasmodiophorid *P. graminis* is associated with wheat roots exhibiting SBWMD symptoms and infected by the detected viral species.

**Materials and methods**

**Sample collection for NGS analysis**

Leaves and stems of wheat (*T. aestivum*) with typical mosaic symptoms (Fig. 1) were collected in Embrapa Trigo in Passo Fundo municipality, Rio Grande do Sul state, southern Brazil where, historically, field trials were carried out to assess the susceptibility and resistance of wheat cultivars to SBWMD (Rio Grande do Sul State Assay). In this area the infected plants are known to contain typical particles (rod-shaped) of the SBWMV described by Caetano et al. (1978). Samples, including leaves and stems, from each of the eight cultivars (Karl 92, Everest, OR 1, CEP 11, TBIO Toruk, TBIO Tibagi, Esporão and LG Oro) were collected and stored in a freezer (−80 °C).

**Double-stranded RNA extraction**

Extraction of double-stranded RNA (dsRNA) was performed as described by Valverde et al. (1990) with some modifications. Some samples with SBWMD symptoms did not have sufficient vegetative tissue; thus, two mixed samples were generated (TBIO Toruk and TBIO Tibagi, Esporão and LG Oro).

Samples of 30 g wheat tissue were ground with liquid nitrogen and incubated with 129 mL extraction buffer (90 mL 2× sodium chloride-Tris-EDTA buffer (STE); 35 mL 10% sodium dodecyl sulphate (SDS); 2 mL bentonite (45 mg mL⁻¹); 2 mL β-mercaptoethanol for 5 min in a shaker at room temperature. After adding 35 mL of Tris-HCl-saturated phenol (pH 8.0) and 35 mL chloroform:isoamyl alcohol (24:1 v/v) to the solution, samples were incubated for a further 1 h. The solution was centrifuged (16 274 g) for 10 min at 4 °C, and the resulting supernatant (nucleic acid solution) was enriched for dsRNA using Whatman CF11 cellulose affinity chromatography. Three cycles of centrifugation and one cycle of cellulose chromatography, all of them using 1× STE buffer with 17% ethanol (v/v) for resuspending the cellulose pellet or washing the column, were used to remove residual ssRNA. The sediment cellulose was collected and resuspended after each centrifugation. After the third centrifugation cycle, the resuspended cellulose was transferred into a column and washed. The dsRNA was eluted with 1× STE buffer, precipitated, centrifuged and resuspended. Subsequently, the dsRNA was cleaned using a RNeasy Plant Mini kit (QIAGEN), following the manufacturer’s protocol.

The purity and concentration of the dsRNA were measured by spectrophotometry (NanoDrop 2000; Thermo Scientific). The dsRNA samples were placed in RNA Stable tubes (Biomatrica) and dried in a Speed Vac (Eppendorf Concentrator Plus) for 1.5 h; NGS was performed at Macrogen Inc.

**NGS and bioinformatic analyses**

NGS sequencing reads were generated from a complementary DNA library (cDNA) using the Illumina HiSeq 2500 platform (Macrogen Inc.). Adapters and low-quality reads were removed by TRIMMOMATIC (Bolger et al., 2014). The NGS data were initially analysed using the TRINITY tool (Grabherr et al., 2011) for RNASeq assembly. The resulting contigs were analysed for similarity to records in public databases (GenBank). Sequences homologous to the wheat genome were removed in order to analyse only those with viral origin. NGS reads that did not map to the wheat genome were again submitted to genome assembly with TRINITY and, in parallel, with VELVET (Zerbino & Birney, 2008). The results were compared using caff (Huang & Madan, 1999) in two stages. In the first stage, the contigs resulting from the initial assemblies of TRINITY and VELVET were analysed individually. In the second stage, the contigs resulting from the first step were again grouped and analysed. The results showed that the VELVET-mounted contigs were contained in those derived from the TRINITY assembly. The contigs were
subsequently compared with the NCBI database using the BLASTN and BLASTX search tools with standard parameters.

