Virus Assessment of Ajuga reptans Cultivars Reveals Alfalfa Mosaic, Tobacco Streak, and Cucumber Mosaic (CMV) Viruses, and a CMV Satellite RNA

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Additional index words. reverse transcription polymerase chain reaction (RT-PCR)

Abstract. Ajuga reptans L. is an herbaceous ornamental mint grown in borders or as a groundcover, and is commonly propagated vegetatively and by seed. Three hundred and fifty-six A. reptans samples were obtained from growers in Washington, Michigan, Iowa, and Ohio, and screened for alfalfa mosaic virus (AMV), tobacco streak ilarivirus (TSV), cucumber mosaic cucumovirus (CMV), tomato aspermy cucumovirus (TAV), tomato spotted wilt tospovirus (TSWV), impatiens necrotic spot tospovirus (INSV), tobacco mosaic tobamovirus (TMV), potato virus X potexvirus (PVX), and 80 potyviruses, using direct antibody sandwich (DAS) and indirect enzyme-linked immunosorbent assay (ELISA). Viral-associated double-stranded ribonucleic acid (dsRNA) analysis was used to detect an apparent satellite (sat) RNA, and northern hybridization using a digoxigenin (DIG) labeled (S) CARNA-5 cDNA probe was used to confirm the identity of the apparent satRNA. No incidences of TAV, TMV, TSWV, INSV, PVX, or potyviruses were detected. CMV was detected in 11%, AMV in 22.2%, TSV in 3.7%, and mixed infections of CMV and AMV in 11.1% of the samples. SatRNA was detected in 36 A. reptans ‘Royalty’, two ‘Rainbow’, and two ‘Burgundy Glow’ samples by dsRNA analysis, and confirmed by hybridization in 29 ‘Royalty’ and one ‘Burgundy Glow’ samples. Sixteen A. reptans ‘Royalty’ seedlings grown from seed harvested from CMV-infected plants were tested by ELISA for CMV, AMV, and TSV. All were positive for CMV, and two were positive for a mixed infection of CMV and AMV. SatRNA was detected in all 16 seedlings by RT-PCR.

Ajuga reptans L. is a perennial ornamental mint (Lamiaceae) grown in borders or as a groundcover. Cultivars include ‘Bronze Beauty’, ‘Burgundy Glow’, ‘Royalty’, ‘Rainbow’, ‘Silver Beauty’, and ‘Catlin’s Giant’, and are commonly propagated vegetatively and by seed. Cucumber mosaic cucumovirus (CMV), alfalfa mosaic virus (AMV), tobacco streak ilarivirus (TSV) and broad bean wilt fabavirus (BB WV) were reported in this host in Australia (Shukla and Gough, 1983), CMV in Denmark (Kristensen, 1956), and AMV in the United States (Schroeder and Provvidenti, 1972). Recently CMV, AMV, and TSV were reported in A. reptans cultivars in Ohio, and a satRNA was also found to be associated with CMV-infected A. reptans ‘Royalty’ plants (Fisher and Nameth, 1997).

CMV, TSV, and AMV are single-stranded RNA viruses belonging to the family Bromoviridae. All three contain three major genome segments, and have very wide host ranges (Francki et al., 1979; Fulton, 1985; Jaspers and Bos, 1980). The AMV is transmitted mechanically, through seed, and by 24 species of aphids in the nonpersistent manner (Edwardson and Christie, 1991; Jaspers and Bos, 1980). The TSV is transmitted mechanically, through seed, and by at least two thrips species [Frankliniella occidentalis (Pergande) and Thrips tabaci Lindeman] (Fulton, 1985), and CMV is transmitted mechanically, through seed, and by >60 aphid species in the nonpersistent manner (Francki et al., 1979). Many CMV isolates also have an additional satellite RNA (satRNA) associated with them. This satRNA is dependent upon the helper virus for replication and encapsidation, often interferes with the replication of the helper virus, and affects symptoms induced by the helper virus (Collmer and Howell, 1992). The effects on symptom expression produced by the presence of satRNA are influenced by the strain of CMV, the host plant, and the satRNA strain. For example, symptoms on tomato (Lycopersicon esculentum Mill.) plants containing satRNA can range from lethal necrosis, to white chlorosis, to disease attenuation, depending on the satRNA strain (Collmer and Howell, 1992). However, the same satRNA strain can have different effects on tomato and tobacco (Nicotiana sp.). The CMV white leaf (WL) strain plus a particular satRNA produces a white leaf syndrome on tomato. On tobacco, the CMV-WL genomic RNAs produce a brilliant chlorosis, but the addition of the satRNA causing white leaf in tomato leads to a symptomless infection (Gonsalves et al., 1982). The CMV satRNAs can also be helped by the related tomato aspermy cucumovirus (TAV) (Collmer and Howell, 1992).

