Sequence diversity in the coat protein gene of *Lettuce big-vein associated virus* and *Mirafiori lettuce big-vein virus* infecting lettuce in Brazil.

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**ABSTRACT**

*Lettuce big vein associated virus* (LBVaV) e *Mirafiori lettuce big vein virus* (MLBVV) têm sido encontrados em infecções mistas no Brasil, causando a doença conhecida como engrossamento das nervuras da alface. Análise de parte do gene da proteína capsidial (CP) de isolados brasileiros de LBVaV coletados em alface, indicou que estes possuem identidade superior a 93% com isolados coletados em diferentes regiões geográficas.

**Additional keywords**: LBVaV, MLBVV, *Olpidium brassicae*.

**RESUMO**

*Lettuce big vein associated virus* (LBVaV) e *Mirafiori lettuce big vein virus* (MLBVV) têm sido encontrados em infecções mistas no Brasil, causando a doença conhecida como engrossamento das nervuras da alface. Análise de parte do gene da proteína capsidial (CP) de isolados brasileiros de LBVaV coletados em alface, indicou que estes possuem identidade superior a 93% com isolados coletados em diferentes regiões geográficas.

**Palavras-chave adicionais**: LBVaV, MLBVV, *Olpidium brassicae*.

**ABSTRACT**

*Lettuce big vein associated virus* (LBVaV) and *Mirafiori lettuce big vein virus* (MLBVV) have been found in mixed infection in Brazil causing the lettuce big vein disease. Analysis of part of the coat protein (CP) gene of Brazilian isolates of LBVaV collected from lettuce, showed at least 93% amino acid sequence identity with other LBVaV isolates. Genetic diversity among MLBVV CP sequences was higher when compared to LBVaV CP sequences, with amino acid sequence identity ranging between 91% to 100%. Brazilian isolates of MLBVV belong to subgroup A, with one unique restriction site RsaI in the gene. There is no indication for a possible geographical origin for the Brazilian isolates of LBVaV and MLBVV.

**Additional keywords**: big-vein, LBVaV, MLBVV, *Olpidium brassicae*.

Lettuce big vein disease was first described in California (5), and is a soil-borne disease found worldwide (9). The name of the disease refers to the appearance of chlorotic regions surrounding the vascular tissue that confers the aspect of an anomalous vein enlargement, commonly accompanied by severe leaf deformations and growth reductions. The economic importance of the disease is a result of the unsightliness of the lettuce foliage, which depresses market value, delays head formation, decreases head size and reduces the proportion of harvestable plants (13).

The big vein symptoms were historically attributed to *Lettuce big-vein associated virus* (LBVaV; genus Varicosavirus), formerly known as *Lettuce big-vein virus* (LBVV), but a causative relationship was never confirmed. According to Roggero et al. (9), the *Mirafiori lettuce big-vein virus* (MLBVV), formerly known as Mirafiori lettuce virus (MiLV), which belongs to the genus Ophiovirus, was reported to be the causal agent of big-vein disease. LBVaV and MLBVV are both transmitted by the soil-borne fungus *Olpidium brassicae* (6).

The resting spores of the fungus can persist for over 20 years in soil and can retain the ability to transmit the disease for over 15 years (2). In Brazil the disease was reported on 2003 and its occurrence was associated to cooler season (3). Frequently plants exhibiting big vein symptoms are coinfected with both viruses (3, 7, 9, 11), but simple infection by LBVaV has also been observed, indicating that LBVaV could also contribute in some manner to the disease (7, 9, 11).

In order to give additional knowledge of the big vein disease occurring in Brazil, the coat protein gene from three MLBVV and four LBVaV isolates collected from lettuce producing areas of São Paulo State, was sequenced and compared with others found in different
areas of the world.

Old leaf pieces from symptomatic lettuce plants were collected from the lettuce producing areas surrounding the cities of Mogi das Cruzes, Campinas and Bauru, São Paulo State during 2004 and 2005. Total RNA extraction was performed as described by Bertheau et al. (1) using leaves stored at 4°C. The RT-PCR assay was carried out in a one-step procedure, using the primer pairs MiLV CP1/CP2 and MiLV CP2/CP3 and LBVV CP1/CP2 and LBVV CP3/CP4 covering the complete nucleotide sequence of the coat protein of MLBVaV and LBVaV, respectively (10). The RT-PCR was made in a 50 ml reaction volume containing 0.75 U AMV reverse transcriptase (Promega Corp., USA); 5 ml of RNA total as template; 10 ml of GoTaq DNA Polimerase Buffer 5x (Promega Corp., USA); 0.5 mM each of primer pairs; 0.5 mM dNTPs; 1.25 U Go Taq DNA Polimerase (Promega Corp., USA). RT-PCR amplification conditions included a reverse transcription at 42°C for 30 min, initial denaturation at 95°C for 5 min followed by 40 cycles of denaturation at 92°C for 30 sec, annealing at 50-54°C for 40 sec, extension at 72°C for 40 sec and final extension at 72°C for 10 min. For primer pairs MiLV CP1/CP2, LBVV CP1/CP2 and LBVV CP3/CP4 annealing was performed at 54°C and for primer pair MiLV CP3/CP4 at 50°C.

The PCR fragments were purified by Wizard SV Gel and PCR Clean-Up System (Promega Corp., USA) according to manufacturer’s recommendations and directly ligated into plasmid pGEM-T Easy Vector (Promega Corp., USA), prior to sequence. Nucleotide and predicted amino acids sequences were aligned using the CLUSTAL W program (12). At aminoacid levels, identities varied from 93% to 99% for LBVaV, and 91 to 100% for MLBVV. Phylogenetic analysis was done using MEGA version 3.1 (www.megasoftware.net) by rooting the trees with the CP gene of the Cytorhabdovirus Northern cereal mosaic virus (NCMV) and the Ophiovirus Citrus psorosis virus (CpsV) for LBVaV and MLBVaV, respectively.

Figure 1. Phylogenetic analysis of Lettuce big vein associated virus (LBVaV) (left) and Mirafiori lettuce big vein virus (MLBVV) (right) isolates based on their CP nucleotide sequences. Trees were constructed with neighbor-joining method with 2000 bootstrap replicates. Internal branches with bootstrap replicates less than 50% were not indicated. Outgroup: Northern cereal mosaic virus (NCMV) (AB030277) for LBVaV and Citrus psorosis virus (CpsV) (AF036338) for MLBVV and its relationship with the isolates of MLBVV from Japan (JAP1, BD418581), Netherlands (LS301-O, AF525935; HOL2 (AY581693), Spain (MUR1, AY366415; GAL1, AY366416; ALM1, AY366417; ALM2, AY366418; ALM3, AY366700; ALM4, AY366701; ALM5, AY366702; SON3, AY366706), Chile (CH 01), Australia (AUS1, AY366706), Denmark (DEN1, AY366702), England (UK1, AY366704), Germany (GER1, AY366705; GER2, AY366706; GER3, AY366707), Italy (ITA1, AY366708), Brazil-Mogi das Cruzes (BR61, DQ530356), Brazil-Campinas (BR60, DQ530357), Brazil-Bauru (BR58, DQ530358) and LBVaV isolates from Japan (Wa, AB190528; Hy, AB190527; A, AB190526; JAP5, AB502725), Netherlands (NL, AY496056), Spain (ALM7, AY366788; MUR1, AY366411; ALM2, AY366413; ALM3, AY366784; ALM4, AY366785; ALM5, AY366786; GAL1, AY366124; SON1, AY366128; MUR2, AY366141; GRA1, AY366142; ALM6, AY366147; ALM7, AY366151), Brazil-Campinas (BR102, DQ530353) and Brazil-Bauru (BR105, DQ530354; BR107, DQ530352).
Variability in the nucleotide and amino acid sequences of the CP gene of LBVaV isolates from São Paulo State and other countries was low. No correlation of isolates and geographic origin was revealed, although Japanese LBVaV isolates formed a group with 70% bootstrap support (Figure 1). These results agree with those described by Hayes et al. (4) that the selective pressure to preserve particular biological functions is stronger than adaptation to specific geographic areas.

The variability in the nucleotide and amino acid sequences of the CP gene of MLBVV isolates is higher than variability of LBVaV isolates. Restriction sequence analysis of the coat protein of MLBVaV Brazilian isolates revealed only one RsaI site, indicating that these isolates belong to the Subgroup A, according to Navarro et al. (8) classification. The MLBVV isolates formed two subgroups with different restriction sites for RsaI, which do not differ in symptom development. The subgroup B is formed only with Spanish isolates (ALM1, ALM4, ALM5 and SON3) previously described by Navarro et al. (8). These results support the suggestion of Hayes et al. (4) that research on big vein disease in the United States, Europe and Japan will likely be relevant for lettuce production in other areas as well, due to nucleotide sequence conservation among lettuce isolates of these pathogens, including countries on South America. Understanding the nucleotide sequence diversity of MLBVV may represent a contribution for future big-vein research in Brazil.

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