

# Biological and Molecular Characterization of an Isolate of *Tobacco streak virus* Obtained from Soybeans in Brazil

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## ABSTRACT

A virus was isolated from soybean (*Glycine max*) plants with symptoms of dwarfing and bud blight in Wenceslau Braz County, Paraná, Brazil. The host range and properties resembled those of *Tobacco streak virus* (TSV). The purified virus showed three peaks in a frozen sucrose gradient. Antiserum was produced and the virus was serologically related to TSV. Electron microscopy detected 28 nm spherical particles. Coat protein (CP) had a Mr of 29.880 Da. A fragment of 1028 nt was amplified, cloned and sequenced. One open reading frame with 717 nt was identified and associated to the CP. The CP gene shared 83% identity with the sequence of TSV CP from white clover (*Trifolium repens*) (GenBank CAA25133). This is the first report of the biological and molecular characterization of TSV isolated from soybeans. It is proposed that this isolate be considered a strain of TSV named TSV-BR.

**Additional keywords:** nucleotide sequence, Elisa, host range.

## RESUMO

### Caracterização biológica e molecular de um isolado de *Tobacco streak virus* obtido de soja no Brasil

Um vírus foi isolado de plantas de soja (*Glycine max*) cultivadas em Wenceslau Braz, Estado do Paraná, com sintomas de nanismo e queima do broto. O vírus foi caracterizado por meio de hospedeiros diferenciais e propriedades biológicas como sendo um isolado do *Tobacco streak virus* (TSV). O vírus purificado apresentou três picos em gradiente de sacarose congelado. Anti-soro produzido contra o vírus foi sorologicamente relacionado ao TSV. Microscopia eletrônica detectou partículas esféricas com 28 nm de diâmetro. A proteína do capsídeo (PC) apresentou massa molecular de 29.880 Da. Um fragmento de 1028 nt foi amplificado, clonado e seqüenciado. Uma ORF com 717 nt foi identificada e associada com a PC, a qual compartilha 83% de identidade com a seqüência da PC do isolado de TSV de trevo branco (*Trifolium repens*) (GenBak CAA25133). Este é o primeiro relato da caracterização biológica e molecular de um isolado de TSV de soja. Propõe-se que o isolado seja considerado uma estirpe do TSV denominada TSV-BR.

**Palavras-chave adicionais:** seqüenciamento de nucleotídeos, Elisa, gama de hospedeiros.

## INTRODUCTION

A recent outbreak of soybean [*Glycine max* (L.) Merrill] bud blight in Wenceslau Braz County, State of Paraná, Brazil, may be an indication of the occurrence of a virus disease. The outbreak occurred in an area close to a region where the presence of *Tobacco streak virus* (TSV), family *Bromoviridae*, genus *Ilarvirus*, is historical. Preliminary evaluations using electron microscopy of infected leaves showed the presence of spherical particles. Additional studies proved that TSV was responsible for the problem. Although no resistance to this virus has been found in the soybean germplasm the disease has been controlled

by delaying the sowing date (Almeida & Corso, 1991; Almeida *et al.*, 1994).

The TSV was identified in tobacco (*Nicotiana tabacum* L.) plants in Brazil in 1940 (Costa, 1945), and it is currently known to infect several cash crops such as cotton (*Gossypium hirsutum* L.), tomato (*Lycopersicon esculentum* Mill.), tobacco (*Nicotiana tabacum* L.), soybean, peanut (*Arachis hypogaea* L.), sunflower (*Helianthus annuus* L.) and some weeds (Costa & Carvalho, 1961). The association of soybean bud blight and TSV was reported for the first time in 1955 (Costa *et al.*, 1955). At that time, the authors mentioned that similar symptoms were described in soybean in the USA, but these symptoms were caused by *Tobacco*

*ringspot virus* (TRSV), family *Comoviridae*, genus *Nepovirus*. The disease was called soybean bud blight by Costa *et al.* (1955). Since then, several other viruses inducing similar symptoms have been described on soybeans in Brazil (Costa, 1982), causing serious confusion in field diagnosis.