In this paper we report the results of a serological screening of A. reptans cultivars for CMV, AMV, TSV, TAV, tomato spotted wilt tospovirus (TSWV), impatiens necrotic spot tospovirus (INSV), tobacco mosaic tobamovirus (TMV), potato virus X potexvirus (PVX), and a potyvirus screen. Ajuga reptans samples were obtained from growers in the United States and screened for viral infection using the enzyme-linked immunosorbent assay (ELISA).

Materials and Methods

Plant material. Samples were obtained from growers in the states of Washington, Michigan, Iowa, and Ohio. Growers were requested to send plants typical of their stock, but any plants displaying symptoms were of special interest. Additionally, samples were collected from several established plantings in Ohio. A total of 356 samples were tested for all nine viruses. Live plants were transplanted and maintained in the greenhouse for the project duration and symptoms were noted prior to ELISA testing. Established plantings were surveyed for virus-like symptoms such as mosaic and ringspots, and samples of the symptomatic plants were collected. If no symptoms were evident then asymptomatic tissue was collected. Freshly collected tissue was prepared immediately for direct antibody sandwich (DAS) ELISA.

DAS and indirect ELISA. Samples were tested using a variation of the method previously described (Clark and Adams, 1977). Commercially produced antibodies (AGDIA, Elkhart, Ind.) against CMV, AMV, TSV, TAV, TSWV, INSV, TMV, and PVX were used in this study, as well as an indirect ELISA potyvirus screen that tests for 80 potyviruses. Asymptomatic, apparently virus-free, A. reptans ‘Bronze Beauty’ tissue collected from an established planting was used as negative control after first testing against all of the antisera using extraction buffer as the negative control.

Received for publication 15 Mar. 1999. Accepted for publication 31 Aug. 1999. This paper is a portion of a thesis submitted by J.R. Fisher toward the completion of the requirements of the Ph.D. degree in Plant Pathology. Salaries and research support provided by state and federal funds appropriated to the Ohio Agricultural Research and Development Center, Ohio State Univ. The cost of publishing this paper was defrayed in part by the payment of page charges. Under postal regulations, this paper therefore must be hereby marked advertisement solely to indicate this fact.

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tate false positives were not obtained due to cross-reactivity. Nune Maxisorp 96 well polystyrene microtitre plates (Thomas Scientific, Swedesboro, N.J.) were used in the study. Each sample was dispensed into two test wells, and the average absorbance of the two wells at 405 nm was determined using a microtitre plate reader (EAR 400SF Plus ELISA plate reader; SLT Lab Instruments, Hillsborough, N.C.). A sample was considered to be positive if its absorbance was twice that of the negative control value established for that plate.

**Viral dsRNA analysis.** Viral-associated double-stranded ribonucleic acid (dsRNA) was extracted and purified as previously described (Jordan et al., 1983; Morris and Dodds, 1979). DSRNA was loaded onto 5% or 10% polyacrylamide gels and analyzed by electrophoresis. Ethidium bromide-stained dsRNA bands were visualized on a transilluminator (302 nm) and photographed with Polaroid-type 667 black and white film.

**Northern hybridization.** Following extraction and purification, dsRNA was heat denatured, subjected to electrophoresis, and visualized on a transilluminator (302 nm) and photographed with Polaroid-type 667 black and white film.

