**EPPO Datasheet: *Caulimovirus venafragariae***

Last updated: 2023-03-08

**IDENTITY**

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| **Preferred name:** *Caulimovirus venafragariae* **Taxonomic position:** Viruses and viroids: Riboviria: Pararnavirae: Artverviricota: Revtraviricetes: Ortervirales: Caulimoviridae: Caulimovirus **Other scientific names:** *SVBV*, *Strawberry vein banding caulimovirus*, *Strawberry vein banding virus*, *Strawberry virus 5* **Common names in English:** leaf curl of strawberry, vein banding of strawberry [view more common names online...](https://gd.eppo.int/taxon/SVBV00/) **EPPO Categorization:** A2 list **EU Categorization:** RNQP (Annex IV) [view more categorizations online...](https://gd.eppo.int/taxon/SVBV00/categorization) **EPPO Code:** SVBV00 | 14826.jpg [more photos...](https://gd.eppo.int/taxon/SVBV00/photos) |

**Notes on taxonomy and nomenclature**

Strains of *Strawberry vein banding virus* (SVBV) that have been identified include: strawberry yellow veinbanding virus, strawberry necrosis virus (Schöninger), strawberry chiloensis veinbanding virus, strawberry eastern veinbanding virus. In North America, most strains found on the west coast are more severe than those found along the east coast.

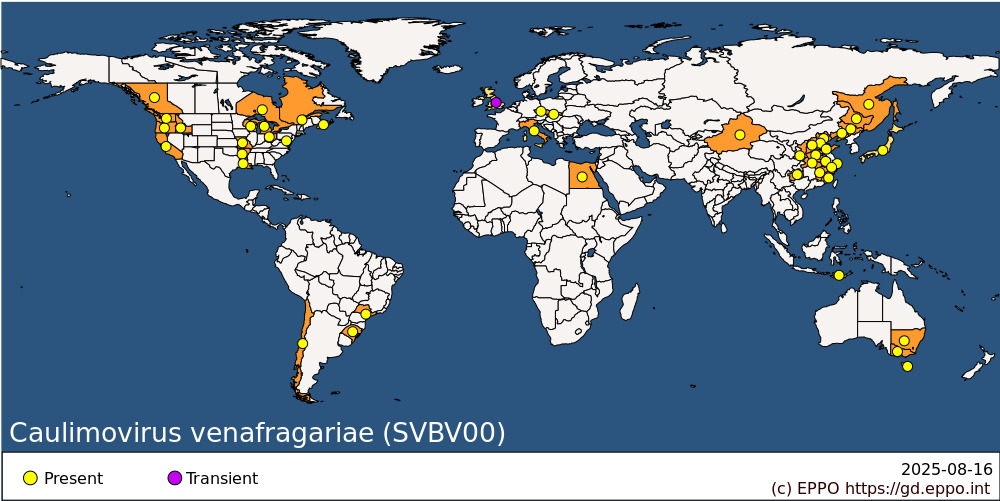
**HOSTS**

The virus is known to occur only on *Fragaria* spp. The main host is *Fragaria vesca* (wild strawberry). Commercial strawberries may also be infected.

**Host list:** *Fragaria chiloensis subsp. chiloensis*, *Fragaria vesca*, *Fragaria virginiana*, *Fragaria x ananassa*

**GEOGRAPHICAL DISTRIBUTION**

The distribution of SVBV shown on the map below is mainly based on references that used nucleic acid-based detection techniques (PCR and western hybridization), but it still contains several records of SVBV made before 1995 that were based solely on disease symptoms. These records have to be considered critically and require further confirmation

 **EPPO Region:** Czechia, Italy (mainland), Russian Federation (the) (Far East), Slovakia, United Kingdom **Africa:** Egypt **Asia:** China (Anhui, Beijing, Fujian, Guizhou, Hebei, Heilongjiang, Henan, Hubei, Jiangxi, Jilin, Liaoning, Shaanxi, Shandong, Shanghai, Shanxi, Xinjiang, Zhejiang), Indonesia (Nusa Tenggara), Japan **North America:** Canada (British Columbia, Nova Scotia, Ontario, Québec), United States of America (Arkansas, California, Idaho, Louisiana, Maryland, Michigan, Missouri, Ohio, Oregon, Washington, Wisconsin) **South America:** Brazil (Rio Grande do Sul, Sao Paulo), Chile **Oceania:** Australia (New South Wales, Tasmania, Victoria)