RT-PCR protocol

For cDNA synthesis, 100 ng purified dsRNA was denatured at 94 °C for 5 min followed by cDNA synthesis using the oligo dT and the iProof II Reverse Transcription System kit (Promega), in accordance with the manufacturer’s recommendations. The PCR mixture contained 5 μL cDNA (c. 1 μg), 10 μL of 5× PCR buffer, 2 μL MgCl2 (25 mM), 1 μL of dNTP (10 mM), 1 μL of 0.25 Go Taq DNA polymerase (5 U μL−1; Promega) and ultrapure water to a final volume of 50 μL. Reaction conditions were 94 °C for 2 min; 34 cycles of 94 °C for 1 min, 42–54 °C (depending on the primer used, Table 1) for 1 min, and 72 °C for 1 min; and a final extension at 72 °C for 10 min. The amplified fragments were subjected to 1% agarose gel electrophoresis, stained with GelRed (Biotium), visualized under a UV light transilluminator, and photographed. PCR fragments were purified using the Gene Jet kit (Fermentas), in accordance with the manufacturer’s recommendations, and sequenced (Helixxa Genomic Services).

Sequence analysis and phylogeny

The contigs were submitted to open reading frame (ORF) prediction using the ORF FINDER program (NCBI, https://www.ncbi.nlm.nih.gov/orffinder/). The identification of conserved and functional domains of proteins was performed using the SMART tool (http://smart.embl-heidelberg.de/). Phylogenetic trees were constructed using the maximum likelihood method implemented in the MEGA v. 6.0 program (Tamura et al., 2013) with Tamura’s three-parameter model, gamma distribution (G), and invariant sites (I). The robustness of each internal branch was estimated from 2000 bootstrap replicates for the coat protein (CP) and for the triple gene block movement 1 (TGB 1) genes, and from 6000 bootstrap replicates for the replicate gene. The alignments used for the phylogenetic analyses were performed using the MUSCLE tool available in MEGA v. 6.0.

Nucleotide and amino acid identity analyses were performed with multiple sequence alignment (https://www.ebi.ac.uk/Tools/msa/muscle/), using the complete genome sequences and coding regions of the species described in the family Benyviridae and the isolates characterized in this study.

Vector detection and characterization

Wheat roots of the cultivars TBIO Toruk and BRS Guamirim were collected at the Embrapa Trigo experimental station in Passo Fundo municipality, Rio Grande do Sul state, southern Brazil. Ten plants with symptoms from each cultivar were collected and the roots were pooled separately.

Wheat roots were washed in running water, dried in filter paper and prepared for visualization of P. graminis structures. The roots were immersed in a solution containing 10 mL lactic acid (85–90%), 20 mL glycerol (99.5%), 10 mL distilled water, 10 g crystallized phenol, and 0.5 g blue cotton dye and heated to 100 °C. Microscope slides were prepared and the P. graminis structures were visualized under an optical microscope at ×400 magnification.
Molecular identification of *P. graminis* was carried out using rDNA as a target. The partial nuclear 5.8S rRNA gene and internal transcribed spacer 1 (ITS1) were amplified by PCR from root samples collected from each cultivar. The washed and dried roots were ground with liquid nitrogen. Total DNA extraction using 100 mg powdered root tissues was performed in accordance with the protocol described by Doyle & Doyle (1987), followed by PCR using the primers Pgfwd2/Pxrev7 (Ward & Adams, 1998). The PCR mixture consisted of 1 μL DNA (100 ng), 5 μL of 5 × buffer, 2 μL MgCl₂ (25 mM), 0.5 μL dNTPs (10 mM), 5 μL each primer (10 mM), 0.25 μL Taq DNA polymerase (5 U/μL; Promega) and ultrapure water to a final volume of 25 μL. PCR conditions were in accordance with the work of Tyagi et al. (2016). The amplified fragments were subjected to electrophoresis in 1% agarose gel, stained with GelRed (Biotium), visualized under UV light, and photographed. PCR fragments were purified using the Gene Jet kit (Fermentas), in accordance with the manufacturer’s recommendations, and Sanger sequencing was performed using the primers Pgfwd2/Pxrev7 (Helixxa Genomic Services). The sequences were compared with sequences from the database available in GenBank. A phylogenetic tree was constructed using the maximum likelihood method implemented in [MEGA](https://www.mega-cambodia.org) v. 6.0, using Kimura’s two-parameter model with invariant sites (I). The robustness of each internal branch was estimated from 1000 bootstrap replicates. The alignments used for the phylogeny were performed with the [MUSCLE](https://muse.tue.msu.edu) tool available in [MEGA](https://www.mega-cambodia.org) v. 6.0.

To determine the presence of the virus in root samples, total RNA extraction from the previously described root pool of the TBIO Toruk and BRS Guamirim cultivars was performed using the AxyPrep Multisource Total RNA Miniprep kit (Axygen Biosciences), in accordance with the manufacturer’s recommendations. The presence of the virus was determined following the RT-PCR protocol described above and using primers for amplification of the CP gene.

**Results**

**NGS and genome assembly**

Double-strand RNA from leaves and stems of six wheat samples with symptoms (cultivars Karl 92, Everest, OR 1, CEP 11, TBIO Toruk and TBIO Tibagi, Esoporão and LG Oro) was extracted and submitted to cDNA library synthesis and NGS analysis. The Illumina HiSeq 2500 platform generated approximately 30 million reads for each analysed sample. The samples yielded a large number of reads, considering the total number of sequenced bases, total number of reads, and CG/AT content analysation. This large number of reads was expected and the quality parameters (Q20) were higher than 99%, indicating excellent NGS quality (data not shown).

**TRINITY** software generated 127 665 contigs (1023 contigs homologous to viral sequences) and [VELVET](https://www.ornl.gov/pub/velvet/) generated 43 353 contigs (393 contigs homologous to viral sequences). The complete genome sequence of a putative novel species of the *Benyviridae* was obtained from all wheat samples analysed when the contig data were compared with already well-characterized viral species. No further reads and/or contigs of other viral species were found and the results indicated that only the novel species reported was present in the analysed samples.

The genome of the putative new virus species was composed of two RNA segments (RNA 1 and RNA 2; Fig. 2). The data indicated that the genomic organization of the characterized virus was very similar to those of members of the family *Benyviridae*. RNA 1 was the largest segment and varied in length from 6583 to 6600 nt, depending on the isolate and excluding the poly-A tail. The 5′- and 3′-UTR regions were 115 nt and 336–353 nt from Ivaiporí municipality (24°19′51.8″S, 51°45′20.9″W) were collected. The 26 samples were stored in a freezer (−80 °C) until processing. Total RNA extraction from each sample was performed using the AxyPrep Multisource Total RNA Miniprep kit (Axygen Biosciences), in accordance with the manufacturer’s recommendations. The presence of the virus characterized in this study was determined following the RT-PCR protocol described above using Beny_TGB2 primers (Table 1).

**Table 1** Primers used for NGS validation and detection of WhSMV.

<table>
<thead>
<tr>
<th>Set</th>
<th>Name</th>
<th>Sequence (5′–3′)</th>
<th>Position alignment (nt)</th>
<th>Annealing temperature (°C)</th>
<th>Amplicon length (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Beny_CP (F)</td>
<td>AAGTGTGCAGACGCCGCG</td>
<td>89</td>
<td>54</td>
<td>641</td>
</tr>
<tr>
<td></td>
<td>Beny_CP (R)</td>
<td>ATCGACCGAGGATGAAGA</td>
<td>730</td>
<td></td>
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</tr>
<tr>
<td>2</td>
<td>Beny_TGB2 (F)</td>
<td>AAGAGTTGTACGGAGGTTGG</td>
<td>3967</td>
<td>52</td>
<td>607</td>
</tr>
<tr>
<td></td>
<td>Beny_TGB2 (R)</td>
<td>CGTCGTCAGGCGCTGAAACTGT</td>
<td>4574</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Beny_Pol (F)</td>
<td>ATCGAGTCTAGCGGTTTACTG</td>
<td>3770</td>
<td>54</td>
<td>465</td>
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<tr>
<td></td>
<td>Beny_Pol (R)</td>
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<td>4234</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Beny_1_3-UTR</td>
<td>AGTGAAGCGTTGCTACTGTCGCG</td>
<td>6243</td>
<td>42</td>
<td>c. 357</td>
</tr>
<tr>
<td></td>
<td>Oligo (dT)₁₅</td>
<td>TTTTTTTTTTTTTTTTTTTTTTTTTTTT</td>
<td>4572</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Beny_2_3-UTR</td>
<td>ACGAAGATCGTCGTTTTTTTTTTTTTTTTTTTTTT</td>
<td>4572</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NGS, next-generation sequencing; WhSMV, wheat stripe mosaic virus; CP, coat protein; Pol, polymerase (replicase); TGB, triple gene block; UTR, untranslated region.

**Virus incidence**

Additional samples of 13 wheat cultivars (Everest, Karl 92, OR 1, CEP 11, TBIO Toruk and TBIO Tibagi, Esoporão and LG Oro) were collected in four areas from southern Brazil. All the collected samples presented typical symptoms of SBWMD. From Rio Grande do Sul state, in Caxixi (28°04′53.2″S, 52°24′55.8″W), Passo Fundo (28°13′40″S, 52°24′30″W) and Santo Augusto (27°54′04.20″S, 53°49′44.53″W) municipalities, 3, 11 and 3 samples, respectively, were collected. In Paraná state, nine samples were collected. In 20 municipalities, 30 samples were collected. In Santa Catarina state, nine samples were collected. In São Paulo state, 20 samples were collected.
in size, respectively, for all characterized isolates (Table 2). RNA 1 had a single ORF with the potential to encode a 231.7 kDa polyprotein with putative viral replication function. The polyprotein of the novel virus was possibly autocleaved into two proteins of 165.6 kDa and 66.1 kDa (Fig. 2) and contained methyltransferase, helicase (Hel), papain-like protease and RNA-dependent RNA polymerase (RdRP) motifs. A conserved motif (Gly-Asp-Asp), which is the signature of all viral RdRPs, was present in the replicase at position 1863–1865 aa of all of the characterized viral isolates. The RNA 2 segment ranged in length from 4879 to 4901 nt (Table 2), excluding the poly-A tail. The 5′- and 3′-UTR regions of RNA 2 were 145 nt and 96–116 nt in size, respectively, for all characterized isolates (Table 2). Six putative ORFs were identified in RNA 2 of the new virus (Table 2; Fig. 2). ORF 2 encoded the 18.9 kDa CP that had a stop codon (UAG). When this codon was translated by read-through protein strategies, an 80.8 kDa read-through (RT) protein (encoded from ORF 2A) could be produced (Fig. 2). In the RT protein, the VTER (Val-Thr-Glu-Arg) motif was found at amino acid positions 638–641 for WhSMV:BR:Everest, WhSMV:BR:Karl92 and WhSMV:BR:OR1; for WhSMV:BR:CEP11 and WhSMV:BR:Esporão-Oro, the amino acid positions were 640–643. ORFs 3, 4 and 5 comprised the putative triple gene block (TGB), associated with viral movement in plants (Fig. 2). ORF 3 (putative TGB 1) encoded a 54.6 kDa protein with a helicase domain. ORF 4 (putative TGB2) encoded a 13.8 kDa protein, which has a conserved domain associated with viral movement. ORF 5 (putative TGB 3) encoded a 12.8 kDa protein and ORF 6 encoded a hypothetical 2.8 kDa protein.

NGS validation

In order to confirm that the putative virus was not an artefact of NGS sequencing, five primers pairs were designed to the different genomic regions (Table 1), based on the sequence analysis. All primer sets provided amplicons in all six samples analysed and they were sequenced (capillary sequencing/Sanger; Fig. S1). Analysis of these sequences showed 99–100% identity with the sequences obtained by NGS. In addition, the use of the primer sets 4 and 5 (Table 1) was essential to confirm the accuracy of the NGS in obtaining the 3′ ends of RNAs 1 and 2, respectively, because fragments of the expected size and nucleotide sequences were obtained. These results confirmed the presence of the new putative virus in the wheat samples showing mosaic symptoms used for NGS.

Sequence analysis

The determined genome sequences of the viral isolates were deposited in GenBank (accession numbers

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**Figure 2** Genome organization and putative translation strategies of Wheat stripe mosaic virus (WhSMV). The scheme indicates putative self-cleavage of the replicase protein (black arrow RNA 1), a read-through stop codon UAG (black star RNA 2), putative m7Gppp (black circle) and the 3′ poly(A)-tails (Aₐ). The WhSMV has seven predicted open reading frames (ORFs): ORF 1 encodes putative viral replicase, ORF 2 encodes putative coat protein, ORF 2A encodes putative read-through domain, ORF 3 encodes putative triple gene block 1 (TGB 1), probably coded from subgenomic RNA a, ORF 4 encodes putative TGB 2, probably coded from subgenomic RNA b, ORF 5 encodes putative TGB 3, probably coded from subgenomic RNA b, and ORF 6 hypothetical protein, probably coded from subgenomic RNA c. Predicted molecular weights of proteins are indicated in kilodaltons (kDa). Dotted lines are used for all putative aspects.
The nucleotide identity of RNA 1 and RNA 2 among Brazilian isolates varied from 97% to 99%, indicating that all of them were isolates from the same viral species (Fig. 3). Sequence analysis indicated significant nucleotide and amino acid identity of Brazilian isolates with species of the family Benyviridae. When the Brazilian isolates, characterized in this study, were compared with the viruses belonging to the Benyviridae, low nucleotide identity was observed, varying from 48% to 51% and 42% to 47% for RNA 1 and RNA 2, respectively (Fig. 3). The Brazilian isolates showed highest identities for both RNA 1 and RNA 2 to BNYVV and BSBMV. When excluding the sequences of the Brazilian isolates, the nucleotide identities of RNA 1 and RNA 2 ranged from 52% to 78% and 45% to 70%, respectively among the species of the Benyviridae (Fig. 3).