**Reverse transcription polymerase chain reaction (RT-PCR).** Two pairs of primers were designed for RT-PCR detection of satRNA in A. reptans by comparing 10 previously published satRNA sequences. The published satRNA sequences used were G-satRNA, B1-satRNA, B2-satRNA, B3-satRNA, WL1-satRNA, WL2-satRNA (Garcia-Arenal et al., 1987), E-satRNA, OY-satRNA (Hidaka et al., 1988), Y-satRNA (Hidaka et al., 1984) and D-satRNA (Kurath and Palukaitis, 1987). The primer sequences S′-GTTTTGTTTTGAACTGCT-3′, which hybridizes to the 3′ end of minus-sense satRNA, and G′-GGAGATGTCCTGGG-3′, which hybridizes to the 3′ end of positive-sense satRNA, were used to reverse transcribe full length cDNAs. The primer sequences S′-GTMTGATGAGATCGCTAG-3′ and G′-CAATGATGAGGTAGCT-3′, which were nested five bases inward from the S′ and 3′ ends, respectively, of the full-length satRNA molecule, were used in the PCR amplification. Samples were prepared by grinding tissue at a 1:4 ratio in 1× TBST buffer (0.025 M tris, pH 8.0; 0.015 M NaCl; 0.05% w/v Tween-20). The macerate was placed in 1.5-mL microcentrifuge tubes, and centrifuged at 11,500 g, in a benchtop microcentrifuge for 10 min, and the supernatant further diluted to 1:16 and 1:32 in TBST buffer. A two-step RT-PCR reaction was performed as previously described (Innis et al., 1990). The RT reaction was performed in a 20-µL volume with 1 µL each of 0.1 µg µL⁻¹ 3′ and 5′ primer, 3 µL 1:16 or 1:32 exp, and 200 units Superscript II (Gibco, BRL Inc., Gaithersburg, Md.) reverse transcriptase and incubated at 42 °C for 1 h. One-tenth volume (2 µL) of the RT reaction was used in the PCR reaction, which was done in a 50-µL volume with 4 mM MgCl₂, 0.1 µg each 3′ and 5′ primer, and 2.5 units Tag polymerase (Gibco, BRL Inc.). Thirty cycles were performed (94 °C denaturation; 52 °C primer annealing; 72 °C extension), and one-half volume of the reaction was analyzed on a 1.5% agarose gel. Tissue known to be positive for CMV and satRNA was used as positive control in the RT reaction and ddH₂O or TBST buffer used as negative control.

**Inoculation of test plants.** Nicotiana rustica L. and N. tabacum ‘Glurk,’ ‘Turk,’ and ‘Samsun’ were rub-inoculated using one of the symptomatic ‘Royalty’ and ‘Rainbow’ plants as the source of inoculum. Tissue was ground to a fine powder, and a 1:16 and 1:32 in TBST buffer. A two-step RT-PCR reaction was performed as previously described (Jordan et al., 1983; Morris and Dodds, 1979). The macerate was placed in 1.5-mL microcentrifuge tubes, and centrifuged at 10,000 × g for 20 min, and the supernatant further diluted to 1:16 and 1:32 in TBST buffer. A two-step RT-PCR reaction was performed as previously described (Innis et al., 1990). The RT reaction was performed in a 20-µL volume with 1 µL each of 0.1 µg µL⁻¹ 3′ and 5′ primer, 3 µL 1:16 or 1:32 exp, and 200 units Superscript II (Gibco, BRL Inc., Gaithersburg, Md.) reverse transcriptase and incubated at 42 °C for 1 h. One-tenth volume (2 µL) of the RT reaction was used in the PCR reaction, which was done in a 50-µL volume with 4 mM MgCl₂, 0.1 µg each 3′ and 5′ primer, and 2.5 units Tag polymerase (Gibco, BRL Inc.). Thirty cycles were performed (94 °C denaturation; 52 °C primer annealing; 72 °C extension), and one-half volume of the reaction was analyzed on a 1.5% agarose gel. Tissue known to be positive for CMV and satRNA was used as positive control in the RT reaction and ddH₂O or TBST buffer used as negative control.

**Results**

Symptoms observed on A. reptans plants were similar to those previously described (Shukla and Gough, 1983). Red and yellow spots, ringspots, mosaic, and oakenleaf patterns developed on ‘Bronze Beauty’ (Fig. 1A). Symptoms on ‘Burgundy Glow’, which normally has pink, green, and white variegated leaves, included yellow streaks, mosaic and oakenleaf-like patterns (Fig. 1B). Several samples of the cultivar ‘Rainbow’ showed severe mosaic and ringspots (Fig. 1D). Some ‘Royalty’ plants displayed red, orange, or yellow spots, mosaic and ringspots (Fig. 1D). Some ‘Royalty’ plants displayed red, orange, or yellow spots, mosaic and ringspots (Fig. 1D). Some ‘Royalty’ plants displayed red, orange, or yellow spots, mosaic and ringspots (Fig. 1D). None of the 356 A. reptans plants were positive for TAV, TMV, TSWV, INSV, PVX, or the potyvirus screen. Thirty-nine samples were positive for CMV (11%), 13 for TSV (3.7%), 79 for AMV (22.2%), and four (1.1%) for a mixed infection of CMV and AMV, using the criterion described above for determining positive results. Several samples had elevated absorbance values that were not high enough to be considered positive, but were clearly higher than the established negative control values. Thirty-eight samples (10.7%) had elevated absorbance values for AMV, one (2.7%) for CMV, and 18 (5.0%) for TSV. No incidences of a mixed infection of AMV and TSV, or of CMV and TSV, were detected. A summary of the overall ELISA data for CMV, AMV, and TSV is presented in Table 1. In some cases, positive results were obtained from asymptomatic plants, or samples collected from plants showing mosaic or ringspot symptoms were negative in all tests. These results are also summarized in Table 1.