**BIOLOGY**

In the field, SVBV is transmitted by aphids in a semipersistent manner, and the following species are cited as vectors: *Acyrthosiphon malvae malvae (*syn*. Macrosiphum pelargonii)*, *Amphorophora rubi*, *Amphorophora agathonica, A. rubifolii*, *Aulacorthum solani*, *Chaetosiphon fragaefolii*, *C. jacobi*, *C. tetrarhodum*, *C. thomasi*, *Macrosiphum rosae*,*Myzus ascalonicus*, *M. ornatus*, *M. persicae*.

Of these species, *Chaetosiphon* spp. are the most efficient vectors in glasshouse experiments, although other genera are probably important vectors when they occur in large numbers and frequently move from plant to plant. Aphids can acquire and transmit the virus in 30-120 min, but persistence in the vector is short, usually less than 8 h (semipersistent type). There are differences in the efficiency of clonal lines of aphids, and evidence that some species will only transmit certain strains of SVBV. *Aphis gossypii*, *A. fabae*, *Aulacorthum solani* and *Macrosiphum euphorbiae* failed to transmit the virus in a limited number of trials.

The virus is transmissible by grafting and by means of *Cuscuta subinclusa*. Attempts to transmit SVBV mechanically have been unsuccessful. The incubation period in the indicator host varies from 2 to 5 weeks depending on the strain.

For additional information, see Frazier (1955), Miller & Frazier (1970), Smith (1972).

**DETECTION AND IDENTIFICATION**

**Symptoms**

***On Fragaria vesca***

Symptoms initially appear on the youngest developing leaf; there is epinasty of the midrib and petiole, a tendency for opposite halves of leaflets to be appressed, irregularly wavy leaflet margins, and slight crinkling of the laminae. Usually, the above symptoms are mild and not all present simultaneously. It is not until the affected leaf expands that clearing, followed by yellowish banding of some or all of the veins, becomes visible. Often, the coloration occurs in scattered discontinuous streaks of varying lengths along the main and secondary veins.

The second and third leaves formed after onset of symptoms are affected more severely than the first or any subsequent leaf; in older leaves, chlorotic streaks are reduced in number, scattered and confined to portions of the leaflets. This may be followed by the appearance of a series of apparently healthy leaves and then reappearance of mild or severe symptoms.

For additional information, see Frazier (1955), Mellor & Fitzpatrick (1961), Miller & Frazier (1970), Smith (1972) and Frasier & Converse (1980).

***On commercial strawberries***

SVBV usually does not induce distinct symptoms in commercial cultivars, and often the only indications of infection are loss of vigour, stunting, lowered yields of a cultivar. In the cv. Marshall, for example, the veinbanding is usually diffuse, commonly located along main veins and may often appear as spots. As affected leaves mature, the veinbanded areas may gradually disappear, or they may become brownish-red or necrotic. In particular on outdoor plants, the veins become discoloured, without previous chlorosis. Affected leaflets characteristically exhibit epinasty, mild crinkling and wavy margins. SVBV rarely occurs singly in strawberry; frequently several viruses are present, and together they cause more severe reductions in productivity (Spiegel and Martin, 1998). Symptoms may be more severe in combination with the strawberry crinkle virus (Martin & Tzanetakis, 2006).

**Morphology**

The particle of this caulimovirus is isometric (40-50 nm in diameter). Native viral DNA is circular and double-stranded (Stengel *et al*., 1988).

**Detection and inspection methods**

Diagnosis can be made or confirmed by use of virus-free *F. vesca* indicator plants. The *F. vesca* clone UC-6, the *F. virginiana* clone UC-12 and *F. vesca semperflorens* are recommended for detecting and diagnosing SVBV (Converse, 1987). A modified leaf grafting technique is used (Frazier, 1974).