Analysis of the coding regions revealed that the nucleotide and amino acid sequence identities among the viral isolates characterized here were 98–99% for ORF 1, 95–100% for ORF 2, 97–99% for ORF 2A, 96–100% for ORF 3, 99–100% for ORF 4, and 96–100% for ORF 5 (Fig. 3). ORF 6 presented 100% nucleotide and amino acid identity among the Brazilian isolates but was not compared with the other species of the family Benyviridae due to differences in size. ORF 1 of the Brazilian isolates shared 53–57% nucleotide identity with other species of the Benyviridae (Fig. 3), and ORF 2, 2A, 3, 4 and 5 shared nucleotide identities of 48–53%, 42–47%, 46–51%, 48–57% and 49–52%, respectively with other members of the Benyviridae (Fig. 3). In general, the amino acid identities were lower than those observed for nucleotides (Fig. 3).

Based on the genomic organization and the nucleotide and amino acid sequences analysed in the viral isolates described in this work, it is proposed that the virus found infecting different wheat cultivars in Brazil could be considered a new putative species in the family Benyviridae. For this species, the name Wheat stripe mosaic virus is proposed.

### Phylogenetic analysis

Phylogenetic trees were generated using the nucleotide sequences of the CP, replicase and TGB 1 genes, which verified the relationship of WhSMV to viruses of the family Benyviridae (Fig. 4). However, WhSMV isolates formed a distinct clade from the species of the genus Benyvirus, with strong bootstrap support (Fig. 4). The topology observed in the phylogenetic trees suggested that WhSMV does not belong to the genus Benyvirus. The Brazilian isolates formed a monophyletic group for all coding regions used (Fig. 4), indicating close relationships among them.

The phylogeny of complete RNA 1 and RNA 2 showed that the WhSMV isolates were closely related (Fig. S2), confirming the results of analysis of the coding regions and supporting that they are all of the same species. Additionally, phylogenetic analyses suggested that...
Figure 3  Two-dimensional plot representing the percentage sequence identities among isolates of wheat stripe mosaic virus (WhSMV) and species of the family Benyviridae for RNA 1, RNA 2, and all open reading frames (ORFs), except ORF 6. Percentage nucleotide and amino acid sequence identities are shown below and above the diagonal, respectively. ORF 1 encodes viral replicase, ORF 2 encodes coat protein, ORF 2A encodes read-through domain, ORF 3 encodes triple gene block 1 (TGB 1), ORF 4 encodes TGB 2, and ORF 5 encodes TGB 3. Members of the family Benyviridae are Beet necrotic yellow vein virus (BNYVV), Beet soil-borne mosaic virus (BSBMV), Burdock mottle virus (BdMV), Rice stripe necrosis virus (RSNV), and Mangifera indica latent virus (MILV). [Colour figure can be viewed at wileyonlinelibrary.com].
A novel virus infecting wheat in Brazil

Wheat stripe mosaic virus is a new species within the family Benyviridae and distinct from the other species of the genus Benyvirus.

Characterization of the plasmodiophorid associated with wheat roots

Members of the family Benyviridae are transmitted by parasites of the root system belonging to the order Plasmodiophorales. The presence of this vector in the wheat roots was confirmed by optical microscopy and it was possible to visualize the encysted zoospores (Fig. 5a) and the formation of zoosporangia (Fig. 5b) in both BRS Guamirim and TBIO Toruk cultivars.

To support the morphological data, parts of the analysed roots were used for detection and molecular characterization of the putative vector, as well as the detection of WhSMV. Using specific primers for the nuclear 5.8S rRNA gene and ITS1 and for the coding region of WhSMV CP, it was possible to amplify DNA fragments of the expected size for P. graminis (c. 240 and 310 bp) and WhSMV (641 bp) from the root samples of the two cultivars analysed (Fig. 5c). The DNA fragments of P. graminis obtained from wheat cultivars BRS Guamirim and TBIO Toruk in this study showed nucleotide identity of 98% among themselves and 100% among the isolates from BRS Guamirim (GenBank no. MH318570) and isolates from Belgium (GenBank no. AM259278) and the United Kingdom (GenBank no. AJ841287). Polynymyx graminis from wheat cultivar TBIO Toruk (GenBank no. MH318569) showed nucleotide identity of 99% with isolates from the Czech Republic (GenBank LT221860) and Australia (GenBank KF535921). The DNA fragments from the WhSMV CP amplification were also sequenced and showed nucleotide identity higher than 99% with the WhSMV isolates characterized in this work.