Disease symptoms caused by TSV started to appear 15-20 days after sowing, when plants exhibited browning buds which later curled downward. Yield losses of up to 100% have been reported in soybean fields in Brazil. Studies have shown that the disease is endemic in specific regions where high populations of the vectors *Thrips tabacci* Linderman and *Frankliniella occidentalis* Pergande (Kaiser *et al.*, 1982) have developed due to the simultaneous presence of a weed (*Ambrosia polystachia* L.) that is a known host of the virus (Costa & Carvalho, 1961). Another vector species (*Microcephalothrips abdominalis* Crawford) has also been reported by Greber *et al.*, 1991. However, the transmission by thrips was firstly demonstrated by Costa & Lima Neto (1976).

TSV has a tripartite single-stranded messenger-sense RNA genome. The RNAs 1 and 2 encode proteins involved in viral RNA replication, whereas RNA 3 encodes a protein required for cell-to-cell movement. The viral coat protein (CP) is expressed by a subgenomic RNA, designated RNA 4, collinear with the 3' end of RNA 3 (Bol, 1999). Studies on ilarviruses revealed that in addition to functioning as a structural protein, the CP is also involved in many steps of virus replication.

In this report we present the biological and molecular characteristics of a Brazilian isolate of TSV as well as a phylogenetic analysis of the CP gene as compared with other ilarviruses.

## MATERIALS AND METHODS

### Virus isolate and host range

A virus was isolated from soybean plants with typical symptoms of bud blight as well as from plants of *A. polystachia* with mosaic symptoms in Wenceslau Braz County. Infected leaves were ground in chilled 0.01 M sodium phosphate buffer pH 7.0 with 0.1% 2-mercaptoethanol.

Mechanical inoculation was performed in leaves previously dusted with fine charcoal. Inoculated leaves were washed with running water and maintained in a greenhouse under temperatures ranging from 18 °C to 35 °C, depending on the season. For the host range experiment plants were kept in sterilized soil with four plants per pot and two pots of each botanical species were inoculated. Plants were assayed two and four weeks after inoculation by symptomatology and indirect ELISA (Koennig, 1981).

### Virus purification, nucleoprotein analysis and western blotting

Infected leaves from tobacco cv. Sansun 'NN' were ground in 0.01 M potassium phosphate buffer, pH 8.0 (2:1

w/v) containing 1% 2-mercaptoethanol. The slurry was filtered in a double cheesecloth, clarified with chloroform (1/2 vol) and stirred for 20 min at 4 °C. The extract was centrifuged at 10,000 g for 10 min at 4 °C and the supernatant was mixed with 6% PEG 8,000 at 4 °C for 2 h. After low speed centrifugation (10,000 g for 10 min) the pellet was dissolved overnight at 4 °C. After another low speed centrifugation the supernatant was centrifuged at 180,000 g for 2 h in a swinging bucket rotor using 25% sucrose frozen gradient (Baxter-Gabbard, 1972; Davis & Pearson, 1978). The virus was fractionated using an ISCO density gradient fractionator and UV analyzer. Fractions were diluted in 0.01 M potassium phosphate buffer, pH 8.0 and centrifuged at 100,000 g for 90 min. The pellet was dissolved and centrifuged at 10,000 g for 10 min and the supernatant rescued and stored. Virus yield was determined by assuming an extinction coefficient of 5.1 (Salazar *et al.*, 1982).

The molecular weight of the CP was estimated by sodium dodecyl sulfate – polyacrylamide gel electrophoresis (SDS-PAGE) on a 3.5% stacking gel and a 12% resolving gel. A mixture of an equal amount of virus preparation and dissociation buffer (0.125 M Tris-HCl, pH 6.7, 3% SDS, 20% glycerol and 10% 2-mercaptoethanol) was boiled for 5 min. After electrophoresis for 2 h at 120 V at room temperature, proteins were stained with coomassie brilliant blue. Protein molecular weight was estimated by comparing their relative mobility with molecular weight standards (Low molecular weight calibration kit-Amersham Biosciences, NJ, USA).

For western blot analysis, the protein was blotted onto nitrocellulose membrane (Sigma N-8142) and transferred by a Hoefer MiniVE device (Amersham Pharmacia, NJ, USA) at 300 mA for 90 min. Immunostaining was carried out according to Lenardon *et al.* (1993). A TSV antiserum was provided by W. J. Kaiser, Washington State University, Pullman, WA, USA. For virus detection alkaline phosphatase conjugated with goat-antirabbit antibodies (A 8025; Sigma, USA) diluted to 1:5,000 and substrate BCIP/NBT (Sigma B-1911) was used.

