**EPPO Datasheet: *Robigovirus robigomaculae***

Last updated: 2023-08-01

**IDENTITY**

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| **Preferred name:** *Robigovirus robigomaculae* **Taxonomic position:** Viruses and viroids: Riboviria: Orthornavirae: Kitrinoviricota: Alsuviricetes: Tymovirales: Betaflexiviridae: Robigovirus **Other scientific names:** *CRMaV*, *Cherry rusty mottle associated virus*, *Cherry rusty mottle virus* [view more common names online...](https://gd.eppo.int/taxon/CRMAV0/) **EU Categorization:** A1 Quarantine pest (Annex II A) [view more categorizations online...](https://gd.eppo.int/taxon/CRMAV0/categorization) **EPPO Code:** CRMAV0 |  |

**Notes on taxonomy and nomenclature**

Cherry rusty mottle associated virus (CRMaV) is a positive-sense single-stranded RNA (+ssRNA) virus with a monopartite genome of approximately 8.4 kb encapsidated flexuous rod-like virions (Villamor *et al.*, 2013). A reference genomic sequence of CRMaV is available in GenBank ([**NC\_020996.1**](https://www.ncbi.nlm.nih.gov/nuccore/NC_020996.1)). The disease cherry rusty mottle was initially described in Washington State (USA) in 1940 (Reeves, 1940). Later on, mild and severe forms of the disease were described in Oregon (USA) and a similarly named disease described in Europe (Posnette, 1951). It was later determined that despite being similarly named, the disease in the USA in Europe were different (Posnette & Cropley, 1961). The causal agent of the cherry rusty mottle (European) disease has not been identified to date but might be a viral complex. CRMaV was initially described from cherry sources affected by cherry rusty mottle (American) disease and consequently named cherry rusty mottle associated virus. The Koch's postulates have been completed following agroinoculation of an infectious cDNA clone (Villamor *et al.*, 2021) demonstrating that CRMaV is a distinct virus and is the causal agent of the disease in North America (Villamor & Eastwell, 2013; Villamor *et al.*, 2015; 2021). This in turn has led some authors to suggest a renaming of the virus to cherry rusty mottle virus (CRMV), a step that has not yet been agreed upon by the ICTV, so that the virus official name remains to date cherry rusty mottle associated virus.

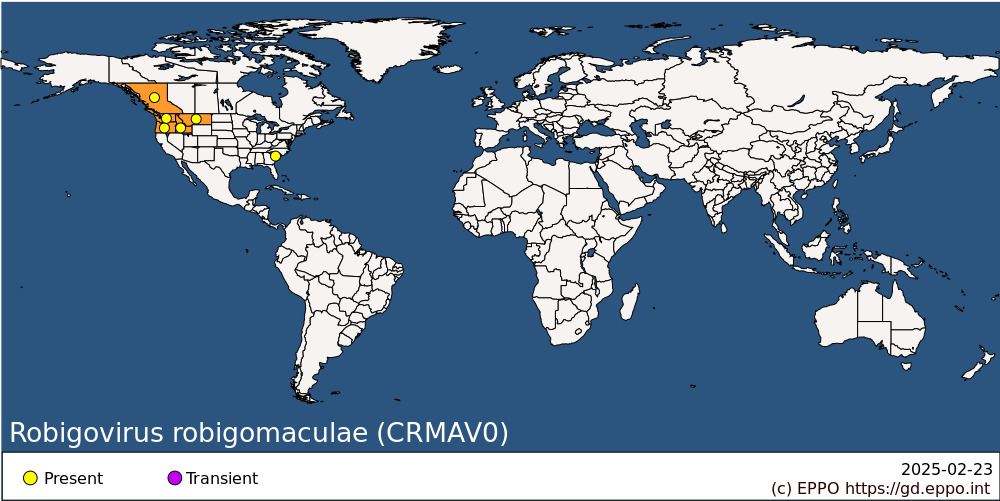
**HOSTS**

The main host of CRMaV is sweet cherry (*Prunus avium*), on which symptoms are most frequently observed. The virus has been anecdotally reported on, or experimentally shown to be able to infect, a few *Prunus* hosts, i.e. *P. lusitanica* (Villamor *et al.*, 2014), *P. mahaleb* (Villamor & Eastwell, 2013), *P. serrulata* and *Prunus x yedoensis* (Poudel & Scott, 2017)*.*

**Host list:** *Prunus avium*, *Prunus lusitanica*, *Prunus mahaleb*, *Prunus serrulata*, *Prunus x yedoensis*

**GEOGRAPHICAL DISTRIBUTION**

Cherry rusty mottle associated virus has so far only been reported from North America, in Canada (British Columbia, Mink, 1995) and in the USA in several west coast states (Mink, 1995; Villamor *et al.*, 2014; Reinhold & Pscheidt, 2023) as well as once on the east coast, in North Carolina (Poudel & Scott, 2017). The virus has to date not been reported from other areas in the world.

 **North America:** Canada (British Columbia), United States of America (Idaho, Montana, Oregon, South Carolina, Washington)

**BIOLOGY**

CRMaV is systemic in its host plants and infects all plant parts. It is transmitted by grafting (Rott & Jelkmann, 2011) and other vegetative propagation techniques. It has no known vector and is not known to be seed- or pollen-transmitted (Rott & Jelkmann; EFSA, 2019). No herbaceous host of CRMaV is known.