Phylogenetic analysis of the nuclear 5.8S rRNA gene and ITS1 supported the classification of the plasmodiophorid found in the wheat roots as P. graminis. Isolates from different countries and hosts formed two distinct groups but there was no division by host or geographical region (Fig. 5d). The two isolates of P. graminis characterized here are in different groups. Together, the results of the morphological, molecular and phylogenetic characterizations indicated the association of P. graminis with roots of wheat plants with mosaic symptoms infected by WhSMV in Brazil.

Prevalence of the new virus in different production areas

To verify the presence of the new virus in different areas, 26 samples of wheat were collected with symptoms in four different municipalities from southern Brazil. RT-PCR and electrophoretic analyses indicated the presence of WhSMV in all samples analysed (Fig. S3). Ivaiporá is 452 km away from Santo Augusto and Passo Fundo municipalities and 436 km away from Coxilha. These results indicate the association of the new virus characterized here with SBWMD in different wheat crop areas in Brazil.

Discussion

The viruses SBWMV and WSSMV have been reported as the causal agents of SBWMD in Brazil, based mainly on the viral particle morphology (Caetano et al., 1978; Lau, 2014). In fact, electron microscopy analyses carried out in 1978 (Caetano et al., 1978) and in 2015 (D. Lau, Embrapa Trigo, Passo Fundo, Brazil, personal communication) indicated the presence of rod-shaped virions of two different sizes, having the same particle morphology as expected for the WhSMV described here. In the present investigation, these two viruses were not present in the wheat plants showing mosaic symptoms. No reads of the SBWMV and WSSMV were detected in the NGS analysis. For the deep sequencing in this study, dsRNAs were used as templates and it is known that RNA viruses form dsRNA either as their encapsidated genome or during the process of replication. In metagenomic studies of plant viruses, the use of purified dsRNA has facilitated in-depth analyses of viruses from different families (Roossinck et al., 2015), supporting the accuracy of the result in the present study that only WhSMV infected the wheat samples analysed.

Members of the family Benyviridae have two to five segments of positive-sense ssRNA (Gilmer et al., 2017). The viral isolates described here showed two RNA segments, equal to the number for BdMV, RSNV and MILV (Lozano & Morales, 2009; Kondo et al., 2013; Sela et al., 2016). However, the type species of the genus Benyvirus, BNYVV, presented four or five RNA segments (Kiguchi et al., 1996; Saito et al., 1996) and BSBMV had four RNA segments (Heidel et al., 1997). No contigs were found that corresponded to viral genomic segments other than the RNA 1 and RNA 2 of benyviruses.

The 3′-UTR of WhSMV RNA 1 (115 nt) was similar in length to that of members of the genus Benyvirus (124–153 nt; Bouzoubaa et al., 1987; Lee et al., 2001; Lozano & Morales, 2009; Kondo et al., 2013). However, the 3′-UTR of WhSMV (336–353 nt) was larger than that found in members of the genus Benyvirus (181–233 nt; Lee et al., 2001; Lozano & Morales, 2009; Kondo et al., 2013).

RNA 1 of WhSMV encodes a 231.7 kDa polyprotein with putative viral replicase function and has a similar size to that observed for members of the genus Benyvirus (Saito et al., 1996; Lee et al., 2001; Lozano & Morales, 2009; Kondo et al., 2013). Proteolytic cleavage of the replicase has been described for members of the family Benyviridae and distinguishes this group from all other viruses with rod-shaped particles (Gilmer et al., 2017). The peptidase domain (papain-like proteinase) is involved in cleavage of the viral polyprotein (Hehn et al., 1997). A conserved GDD motif (Gly-Asp-Asp) is present in all WhSMV isolates and is also present in the same position within Benyviridae members (Bouzoubaa et al., 1987; Poch et al., 1989; Lee et al., 2001; Lozano & Morales, 2009).
The length of the RNA 2 of WhSMV (4879–4901 nt) was similar to that of members of the genus Benyvirus (4314–4635 nt; Bouzoubaa et al., 1986; Lee et al., 2001; Lozano & Morales, 2009; Kondo et al., 2013). The 5' UTR of WhSMV RNA 2 (145 nt) was also similar in length to that of members of the genus Benyvirus (144–
179 nt; Bouzoubaa et al., 1986; Lee et al., 2001; Lozano & Morales, 2009; Kondo et al., 2013). However, the 3′-UTR of WhSMV RNA 2 (96–116 nt) was smaller than those of of BNYVV, BdMV, BSBMV and RSNV (189–227 nt; Bouzoubaa et al., 1986; Lee et al., 2001; Lozano & Morales, 2009; Kondo et al., 2013).