### Antisera production and serological determinations

Two New Zealand rabbits were immunized with intramuscular injections of 100 µg of purified virus, which had been suspended in saline (0.15 M NaCl) and emulsified in an equal volume of Freund's complete adjuvant. Second, third and fourth injections were prepared with incomplete adjuvant, containing the same amount of virus at ten-day intervals. Bleedings began four weeks after the first injection and continued for six weeks. Antisera were stored at -5°C.

### Enzyme-linked immunosorbent assay analysis

Immunoglobulin was prepared according to Clark & Bar-Joseph (1984) using ammonium sulphate precipitation. Virus diagnosis was done by indirect ELISA (Koenig, 1981). The uppermost fully developed soybean leaves were ground in ELISA extraction buffer (4 ml of

buffer/ g leaf). Microtiter plates were covered with antigen previously prepared in coating buffer at 1:200 dilution and incubated for 2 h at 37 °C. Immunoglobulin was used at 0.8 µg/ml. Goat anti-rabbit IgG alkaline phosphatase conjugate (A-8025, Sigma Immuno-Chemicals, USA) was used at 1:8,000 dilution and incubated for 2 h at 37 °C. Twenty minutes after adding the substrate (p-nitrophenil phosphate) absorbance ( $A_{405\text{ nm}}$ ) was read on a ELISA reader (Microplate Reader, Mod. 3550 UV, BioRad, USA). A reaction was considered positive if the reading exceeded two times the mean value for healthy tissue.

### Electron microscopy

Viral particles were also observed in purified preparations negatively stained with uranyl acetate (Kitajima & Nome, 1999).

### RNAs extraction and reverse transcription.

Infected young tobacco 'Sansun' leaves were frozen with liquid nitrogen and pulverized in a mortar. Nucleic acid preparations enriched in dsRNAs were obtained by extraction with buffer-saturated phenol and fractionated by column chromatography on non-ionic cellulose (CF-11, Whatman) as described by Dodds & Bar-Joseph (1983). Reverse transcription was done using CF11-prepared RNA as a template. Ten µl RNA samples were taken and mixed with 2 µl (10 µM) of primer 3TbS3 (5'-GCATCTCCTATAA AGGAGGC-3'), incubated at 80 °C for 3 min and quenched on ice. In sequence, other reaction components were added in the following order: 6 µl of 5X RT first strand buffer (200 mM Tris-HCl, pH 8.4), 1 µl of dNTP mix (10 mM), 2 µl of DTT (0.1M), 2 µl M-MLV reverse transcriptase (20 U) (Gibco BRL), and DEPC treated water to give a reaction volume of 30 µl. The tubes were incubated for 1 h at 37 °C on a thermocycler MJ model PTC 200 (MJ Research Inc., Waltham, MA, USA) followed by 5 min of heating at 99 °C to denature the enzyme.

### Amplification of coat protein gene by PCR

The PCR reactions were performed in 50 µl volume containing 5 µl of PCR buffer (10X), 5 µl of MgCl<sub>2</sub>, 10 µl of dNTP mix (10 mM), 1 µl Taq DNA polymerase (5U/µl), 5µl reverse transcribed sample, 20 µl DEPC water and 2 µl of each primer (10 µM) 3TbS3 and TbCP5U (5'-GCTTCTC GGACTTACCTGAG-3'). The reaction mixture was heated at 94 °C for 3 min followed by 35 cycles at 94 °C for 30 sec, 55 °C for 1 min and 72 °C for 2 min followed by 10 min incubation at 72 °C. The PCR amplified fragment was cloned into a plasmid vector, pBluescript II KS+ (STRATAGENE) according to the manufacturer's protocol.

### Nucleotide sequencing, sequence comparison and phylogenetic analysis

Nucleotide sequences from three clones were performed by comparison with those available in GenBank and EMBL databases. Multiple sequence alignments and

dendrograms of sequence relationship were produced using CLUSTAL W (Thompson *et al.*, 1994). The translation product was estimated through the translate routine from ExPasy software (<http://us.expasy.org/tools/>). The open reading frame was evaluated through the <http://www.ncbi.nih.gov/gorf/gorf.html>. The calculation of identity and similarity was performed by analysis on [http://bioinformatics.org/sms/ident\\_sim.html](http://bioinformatics.org/sms/ident_sim.html). The phylogenetic relationships were established by maximum parsimony, with software PAUP v.4.0b10. The reliability of the tree was examined by a bootstrap test with 1000 replicates.