**DETECTION AND IDENTIFICATION**

**Symptoms**

Affected *P. avium* trees begin to show symptoms of light green or yellow mottling on small basal leaves a few weeks after blooming (Rott & Jelkmann, 2011; Villamor and Eastwell, 2013). Chlorotic areas then turn bright yellow, brownish or show late-season reddening, hence the name ‘rusty’ mottle given to the disease. Affected leaves fall rapidly leading to partial tree defoliation (Rott & Jelkmann, 2011). Remaining leaves develop chlorotic mottling. In severely affected trees, leaf colouring and early senescence resembling fall colours may occur, together with fruit size reduction, late ripening and quality loss (tasteless fruits). In milder forms leaf bronzing occurs without early leaf fall, and fruits’ size and quality are less affected. Trees also show decline and dieback (Rott & Jelkmann, 2011). Typical cherry rusty mottle symptoms were reproduced in the Mazzard sweet cherry indicator following grafting of the Krymsk6 cherry rootstock [*P. cerasus* x (P*. cerasus x P. maackii*)] following its agroinoculation of a cloned CRMaV infectious cDNA construct (Villamor *et al.*, 2021).

There are indications that all varieties of *P. serrulata* may not be equally susceptible and develop symptoms of CRMaV infection (Poudel & Scott, 2017). However at least some varieties are known to express light green or yellow mottling symptoms, such as the cv. Kwanzan *P. serrulata* indicator (Villamor & Eastwell 2013).

Chlorotic yellow leaf blotch symptoms were observed on Portuguese laurel (*Prunus lusitanica*) infected by CRMaV (Villamor *et al.*, 2014). However, the presence of additional viruses that may have contributed to these symptoms cannot be completely excluded.

**Morphology**

The viral particles of CRMaV have yet to be observed by electron microscopy. However, by analogy with other *Betaflexiviridae* members, they would be expected to be flexuous and elongated particles of 10-15 x 600-1000 nm.

**Detection and inspection methods**

Visual examination may allow the detection of symptoms but is not considered reliable enough since symptoms are not highly specific and are not always obvious in infected plants. A procedure for inspection of places of production of *Prunus* trees is provided in Standard PM 3/76 (EPPO, 2021).

CRMaV can be detected by biological indexing on woody indicators such as *Prunus avium* cv. Bing, Sam or F12/1 Mazzard and *P. serrulata* cv. Kwanzan (Rott & Jelkmann, 2011; Villamor & Eastwell, 2013; Villamor *et al.*, 2021). However, such tests are relatively lenghty and low-throughput and results may be complicated to interpret in the case of mixed infections.

There are no commercially available antisera that could be used for detection of CRMaV using serological tests. The molecular characterization of CRMaV and the sequencing of the genome of several isolates have allowed the development of primer pairs that can be used in RT-PCR tests. The broad-spectrum nested RT-PCR tests developed by Foissac *et al.* (2005) and Villamor *et al.* (2013), or the specific test developed by Villamor & Eastwell (2013) may be used. However, the analytical specificity of these tests requires further evaluation. HTS based approaches can also be used for the detection of CRMaV.

**PATHWAYS FOR MOVEMENT**

Movement and trade of contaminated propagation materials is considered the most significant pathway for movement since CRMaV is readily transmitted by grafting.

**PEST SIGNIFICANCE**

**Economic impact**

Trees of sweet cherry infected with CRMaV show defoliation, decline and dieback. More or less pronounced fruit size reduction and fruit quality loss are also observed. Detailed data on yield losses caused by CRMaV are not available but the intensity of defoliation reported in severe cases (30 to 70% leaf loss by fruit maturity, Wadley & Nyland, 1976) suggest the potential for severe impact on sweet cherry production.

**Control**

The most efficient control strategy is the development and use of CRMaV-free propagation material, as described in EPPO Standard PM 4/29(1) *Certification scheme for cherry* (EPPO, 2001). No control measures are known in the field, besides the destruction of infected plants and the limitation of movement of host plants outside the infected area.

**Phytosanitary risk**

The virus typically infects and has its main impact in sweet cherry (*P. avium*). Sweet cherry is widely grown in the EPPO region and represents an important fruit crop. A few other *Prunus* species are known to be susceptible. There are no known ecoclimatic constraints for CRMaV establishment, except those affecting its hosts; and sweet cherrycultivation occurs widely in Europe (EFSA, 2019). It was therefore considered justified by some EPPO countries (e.g. in the EU) to prevent establishment and spread of CRMaV.

**PHYTOSANITARY MEASURES**

Appropriate phytosanitary measures to import plants for planting (excluding seeds and pollen) of Prunus hosts into the EPPO region could require that these plants are produced in a pest free area, in a pest free place/site of production, or shown to be free from CRMaV by appropriate diagnostic methods. A number of EPPO countries already ban the import of Prunus (other than fruits and seeds) from areas where the pest is present (EU, 2019).

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**How to cite this datasheet?**

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**Datasheet history**

This datasheet was first published online in 2023. It is maintained in an electronic format in the EPPO Global Database. The sections on 'Identity', ‘Hosts’, and 'Geographical distribution' are automatically updated from the database. For other sections, the date of last revision is indicated on the right.