Six putative ORFs were identified in RNA 2 of the new virus, as also observed for RNA 2 of BNYVV,
BSBMV and RSNV (Bouzoubaa et al., 1986; Lee et al., 2001; Lozano & Morales, 2009). ORF 2 encodes the viral CP responsible for protecting the genetic material and for long-distance movement. The KTER (Lys-Thr-Glu-Arg) motif found in the RT protein of BNYVV, BSBMV and RSNV is required for virus transmission by *P. betae* and *P. graminis* (Tamada et al., 1996; Lozano & Morales, 2009). A similar motif (VTER) was found in the amino acid sequences of the WhSMV. The involvement of the VTER motif in transmission by *P. graminis* of the new virus isolates characterized here needs to be verified.

ORFs 3, 4 and 5 comprise the putative triple gene block (TGB) associated with viral movement in plants. Putative TGB 1 (encoded from ORF 3) of the new virus isolates has a helicase domain, also found in BdMV, BNYVV and MILDV (Peltier et al., 2008; Kondo et al., 2013; Sela et al., 2016). The putative TGB 2 encoded from ORF 4 has a conserved domain associated with viral movement, as described for BdMV (Kondo et al., 2013). ORF 5 encodes a putative TGB 3 protein, like the other members of the family Benyviridae. These proteins are important for cell-to-cell and long-distance movement of the virus in plants infected with BNYVV (Gilmer et al., 1992). Given the similarities found and the conserved domains, it is suggested that the TGB discovered in isolates of the new virus is involved in cell-to-cell movement. The size of ORF 6 was the most divergent among the characterized isolates and species of the Benyviridae. The protein encoded by ORF 6 has been reported to be associated with the suppression of gene silencing (Chiba et al., 2013). The function of the hypothetical protein encoded by ORF 6 of WhSMV should be verified. The TGB and the hypothetical protein are probably expressed by subgenomic RNAs, which characterize the viruses of the Benyviridae (Gilmer et al., 2017).

The current taxonomic criterion used for species demarcation in the genus Benyivirus is amino acid sequence identity of CP <90% (Gilmer et al., 2017). Based on the genomic organization and the nucleotide and amino acid sequences identified in the viral isolates described in this work, it is proposed that wheat stripe mosaic virus can be considered a new putative species of the family Benyviridae.

Phylogenetic analyses indicated that WhSMV isolates formed a distinct clade when trees were constructed employing complete RNA segments and all used coding regions, indicating a close relationship among Brazilian isolates; this corroborates the sequence analysis data and supports the idea that the viral isolates described here belong to the same species and are distinct from other species of the genus Benyivirus.

An important biological feature of members of the Benyviridae is their transmission by parasites of the root system. BNYVV and BSBMV were reported to be transmitted by *P. betae* (Tamada et al., 1996). RSNV was detected infecting rice in the Rio Grande do Sul State and was transmitted by *P. graminis* (Morales et al., 1999). The presence of *P. graminis* in the wheat roots of TBIO Toruk and BRS Guamirim cultivars has also been confirmed in the present investigation. The morphological structures found correspond to those described for *P. graminis* by Ledingham (1939) and are similar to those observed by Tyagi et al. (2016), who analysed the roots of barley and observed the different phases of the *P. graminis* life cycle. The primer pair Pgfwd2/Pxrev7 is specific for *P. graminis* and amplification using this primer pair resulted in amplified fragments of the expected sizes 310 bp and 240 bp (Ward & Adams, 1998). Nucleotide identity and phylogenetic analyses of these DNA sequences supported the classification of the plasmidiphorid found in the wheat roots as *P. graminis*.

Phylogenetic analysis also indicated that the two isolates of *P. graminis* characterized in this study are in different groups, supporting the nucleotide differences found. These differences were expected, because molecular studies revealed high variability of *P. graminis* (Legreve et al., 2002). Another recent study in China using the ITS region of *P. graminis* ribosomal DNA from 63 samples of wheat roots indicated considerable variation of the sequences (Xu et al., 2018). The morphological, molecular and phylogenetic characterization indicated the association of *P. graminis* with roots of wheat plants infected by WhSMV in Brazil and suggested that WhSMV could be transmitted by *P. graminis*.