## RESULTS

### Virus isolate and host range

After mechanical inoculation several plant species from seven families were infected by the TSV isolate used in this study (Table 1). The TSV infected 18 out of 26 plant species. Soybean plants reacted with the same symptoms observed in the field, exhibiting bud blight seven to ten days after inoculation. No symptoms were observed in *Amaranthus* sp., abobrinha (*Cucurbita pepo* L.), *Pisum sativum* L., *Crotalaria pallida* Ait, and *C. spectabilis* Roth. These inoculated plants were indexed by mechanical inoculation in *N. tabacum* cv. Samsun and no symptoms were observed.

### Virus purification and nucleoprotein analysis

The purification method yielded 1.4-2.6 mg virus/ 100 g of leaves. The 260/280 ratios ranged from 1.43 to 1.56, typical of spherical viruses. Viral preparation showed a specific profile after sucrose gradients typical of ilarviruses with three peaks corresponding to top, medium and bottom components.

The CP migrated in SDS-PAGE as a single band with a molecular mass of 29.880 kDa (Figure 1A). Western blots that were probed with TSV polyclonal using antiserum prepared by W. J. Kaiser (Washington State University, USA) showed a single band in the same position at the SDS-PAGE (Figure 1B).

### Antisera production and serological determinations

Antibodies raised to TSV on rabbits reacted lightly with healthy soybean sap and were used in indirect ELISA only after cross absorption with healthy plant sap (Figure 2). Cross absorbed IgG decreased the absorbance values without significantly affecting the detection of TSV.

### Electron microscopy

Electron microscope analysis of infected leaves showed spherical virus particles measuring 28 nm diameter (Figure 3).

### Nucleotide sequencing, sequence comparison and phylogenetic analysis

The PCR tests using specific primers for the TSV

**TABLE 1** - Symptoms induced in plants by *Tobacco streak virus*-Brazilian strain, isolated from soybean (*Glycine max*)

Botanical family	Species	Reaction <sup>1</sup>
Amaranthaceae	<i>Gomphrena globosa</i> L.	CLL
	<i>Amaranthus</i> sp.	-
Asteraceae	<i>Emilia sonchifolia</i> (L.) DC	M
	<i>Bidens pilosa</i> L.	M
Chenopodiaceae	<i>Chenopodium amaranticolor</i> Coste & Reyn.	NLL
	<i>C. quinoa</i> Willd.	CLL
Cucurbitaceae	<i>Cucurbita pepo</i> L. 'Caserta'	-
Fabaceae	<i>Glycine max</i> L. Mer.	
	c.v. Santa Rosa	SN
	c.v. Davis	SN
	<i>Phaseolus vulgaris</i> L.	
	c.v. Rosinha	NLL
	c.v. Carioca	NLL
	c.v. Tibagi	CLL
	<i>Lupinus albus</i> L.	M
	<i>Crotalaria pallida</i> Ait	-
	<i>C. spectabilis</i> Roth.	-
Gramineae	<i>Arachis hypogaea</i> L.	M
	<i>Vigna unguiculata</i> (Walp.) Walp. Pitiyba	-
	<i>Pisum sativum</i> L.	-
	<i>Zea mays</i> L.	-
	<i>Lycopersicon esculentum</i> Mill.	M
Solanaceae	<i>N. tabacum</i> L. Sansun NN	WCN
	<i>N. glutinosa</i> L.	WCN
	<i>N. debneyi</i> Domin.	M
	<i>N. benthamiana</i> Domin.	M
	<i>Gossypium hirsutum</i> L.	M

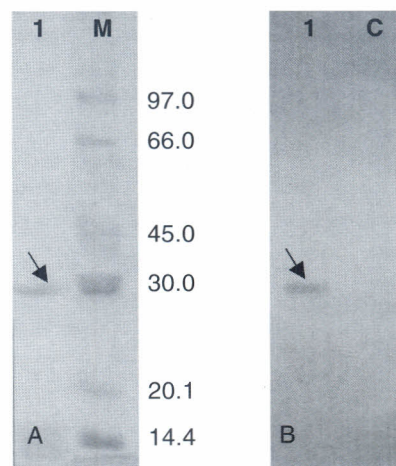
<sup>1</sup>WCN= white circle necrosis; M= mosaic; NLL= necrotic local lesion; CLL= chlorotic local lesion;

isolate from white clover (*Trifolium repens* L.) (TSV WC) yielded a fragment of 1028 nt. The fragment included the gene of the 717 nucleotide CP (ORF) and a 287 nucleotide 3' untranslated region. The determined sequence was deposited at GenBank under the accession number AY354406.