An ELISA test can be performed using Cauliflower mosaic virus antisera which cross-reacts with SVBV (Honetslegrova *et al*., 1995). However, no SVBV specific antiserum or commercial ELISA kit is available because of low yield of SVBV virus particles embedded in inclusion bodies and their low immunogenicity.  Cloning and prokaryotic expression of SVBV genes seems to be a promising state-of-the-art method for the production of specific antisera in future (Jiang *et al*., 2020)

SVBV variability has been studied extensively. Several full length (Petrzik *et al.,* 1998) and dozens of partial sequences have been published in databases and molecular biology-based methods give the best detection results. The gene coding for the coat protein of the virus is highly conserved, and SVBV can be detected readily by PCR using primers in the coat protein open reading frame. Nucleic acid isolation from fresh plant tissue was the best template for PCR, and primers amplifying the shortest product should be recommended. Vaskova *et al.* (2004) developed a test for the detection of SVBV in *Fragaria*spp*.* based on nucleic acid sequence-based amplification (NASBA) and real-time detection using molecular beacons (real-time NASBA). Several multiplex RT-PCR (Thompson *et al*., 2003; Zhang *et al*., 2009) as well as quantitative PCR (q-PCR) (Diaz-Lara et al., 2021) and loop-mediated isothermal amplification (LAMP) (Ren *et al*., 2022) tests were developed for routine detection of SVBV.

**PATHWAYS FOR MOVEMENT**

In the field, the virus is transmitted by aphid vectors. Because of the ability of certain aphid species to undertake long, high-altitude flights, wide natural dissemination is possible. This is, however, limited by the relatively short persistence of the virus in the vector.

In international trade, SVBV is liable to be carried on infected plants and propagating material of strawberries.

**PEST SIGNIFICANCE**

**Economic impact**

Because of the sporadic occurrence and low incidence of SVBV, the disease is only of minor importance but under extreme aphid pressure, the incidence can approach 100% in third-year fields (Martin & Tzanetakis, 2006). Fruit yield and size are affected, and runner production reduced. In combination with strawberry latent C disease, SVBV reduced yield by 17% in the first fruiting year, and total and saleable fruit by 88% and 100%, respectively, in the third year (Bolton, 1974; EPPO/CABI 1996)*.*

**Control**

There are no specific control measures (Martin & Tzanetakis, 2013). Nevertheless, it is important, even in annual production systems, to control the aphid vectors (primarily *Chaetosiphon fragaefolii*) in order to reduce virus infections (Martin & Tzanetakis, 2006). SVBV is highly resistant to inactivation by heat therapy but it can be eliminated from plants by means of meristem tip culture. As a consequence, the use of certified planting material is the best control procedure, and certification schemes for the production of healthy planting material of strawberry are in operation in several EPPO countries. An EPPO certification scheme for strawberries is available and provides guidelines on how to produce healthy planting material (EPPO, 2008). Control of aphids with insecticides could reduce the incidence of the disease.

**Phytosanitary risk**

The most important factors in evaluating the potential impact of SVBV in a new area are the presence of aphid vectors and their mobility. Because of the variety of vectors, conditions can be defined only in so far as they affect aphids in general, e.g. extremely low winter temperatures killing overwintering nymphs and adults; windy climates restricting activity of alatae. In its evaluation for the European Union, EFSA (2014) considered that SVBV presented a minor risk to strawberry production under the current cultivation practices, in particular with the use of certified planting material and short crop cycles which have reduced the impact of strawberry viruses. EFSA concluded that SVBV met the criteria of a regulated non-quarantine pest (RNQP).

**PHYTOSANITARY MEASURES**

Importing countries may require that plants for planting of *Fragaria ananassa*, from countries where the pest occurs, should be derived from mother plants tested and found free from SVBV during the last three growing seasons and should have been maintained under conditions preventing their reinfection; the consignment must come from a field found free (along with its immediate vicinity) of the virus during the last growing season.

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**CABI resources used when preparing this datasheet**

CABI Compendium on Strawberry vein banding virus (vein banding of strawberry). (2021) [https://doi.org/10.1079/cabicompendium.52407](http://%20https%3A/doi.org/10.1079/cabicompendium.52407)

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**How to cite this datasheet?**

EPPO (2025) *Caulimovirus venafragariae*. EPPO datasheets on pests recommended for regulation. Available online. <https://gd.eppo.int>

**Datasheet history**

This datasheet was first published in the EPPO Bulletin in 1978 and revised in the two editions of 'Quarantine Pests for Europe' in 1992 and 1997, as well as in 2023. It is now 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.

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