Each of 16 seedlings grown from seed collected from CMV-infected ‘Royalty’ plants was positive for CMV by ELISA. Additionally, two of 16 (11.8%) were positive for a mixed infection of CMV and AMV, but none of the seedlings was positive for TSV. Only two of the 16 seedlings showed a slight mosaicism of symptoms at the time of testing. One of the symptomatic seedlings was positive for CMV, and the other for CMV and AMV. All of the remains remained symptomless over the next 60 d.

**dsRNA analysis.** dsRNA analysis of symptomatic and asymptomatic ‘Royalty’ tissue, and of symptomatic ‘Rainbow’ tissue, produced a banding pattern consistent with infection by CMV and an associated satRNA (Dodds et al., 1984; Valverde et al., 1990) (Fig. 2). The apparent satRNA was present in all 36 ‘Royalty’ samples tested and both ‘Rainbow’ samples tested. In addition, double stranded satRNA was detected in two of the three ‘Burgundy Glow’ samples with a mixed infection of CMV and AMV. The identity of the apparent satRNA was confirmed by hybridization to the (S)CARN-5 probe in 29 of the ‘Royalty’ samples (Fig. 3). One of the ‘Burgundy Glow’ samples showing double stranded satRNA was probed and it reacted faintly (data not shown). The remaining seven ‘Royalty’ and two ‘Rainbow’ samples were not tested by hybridization. All 16 ‘Royalty’ seedlings that were ELISA-positive for CMV were assayed for the presence of satRNA using RT-PCR. A PCR product corresponding to the satRNA was detectable in all (Fig. 4). The identity of the PCR products was confirmed by hybridization to the DIG-labeled (S)CARN-5 probe (not shown).

**Nicotiana rustica,** mechanically inoculated with ‘Royalty’ and ‘Rainbow’ sap, never developed symptoms. N. tabacum ‘Glurk,’ ‘Turk,’ and ‘Samsun’ inoculated with ‘Royalty’ sap all developed similar symptoms ranging from mild mosaic to ringspots to oakenleaf to a systemic line pattern. The line pattern and ringspots were more prevalent on older leaves and often disappeared, while the mosaic and oakenleaf were more prevalent on younger leaves. Nicotiana tabacum ‘Glurk,’ ‘Turk,’ and ‘Samsun’ inoculated with ‘Rainbow’ sap developed symptoms similar to those observed on the plants inoculated with ‘Royalty’ sap. When tested by ELISA for CMV, AMV, and TSV, several symptomatic ‘Turk’ plants inoculated with ‘Royalty’ sap and several ‘Samsun’ inoculated with ‘Rainbow’ sap were positive for CMV. However, the majority of the tobacco plants showing symptoms were negative for all three viruses. dsRNA analysis of ‘Glurk’ inoculated with ‘Rainbow’ sap, and showing line pattern symptoms, produced several faint bands. dsRNA analysis of ‘Samsun’ inoculated with ‘Rainbow’ sap, also showing line pattern, produced a CMV-like banding pattern.
Fig. 1. Virus-induced symptoms observed on *Ajuga reptans* cultivars. (A) ‘Bronze Beauty’, (B) ‘Burgundy Glow’, (C) ‘Royalty’, and (D) ‘Rainbow’.