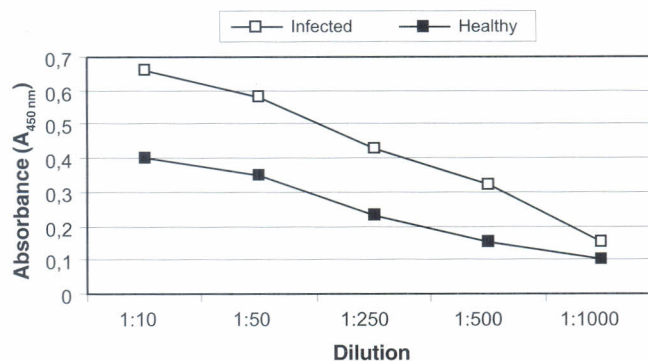
The initiation codon for the CP ORF was at position 25, and translation of this ORF resulted in a polypeptide of 238 amino acids which ends at a UAG termination codon at position 741. The predicted molecular weight of the protein is 30,563 Da, slightly larger than the 29,880 Da determined from the SDS-PAGE (Figure 1A). The deduced amino acid sequence shares 82.8% identity with TSV isolated from white clover (GenBank NC 3845) (Cornelissen *et al.*, 1984) (Table 2).

Phylogenetic analysis was performed and the trees for nucleotides and amino acids showed similar results (Figure 4). The TSV-BR, TSV- WC (this is designated as TSV-NC in Figure 4 and needs to be changed there) and TSV- mungbean [*Vigna radiata* L. (Wilzed)] (AF515823) cluster in the same branch. The TSV-BR was more distantly related with SNSV and HdMV and had a limited homology with other ilarviruses compared.

Analysis of the CP amino acid sequence exhibited the cysteine motif C-X<sub>2</sub>-C-X<sub>10</sub>-C-X<sub>2</sub>-H in the amino acid positions 26 to the 44 which can form a possible "zinc finger" type-binding domain (Figure 5).



**FIG. 1** - (A) SDS-PAGE of proteins from dissociated *Tobacco streak virus* (TSV) on 12.5% polyacrilamide gel. Lane 1, TSV-BR (29,8kDa). Lane M, molecular weight markers (LMW Marker Kit; Amersham Biosciences). B= western blot from the gel (A) of infected (1) and healthy extract (C).



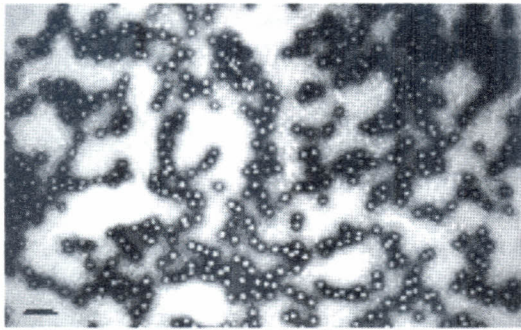
**FIG. 2** - Indirect ELISA reaction against a polyclonal *Tobacco streak virus* (TSV) antiserum with infected and healthy samples of soybean (*Glycine max*) leaves, absorbed with healthy tissue.

The 287 nucleotide 3' untranslated region (UTR) of TSV-BR RNA 3 shows a high degree of similarity with TSV-WC (85%) and SNSV (70.3%) (data not shown). It contains a potential stem-loop structure with interspersed AUGC motifs, characteristic for ilarvirus and alfamovirus (Figure 6).

## DISCUSSION

The virus analyzed in these studies shows the biological and molecular properties similar to those described for TSV (Fulton, 1971). Therefore, the outbreak of soybean bud blight that occurred in Wenceslau Braz County was definitely caused by an isolate of TSV.

