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ICTV Virus Taxonomy Profile: *Caulimoviridae*

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Abstract

*Caulimoviridae* is a family of non-enveloped reverse-transcribing plant viruses with non-covalently closed circular dsDNA genomes of 7.1–9.8 kbp in the order Ortervirales. They infect a wide range of monocots and dicots. Some viruses cause economically important diseases of tropical and subtropical crops. Transmission occurs through insect vectors (aphids, mealy-bugs, leafhoppers, lace bugs) and grafting. Activation of infectious endogenous viral elements occurs in *Musa balbisiana*, *Petunia hybrida* and *Nicotiana edwardsonii*. However, most endogenous caulimovirids are not infectious. This is a summary of the International Committee on Taxonomy of Viruses (ICTV) Report on the family *Caulimoviridae*, which is available at ictv.global/report/caulimoviridae.

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<th>Table 1. Characteristics of members of the family <em>Caulimoviridae</em></th>
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<td><strong>Typical member:</strong> cauliflower mosaic virus-Cabb-S (V00141), species Cauliflower mosaic virus, genus Caulimovirus</td>
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**VIRION**

Virions are either isometric of 45–52 nm in diameter or, in the case of members of the genera *Badnavirus* and *Tungrovirus*, bacilliform particles of 30 nm × 60–900 nm (Table 1, Fig. 1). Virion sedimentation coefficient (S₂₀,₅₀) is 200–220 S; density in CsCl is 1.37 g cm⁻³. No envelope is present.

**GENOME**

Virions contain a single molecule of non-covalently closed circular dsDNA of 7.1–9.8 kbp [1, 2] with discontinuities at specific sites in the negative-sense (one) and positive-sense strand (one to three). Genomes contain 1–8 ORFs encoding 5–6 conserved protein domains (Fig. 2), depending on the genus.
REPLICATION

Following entry into the cell, the virion is targeted to the nucleus by a nuclear localization signal in the N-terminus of the capsid protein. Discontinuities in the genome are sealed to give supercoiled DNA, which associates with histone proteins to form mini-chromosomes in the nucleus. These are transcribed by host DNA-directed RNA polymerase II to give a greater-than-genome length transcript (35S or 34S RNA) that has a terminal redundancy of 35 to 270 nt. This transcript (pregenomic RNA) serves as a template for reverse transcription to give the negative-sense strand DNA and as a polycistronic mRNA for expression of at least some of the ORFs [3].

Unlike retroviruses, the episomal replication cycle does not involve an integration phase [4–6]. Negative-sense strand DNA synthesis is primed by host cytosolic tRNA\textsuperscript{met}. Synthesis of both strands is performed by the viral reverse transcriptase and RNase H1. RNase H1-resistant polypurine stretches serve as primer for positive-sense DNA synthesis. The site-specific discontinuities are at the priming sites for both negative- and positive-sense strand DNA synthesis and are made by the oncoming strand displacing the existing strand for a short distance and not ligating to form a closed circle [2].

TAXONOMY

Current taxonomy: ictv.global/report/caulimoviridae. Members of the genera Badnavirus and Tungrovirus have bacilliform virions whereas members of the genera Caulimovirus, Cavemovirus, Petuvirus, Rosadnavirus, Solendovirus and Soymovirus have isometric virions. The number of ORFs ranges between one (petuviruses and vacciniviruses), three or more (badnaviruses), four (cavemoviruses, dioscoviruses, solendoviruses and tungroviruses), seven (caulimoviruses), seven or eight (soymoviruses) and eight (rosadnaviruses). Insect-mediated transmission has been reported for badnaviruses, caulimoviruses and tungroviruses. Infectious endogenous viral elements (EVEs) have been reported for several banana streak viruses (Badnavirus), petunia vein clearing virus (Petuvirus) and tobacco vein clearing virus (Solendovirus).

RESOURCES

Current ICTV Report on the family Caulimoviridae: ictv.global/report/caulimoviridae

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Conflicts of interest

The authors declare that there are no conflicts of interest

References