Table 1. Summary of symptomatic and asymptomatic *Ajuga reptans* cultivars tested for CMV, AMV, and TSV, by ELISA.

| Cultivar* | No. tested | Total | CMV+ | AMV+ | CMV and AMV+ | TSV+ | CMV+ | AMV+ | TSV+ | Total* |
|-----------|------------|-------|------|------|--------------|------|------|------|------|--------|
| BB        | 189        | 88(47)| 0    | 22(12)| 0            | 10(5)| 0    | 0    | 0    | 11(6)  |
| BG        | 27         | 5(19)| 0    | 22(82)| 3(11)        | 0    | 0    | 0    | 0    | 20(74) |
| ROY       | 36         | 6(17)| 35(97)| 0    | 0            | 0    | 0    | 0    | 0    | 0      |
| RB        | 2          | 2(100)| 0    | 0    | 0            | 0    | 0    | 0    | 0    | 0      |
| GR        | 35         | 2    | 0    | 22(63)| 3(9)         | 0    | 0    | 0    | 0    | 20(57) |
| GTY       | 24         | 0    | 1(4) | 5(21)| 0            | 0    | 1(4)| 5(21)| 0    | 0      |
| SQ        | 24         | 0    | 1(4) | 7(29)| 1(4)         | 0    | 0    | 6(25)| 0    | 0      |
| SB        | 6          | 0    | 0    | 0    | 0            | 0    | 0    | 0    | 0    | 0      |
| CG        | 7          | 4(57)| 0    | 1(14)| 0            | 0    | 0    | 0    | 0    | 4(57)  |
| MCR       | 4          | 0    | 0    | 0    | 0            | 0    | 0    | 0    | 0    | 0      |
| AG        | 2          | 0    | 0    | 0    | 0            | 0    | 0    | 0    | 0    | 0      |
| Total     | 356        | 107  | 39   | 79   | 4            | 13   | 30   | 62   | 11   | 36     |

% of Total 30 11 22 1.1 4 8 17 3 10

*Figures in parentheses = percentages of samples of that cultivar tested.

*Cultivar abbreviations: BB = Bronze Beauty; RB = Rainbow; SQ = Silver Queen; AG = Arboretum Giant; BG = Burgundy Glow; GTY = Gaiety; CG = Catlin’s Giant; GR = Green; ROY = Royalty; SB = Silver Beauty; MCR = Mini Crispa Red.

*Mosaic; red, yellow spots; ringspots; oakleaf pattern.

*Total of samples with symptoms that tested negative for all viruses.
Fig. 2. Double-stranded ribonucleic acid analysis (dsRNA) banding patterns obtained from Lane 1: cucumber mosaic virus white-leaf isolate (CMV-WL); Lane 2: Ajuga reptans ‘Royalty’; and Lane 3: tobacco mosaic virus (TMV) common strain. Numbers next to lane 1 indicate approximate dsRNA molecular weights (2.0, 1.5 × 10^6). Arrow indicates double-stranded satRNA band.

Fig. 3. Northern blot analysis of Ajuga reptans ‘Royalty’ samples using digoxigenin-labeled (S)CARNAA-5 cDNA; Lane 1: tobacco mosaic virus (TMV) common strain (negative control); Lane 2: unlabeled (S)CARNAA-5 cDNA (positive control). Lanes 3–8: A. reptans ‘Royalty’ samples. Arrow indicates satRNA band.

Fig. 4. Reverse transcription polymerase chain reaction (RT-PCR) analysis of Ajuga reptans ‘Royalty’ seedling samples using primers specific for cucumber mosaic virus (CMV) satellite RNAs. Lanes 1–2: A. reptans ‘Royalty’ seedling samples; Lane 3: CMV isolate from A. reptans ‘Royalty’ in N. tabacum ‘Glurk’ (positive control); Lane 4: double-distilled water (negative control); Lane 5: (S)CARNAA-5 cDNA marker (faint band between 0.3 and 0.4 Kb markers) digested from pSP65 plasmid (3.0 Kb band) with EcoRI; and Lane 6: 1 Kb Plus DNA markers. Arrow indicates PCR product between 0.3 and 0.4 Kb marker.

Fig. 5. ‘Turk’ inoculated with ‘Royalty’ sap and showing systemic line pattern/ringspots was used as inoculum to inoculate a series of ‘Glurk’ plants. The systemic line pattern and ringspot symptom developed on the ‘Glurk’ plants, and the symptomatic tissue was tested by ELISA for CMV, AMV, tobacco ringspot Nepovirus (TbRSV) and tomato ringspot Nepovirus (TRSV). All of the inoculated ‘Glurk’ samples were positive for AMV and negative for the other viruses.