The host range data is in agreement with several reports for TSV (Costa & Carvalho, 1961; Salazar *et al.*, 1982; Kaiser *et al.*, 1982), despite the lack of infection of

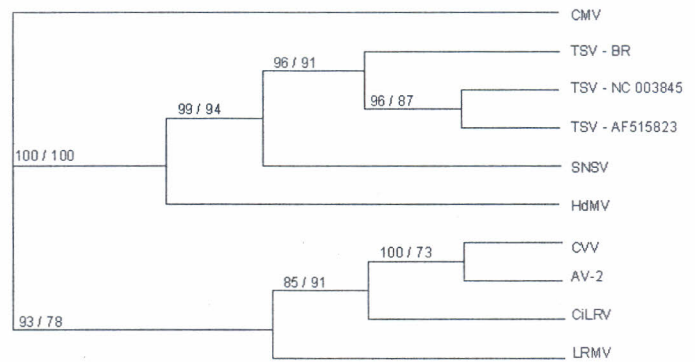


**FIG. 3** - Transmission electron micrograph of purified virus particles of TSV-BR. Barr = 120 nm.

*Amaranthus* sp., *Crotalaria striata*, *C. mucronata*, *C. spectabilis*, cowpea [*Vigna unguiculata* (L.) Walp.] and *P. sativum* previously mentioned as susceptible species by Costa & Carvalho (1961).

Profiles from sedimentation of virus particles on sucrose frozen gradients were typical of ilarvirus, particularly TSV (Lister & Bancroft, 1970). Virus purifications were always done using infected leaves from tobacco cv. Sansun 'NN'. However, the yield of virus as well as the proportion of each component varied among purifications throughout the year. Lister & Bancroft (1970) also reported that alterations in TSV component ratios and yield were influenced by hosts and extraction procedure. In our case, only one species of tobacco and one method were used, and therefore, it is believed that other factors such as the effect of environmental conditions on plants could be responsible for the differences. For example, temperatures in the greenhouse ranged from 16°C in the winter up to 38°C in the summer.

The CP from purified virus had a relative molecular mass close to those reported for ilarviruses (<http://image.fs.uidaho.edu/vide/descr811.htm>). Amino acid analysis of the protein showed a molecular mass of 30,563 Da, close to the values obtained from electrophoresis analysis



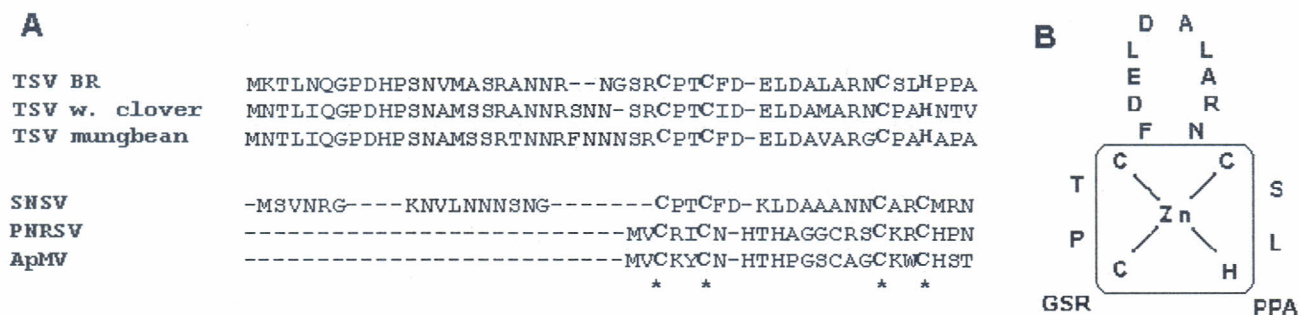
**FIG. 4** - Phylogenetic analysis produced by Paup v.4.0b10 program, using the parsimony maximum method, indicating relationships among isolates of *Tobacco streak virus* (TSV) and others members of *Ilarvirus* genus. The analysis was made based on the complete sequence of coat protein. Numbers at each ramification indicated bootstrap percentages (based on 1,000 replications) for data from nucleotide and deduced amino acid analysis, respectively (nt/ aa). The sequences for comparison were obtained from GenBank with accession numbers for nucleotide and deduced amino acids respectively: TSV - white clover (*Trifolium repens*) (NC003845/CAA25133) TSV- mungbean (*Vigna radiata*)(AF515823/AAM76049), *Strawberry necrotic shock virus* - SNSV (AY363242/AAQ76590). *Hydrangea mosaic virus* - HdMV (U35145/AAA80345), *Spinach latent virus* - SpLV (NC003810/NP620681), *Citrus leaf rugose virus* - CiLRV (NC3546/ NP613280), *Lilac ringspot mosaic virus* - LRMV (U17391/AAA64840), *Citrus variegation virus* - CVV (AF434918/ AAL37957), *Asparagus virus 2* - AV-2 (X86352/ S54138).