The dataset obtained in this study indicates that the aetiological agent of SBWMD in Brazil is a distinct virus. It cannot be ruled out that SBWVMV and other viruses occur in wheat in Brazil. However, at present, the most probable hypothesis is that the virus described by Caetano *et al.* (1978) is in fact the new species described here and that, using the tools available in the 1970s, accurate diagnosis was not possible. This hypothesis is corroborated by the fact that numerous researchers who attempted to characterize and detect Brazilian isolates in recent years with primers and antiserum for SBWMV obtained negative results. In addition, the association between WhSMV and wheat plants with SBWMD of different cultivars and areas was demonstrated. The WhSMV was detected in different wheat cultivars in four different areas (Coxilha, Passo Fundo, Santo Augusto and Ivaiporã municipalities), confirming the presence of the virus described here in all the 26 wheat samples with mosaic symptoms collected.

This new virus is so distinct from other already described viruses infecting wheat that an exciting hypothesis and/or questions about its origin are raised. In future work, it will be necessary to characterize isolates from other geographic regions and other hosts in an attempt to decipher its evolutionary history. Is it a virus that came with wheat to Brazil and then evolved in this country? Alternatively, is it a native virus that acquired the ability to infect wheat? The virus might infect other grasses and alternative hosts and further studies should be carried out to test this possibility. The fact that it is a virus transmitted by a soilborne microorganism and, therefore, has population dynamics slower than that of airborne pathogens, may have preserved populations that contain steps in the evolution of this virus.
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References


Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site.

Figure S1. Electrophoretic analyses of the PCR products for the detection of wheat stripe mosaic virus (WhSMV) using specific primers. Products amplified using primers for detection of (a) polymerase (replicase; amplicon 464 bp), (b) TGB 2 (amplicon 607 bp). TGB 2-1, (c) RNA 1 3′-UTR (amplicon 357 bp), (d) RNA 2 3′-UTR (amplicon 329 bp) and (e) coat protein (amplicon 641 bp). 1: from wheat cultivar Karl 92, 2: from wheat cultivar Everest, 3: from wheat cultivar OR-1, 4: from wheat cultivar CEP11, 6: from wheat cultivars TBIO Toruk + TBIO Tibagi, 7: from wheat cultivars Esporão + LG Oro, MM: 1 kb molecular ladder, B: negative control (water).

Figure S2. Phylogenetic relationships of the family Benyviridae, based on complete genome sequences of RNA 1 (a) and RNA 2 (b). The trees were constructed by the maximum likelihood method implemented in the MEGA v. 6.0 program (Tamura et al., 2013) using the general time reversible (GTR) model with gamma distribution (G) and bootstrap support of 6000 replications. Numbers on branches indicate bootstrap values. Alignments were performed with Muscle. Soil-borne wheat mosaic virus (SBWMV) was used as an out-group. Bar indicates the number of substitutions per site. Members of the family Benyviridae are wheat stripe mosaic virus (WhSMV), Beet necrotic yellow vein virus (BNYYV), Beet soil-borne mosaic virus (BSBMV), Burdock mottle virus (BdMV), Rice stripe necrosis virus (RSNV), and Mangifera indica latent virus (MILV).

Figure S3. Electrophoretic analyses of the PCR products for detection of wheat stripe mosaic virus (WhSMV) from wheat samples collected in different areas of Brazil, using TGB 2 primers (amplicon 607 bp). MM: 1 kb molecular ladder; C+: positive control (wheat TBIO Toruk cultivar with symptoms); C−: negative control (symptomless wheat TBIO Toruk cultivar); B: blank (water); 1–3: samples from Coxilha municipality (BRS Pastoreio, LG Oro and Amplitude cultivars, respectively); 4–12: samples from Ivaiporã municipality (Everest, OR 1, CEP 11, BRS Reponte, Esporão, TBIO Itaipu, TBIO Toruk, ORS 1401 and LG Oro, respectively); 13–15: samples from Santo Augusto municipality (BRS Reponte, ORS 1401 and LG Oro, respectively); 16–26: samples from Passo Fundo municipality (Everest, Karl 92, OR 1, CEP 11, BRS Reponte, Esporão, TBIO Itaipu, TBIO Toruk, ORS 1401, LG Oro and BRS Guaimirim, respectively).