Discussion

These ELISA results show that only CMV, AMV, and TSV were present in the cultivars of A. reptans tested. CMV was found in 11.0%, AMV in 22.2%, TSV in 3.7%, and a mixture of CMV and AMV in 1.1% of the samples tested. When the data were broken down by cultivar, CMV was more prevalent in ‘Royalty’ and ‘Rainbow’; AMV in ‘Bronze Beauty’, ‘Burgundy Glow’, and ‘Green’; and TSV in ‘Bronze Beauty’ and ‘Green’ (Table 1).

Thirty-five (18.5%) ‘Bronze Beauty’, one (2.9%) ‘Green’, one (4.2%) ‘Gaiety’, and one (4.2%) ‘Silver Queen’ sample had elevated absorbance values for AMV, but these were not high enough to be considered true positives. The AMV often causes a symptomless infection in many of its hosts (Jaspars and Bos, 1980), and in tobacco AMV quickly reaches peak titre, but then declines to a very low level, which coincides with recovery from symp-
toms (Jaspers and Bos, 1980; Ross, 1940). That this occurs in *Ajuga reptans* as well is very likely, and the samples with elevated absorbance for AMV may represent cases where the virus titre was too low to detect or that there was an infection by another, still unidentified, virus.

The situation may be very different in ‘Royalty’, where 30 of 36 plants were asymptomatic, but 35 of 36 (97.2%) were positive for CMV, and one had a negative, but elevated, absorbance value. As indicated, a satRNA was associated with CMV in ‘Royalty’ and ‘Rainbow’ samples. An explanation for the high number of symptomless CMV infections in ‘Royalty’ could be a symptom-modulating effect of the satRNA. That there are two main satRNA phenotypes is generally accepted. Symptoms induced by the helper virus are either attenuated or exacerbated by the satRNA (Collmer and Howell, 1992). The satRNA is probably attenuating symptoms in the asymptomatic ‘Royalty’ plants.

None of the samples tested were positive for TAV, which can also act as the helper virus for CMV satRNAs (Collmer and Howell, 1992). In the case of the two ‘Burgundy Glow’ plants with a mixed infection by CMV and AMV, and where dsRNA was detectable, this is the first time we have encountered satRNA in this cultivar. Previously we could detect CMV genomic dsRNAs, but never dssatRNA (Fisher and Nameth, unpublished). The results of the mechanical inoculations of tobacco using ‘Royalty’ and ‘Rainbow’ as inoculum sources indicate that AMV may also be prevalent in those cultivars.

The ELISA results obtained from the ‘Royalty’ seedlings grown from the seed of CMV-infected ‘Royalty’ plants suggest that CMV and AMV may be seed transmitted in this host as well. A more detailed study is required to determine if the seed is just surface-contaminated or if the embryo is infected. The mixed infection by CMV and AMV in two of the 16 ‘Royalty’ seedlings tested is the first instance we have observed of these two viruses being detected together directly from ‘Royalty’ tissue. A possible explanation could be that the young tissue collected was actively growing, and this facilitated the replication of the AMV. The RT-PCR results confirmed that the satRNA was also present in all 16 CMV-infected ‘Royalty’ seedlings. Because there was very little tissue to work with, given the small size of the seedlings, RT-PCR was an excellent tool for detection of satRNA.

These results have several implications for the commercial perennial plant grower. First, since *Ajuga reptans* is an herbaceous perennial, it can serve as a reservoir for CMV and its satRNA, AMV, and TSV. This becomes even more important when one takes into consideration how many asymptomatic *Ajuga reptans* samples were positive for CMV, AMV or TSV (Table 1). Since all three of these viruses have an insect vector, there is a real possibility of virus spread from asymptomatic *Ajuga reptans* plants or plantings to nearby perennial stock of the same and other species. Without symptoms to draw the grower’s attention, all may appear well. The second implication for the grower is that since *Ajuga reptans* is largely propagated vegetatively, infected but asymptomatic stock plants may be infected. This seems to be a likely scenario since we have been unable to locate any source of *Ajuga reptans* ‘Royalty’ that is virus-free. The striking coloration present in certain cultivars may be caused by viral infection. Yet a third implication may be derived from the presence of CMV satRNA in ‘Royalty’. Although it appears not to cause severe disease in *Ajuga reptans*, it may cause very different symptoms in other hosts. As mentioned previously, the CMV-WL helper strain and satRNA can cause a white leaf syndrome in tomato, but a symptomless infection in tobacco (Gonsalves et al., 1982). A CMV satRNA is also responsible for a lethal necrosis disease of tomato (Kaper and Waterworth, 1977), so being aware of potential sources and reservoirs of both CMV and satRNA is important.

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