Review

Biological control for grapevine crown gall using nonpathogenic *Rhizobium vitis* strain ARK-1

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Abstract: Crown gall of grapevine, which is caused by tumorigenic *Rhizobium vitis*, is the most important bacterial disease of grapevine throughout the world. Screening tests of biological control agents resulted in the discovery of a nonpathogenic *R. vitis* strain ARK-1. By soaking grapevine roots with a cell suspension of strain ARK-1 prior to planting in the field, ARK-1 treatment significantly reduced the number of plants with crown gall symptoms. Several field trials result indicated that ARK-1 was very useful in the field, not only for grapevine but also for various other plant species. In experiments where a mixture of ARK-1 and a tumorigenic strain at a 1 : 1 cell ratio was examined *in vitro* and *in planta*, expression levels of the virulence genes *virD2* and *virE2* of the tumorigenic strain were significantly lower. The suppression of virulence genes, which can result in a reduction of gall formation and the pathogen population, seems to be a unique mechanism of ARK-1. These results indicated that ARK-1 is a promising new agent to control grapevine crown gall.

Keywords: *Rhizobium vitis*, grapevine crown gall, biological control, suppression of virulence genes

1. Introduction

Crown gall disease is one of the most economically important soil-borne bacterial plant diseases around the world.1) Symptoms of crown gall are identified as hypertrophy (increase in size) and hyperplasia (increase in the number) of plant cells, which results in the formation of “galls” or tumors on roots and/or at the base or “crown” of woody plants such as pome fruit (e.g., apple, pear), stone fruit (e.g., cherry, apricot), and nut (e.g., almond, walnut) trees.1) Abnormal cellular development is due to DNA transfer from the crown gall pathogen into the plant.1) The transferred bacterial DNA is incorporated into and expressed in the plant genome (Fig. 1), and it represents the only known example of DNA transfer from a bacterium to a plant in nature.1) Other hosts of crown galls include ornamental woody crops such as roses, Marguerite daisies, and *Chrysanthemum* spp., as well as vines and canes such as grapevines and raspberries.1) Occasionally, field crops such as cotton, sugar beets, tomatoes, beans, and alfalfa can develop crown galls, but the economic impact on these crops is not very high.1)

Grapevine crown gall, as one of the most important diseases of grapevine around the world,2)3) is caused mainly by *Rhizobium vitis* (Ti) [= *Agrobacterium vitis* (Ti), *A. tumefaciens* biovar 3], where “Ti” indicates “tumor-inducing” or “tumorigenic”. Today, *R. vitis* (Ti) is recognized as the predominant species causing grapevine crown gall. Based on reports over the past 30 years, the list of countries with grapevine crown gall includes China, Japan, South Africa, Chile, Israel, many countries in Europe, the Middle East, and many countries in North and South America.3) Whereas in the majority of grape-
growing regions, freezing injuries provide sites for initiating crown galls, in South Africa and Israel, high temperatures and humidity are considered equally important. Crown galls commonly form on the trunks and cordons of grapevines (Fig. 2a, b); however, symptoms can even develop on 1-year-old seedlings (Fig. 2c). Infected grapevines often produce inferior growth, and in some cases, the disease may cause partial or complete grapevine death (Fig. 2d).

Virulent Rhizobium strains transfer single-stranded forms of T-DNA and several virulence effector proteins through a bacterial type IV secretion system into plant host cells. Virulent Rhizobium Ti strains transfer single-strand forms of T-DNA and several virulence effector proteins through a bacterial type IV secretion system into plant host cells. Pathogenicity genes of Rhizobium Ti strains are mostly located on large tumor-inducing plasmids (pTi). A part of this plasmid (T-DNA) is transferred and inserted into the nuclear DNA of the plant during infection. