ICTV Virus Taxonomy Profile: Ophioviridae

María Laura García,1,* Elena Dal Bo,2 John V. da Graça,3 Selma Gago-Zachert,4 John Hammond,5 Pedro Moreno,6 Tomohide Natsuaki,7 Vicente Pallás,8 Jose A. Navarro,8 Carina A. Reyes,1 Gabriel Robles Luna,1 Takahide Sasaya,9 Ioannis E. Tzanetakis,10 Anna María Vaira,11 Martin Verbeek12 and ICTV Report Consortium

Abstract

The Ophioviridae is a family of filamentous plant viruses, with single-stranded negative, and possibly ambisense, RNA genomes of 11.3–12.5 kb divided into 3–4 segments, each encapsidated separately. Virions are naked filamentous nucleocapsids, forming kinked circles of at least two different contour lengths. The sole genus, Ophiovirus, includes seven species. Four ophioviruses are soil-transmitted and their natural hosts include trees, shrubs, vegetables and bulbous or corm-forming ornamentals, both monocots and dicots. This is a summary of the International Committee on Taxonomy of Viruses (ICTV) Report on the taxonomy of the Ophioviridae, which is available at http://www.ictv.global/report/ophioviridae.

Table 1. Characteristics of the family Ophioviridae

<table>
<thead>
<tr>
<th>Typical member</th>
<th>citrus psorosis virus P-121 (RNA1 AY654892; RNA2: AY654893; RNA3: AY654894), species Citrus psorosis virus, genus Ophiovirus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Virion</td>
<td>Non-enveloped, nucleocapsids 3 nm in diameter, and 700 nm or 2000 nm long. Pseudo-linear duplex structures are 9–10 nm in diameter</td>
</tr>
<tr>
<td>Genome</td>
<td>11.3–12.5 kb of negative-sense, segmented RNA (3 or 4 segments)</td>
</tr>
<tr>
<td>Replication</td>
<td>Not characterized</td>
</tr>
<tr>
<td>Translation</td>
<td>From mRNAs, which are complementary to the vRNAs</td>
</tr>
<tr>
<td>Host range</td>
<td>Citrus, blueberry, pittosporum, lettuce, sowthistle, tulip, ranunculus, anemone, lachenalia and freesia</td>
</tr>
<tr>
<td>Taxonomy</td>
<td>One genus, Ophiovirus, including seven species</td>
</tr>
</tbody>
</table>

VIRION

Particles are non-enveloped, naked filamentous nucleocapsids forming either circles (open form) or pseudo-linear duplex (collapsed form) (Table 1, Fig. 1).

GENOME

The genome of ophioviruses consists of three or four individually encapsidated RNA segments (Fig. 2). Members of the species Citrus psorosis virus [1], Ranunculus white mottle virus, Freesia sneak virus and Blueberry mosaic associated virus.

---

Received 10 May 2017; Accepted 12 May 2017

Author affiliations: 1Instituto de Biotecnología y Biología Molecular, Universidad de La Plata, La Plata, Argentina; 2Facultad de Ciencias Agrarias y Forestales, Universidad de La Plata, La Plata, Argentina; 3Texas A and M University-Kingsville, Citrus Center, Weslaco, USA; 4Department of Molecular Signal Processing, Leibniz Institute of Plant Biochemistry, Halle (Saale), Germany; 5U.S. Department of Agriculture, Agricultural Research Service, Beltsville, Maryland, USA; 6Centro de Protección Vegetal y Biotecnología, Instituto Valenciano de Investigaciones Agrarias, Moncada, Valencia, Spain; 7Faculty of Agriculture, Utsunomiya University, Utsunomiya, Japan; 8Instituto de Biología Molecular y Celular de plantas (IBMCP), Universidad Politécnica de Valencia-Consejo Superior de Investigaciones Científicas, Valencia, Spain; 9Department of Planning and Coordination, National Agriculture and Food Research Organization, Tsukuba, Japan; 10Department of Plant Pathology, Division of Agriculture, University of Arkansas, USA; 11Institute for Sustainable Plant Protection (IPSIP) – CNR, Torino, Italy; 12Wageningen Plant Research, Wageningen University and Research, Wageningen, The Netherlands.

*Correspondence: María Laura García, marialauragarcia2010@gmail.com

Keywords: Ophioviridae; ICTV; taxonomy; citrus psorosis virus; Miraflor lettuce big-vein virus; lettuce ring necrosis virus; blueberry mosaic associated virus.

Abbreviations: CP, coat protein; RdRp, RNA-dependent RNA polymerase.

---

This is an open access article under the terms of the http://creativecommons.org/licenses/by/4.0/, which permits unrestricted use, distribution and reproduction in any medium, provided the original author and source are credited.
virus have three RNAs (named RNA1, RNA2 and RNA3) whereas members of the species *Mirafiori lettuce big-vein virus* and *Lettuce ring necrosis virus* have a fourth RNA (RNA4) [2]. The genome organization is unknown for members of the species *Tulip mild mottle mosaic virus*.

Virions encapsidate both the minus- and positive-sense RNAs, but a larger amount of RNA of negative polarity is detected. As virions appear circularized, the presence of pan-handle structures has been suggested. RNA1 contains two ORFs in the viral complementary strand (vcRNA) encoding the 22–25K protein and the RdRp. The 24K protein of citrus psorosis virus is involved in the misprocessing of miRNA and RNA silencing suppressor activity. vcRNA2 encodes the cell-to-cell movement protein (MP), which also has RNA silencing suppressor activity. *Mirafiori lettuce big-vein virus* has a second putative ORF in vRNA2 that encodes a protein of unknown function. The coat protein (CP) is encoded by vcRNA3. A fourth genomic RNA has been reported for *Mirafiori lettuce big-vein virus* and *lettuce ring necrosis virus*, encoding putative proteins of unknown function.

**TAXONOMY**

*Ophiovirus* is the only genus in the family, including seven species. CP amino acid sequence alignments show 31–52% identity between isolates of different species, and 92–100% identity within a species. The CPs of *Mirafiori lettuce big-vein virus* and tulip mild mottle mosaic virus (partial CDS) share about 80% amino acid sequence identity, thus warranting an ophiovirus species decamation criterion of <85% CP amino acid sequence identity. The RdRp contains the SDD sequence in motif C, a signature for segmented negative-stranded RNA viruses. Phylogenetic reconstructions using sequences of the conserved core module from RdRps of ophioviruses and representative negative-stranded RNA viruses reinforce their separation as a monophyletic group. Citrus psorosis virus is present worldwide, transmitted by vegetative propagation of the host, and natural dispersion has also been observed. No vector is known for ranunculus white mottle virus and blueberry mosaic associated virus. *Olpidium virulentus* and *Olpidium brassicae* fungi transmit *Mirafiori lettuce big-vein virus*, tulip mild mottle mosaic virus and lettuce ring necrosis virus; freesia sneek virus is presumably transmitted by a member of the genus *Olpidium*. Ranunculus white mottle virus has been reported in Italy, France and Germany [3], and tulip mild mottle mosaic virus in Japan [4]. *Mirafiori lettuce big-vein virus*, the causal agent of big-vein disease in lettuce, probably occurs worldwide. Lettuce ring necrosis virus is closely associated with lettuce ring necrosis disease in The Netherlands, Belgium and France [5]. Freesia sneek virus [6] has been reported in Europe, South Africa, North America, South Korea and New Zealand. Blueberry mosaic associated virus is associated with blueberry mosaic disease found in North and South America, Europe, New Zealand and Japan [7].

**RESOURCE**


**Funding information**

Production of this summary, the online chapter and associated resources was funded by a grant from the Wellcome Trust (WT108418AA).

**Acknowledgements**

We thank Ulises Mancini (Universidad de La Plata) for helping in the phylogenetic analysis. Members of the ICTV Report Consortium are Elliot J. Lefkowitz, Andrew J. Davison, Stuart G. Siddell, Peter Simmonds, Michael J. Adams, Donald B. Smith, Richard J. Orton and Hélène Sanfaçon.

**Conflicts of interest**

The authors declare that there are no conflicts of interest.

**References**