(29,880 Da). In addition, particle measurements estimated by electron microscopy showed average sizes (28 nm) similar to those described for TSV.

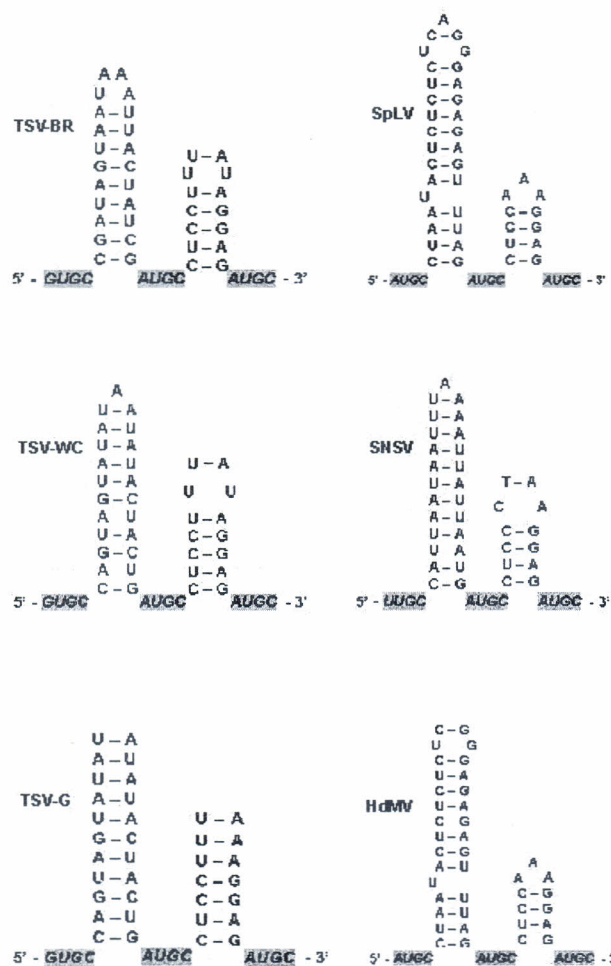
The nucleotide sequence of the RNA3 from clones TSV-BR28, TSV-BR29 and TSV-BR39 permitted identification of a 1028 nt sequence that revealed one open reading frame flanked by a 25 nucleotide portion of the intercistronic region and a 287 nucleotide 3'-NTR. The folded tRNA-like motif of the 3'-NTR observed in this isolate

**TABLE 2** - Comparison on the percentage of identity among nucleotide sequences (below diagonal) and deduced amino acids (above diagonal) of the CP gene of *Tobacco streak virus* (TSV) and species of *Ilarvirus* genus deposited at GenBank: TSV<sup>(1)</sup> (Brazilian isolate, AY354406), TSV<sup>(2)</sup> [TSV-isolado de mungbean (*Vigna radiata*)(AF515823)], TSV<sup>(3)</sup> (NC003845), *Strawberry necrotic shock virus* (SNSV) (AY363242), *Hydrangea mosaic virus* (HdMV) (U35145), *Citrus variegation virus* (CVV) (AF434918), *Asparagus virus-2* (AV-2) (X86352), *Lilac ring mottle virus* (LRMV) (U17392), *Citrus leaf rugose virus* (CiLRV) (NC003546)

Nt/na	TSV <sup>1</sup>	TSV <sup>2</sup>	TSV <sup>3</sup>	SNSV	HdMV	CVV	AV-2	LRMV	CiLRV
TSV <sup>1</sup>		80.8	82.8	62.3	46.9	18.4	18.4	15.7	17.3
TSV <sup>2</sup>	80.7		89.8	63.3	47.3	18.4	18.8	17.3	17.7
TSV <sup>3</sup>	81.3	88.9		61.9	46.7	17.6	18.4	16.1	17.0
SNSV	63.9	65.7	64.8		44.7	22.7	22.3	17.4	19.1
HdMV	56.8	58.6	58.1	53.8		20.5	20.5	16.3	18.1
CVV	41.0	41.0	40.7	40.1	40.1		75.1	34.1	63.5
AV-2	39.7	40.3	39.9	40.9	40.0	76.3		34.1	59.5
LRMV	31.2	31.8	32.2	31.1	30.5	48.0	49.7		36.1
CiLRV	37.6	37.7	38.1	37.4	38.3	65.1	62.0	47.6	



**FIG. 5 - A:** Alignment of amino acid sequences of the CP N-terminal region of several *Tobacco streak virus* (TSV) isolates and other ilarviruses. The positions of the conserved cysteine amino acids in the CP are marked with asterisks. **B:** Putative “zinc finger” motif present at the amino-terminus of the TSV-BR CP. The sequences for comparison were obtained from GenBank with accession numbers for deduced amino acids: TSV white clover (*Trifolium repens*) (CAA25133); TSV mungbean (*Vigna radiata*) (AAM76049); TSV cotton (*Gossypium hirsutum*) (AAM76050); TSV sunn-hemp (*Crotalaria juncea*) (AAM76051); TSV sunflower (*Helianthus annuus*) (AAL31701); SNSV (AAQ76588); *Prunus necrotic shock virus* (PNRSV) (NP733825); *Apple mosaic virus* (ApMV) (NP604485).



**FIG. 6 - Possible secondary structures at the 3'-termini of *Tobacco streak virus* brazilian isolate (TSV- BR) and other ilarviruses. The conserved AUGC sequences are boxed and shaded. Structures were designed based on sequence data from GenBank: TSV - white clover (*Trifolium repens*) (NC003845), TSV - G [groundnut (*Arachis hypogaea*), Cook *et al.*, 1999], *Spinach latent virus* (SpLV) (NC003810), *Citrus leaf rugose virus* (CiLRV) (NC3546), *Hydrangea mosaic virus* - HdMV (U35145), *Strawberry necrotic shock virus* (SNSV) (AY363242).**

is predicted to have an important role in the virus infection by binding the CP as it does for other ilarviruses and alfamoviruses (Vloten-Doting, 1975; Bol, 1999). The ORF codes for a Mr 29,880 product that corresponds to the viral CP. This value is larger than the Mr 26,346 described by Cornelissen *et al.* (1984) for the CP of an isolate of TSV obtained from white clover but is close to the value mentioned by Salazar *et al.* (1982). The CP of this strain shares an 82% amino acid sequence identity with that of the TSV strain WC.

Parsimony phylogenetic analyses for nucleotide sequences were obtained. The nucleotide sequence of this virus showed close association to TSV-WC (NC003845). Based on the proposal of Regenmortel *et al.* (1997) and Bhat *et al.* (2002) and assuming a threshold level of 90% similarity in the CP sequence, we consider the soybean virus to be a new strain of TSV isolated in Brazil.

A cysteine motif C-X<sub>2</sub>-C-X<sub>10</sub>-C-X<sub>2</sub>-H that may form a “zinc finger” type-binding domain as described by Berg (1986) is found in the TSV-BR CP sequence can be between the 28<sup>th</sup> and 46<sup>th</sup> amino acids, as it is for all sequenced isolates (Sehnke *et al.*, 1989). A similar structure also occurs in SNSV, APMV and PNRSV (Guo *et al.*, 1995) but is located in the position of amino acids 2 to 13 and presents C-X<sub>2</sub>-C-X<sub>10</sub>-C-X<sub>2</sub>-C pattern.

The N-terminal region of the CP is highly conserved in sequenced TSV isolates and has been shown to be necessary for binding the CP to the 3' end of ilarvirus and AMV RNAs (Baer *et al.*, 1994). This indicates the possibility of involving a process of genome activation and/or interaction of the protein with genomic RNAs for virion formation.

The folded stem-loop structure of the 3'NTR is flanked by tetranucleotide AUGC motifs in this isolate as well as in other ilarviruses and AMV, and this motif has an important role in the virus infection by binding to CP (Vloten-Doting, 1975; Bol, 1999). Although several AUGC motifs are conserved among ilarviruses and AMV RNAs,

TSV isolates have two AUGC motifs as well as third GUGC motif in a proximal position in the 3' terminal region.

Nucleotide and amino acid sequences of TSV-BR present serological, ultrastructural and biological characteristics that demonstrate structural and molecular similarities of TSV-BR with other TSV isolates, indicating a close relationship between them. This is the first report of the biological and molecular characterization of TSV from soybean, and we believe that the PCR primers designed for use in this work will be helpful as a primary diagnostic tool for TSV-BR.

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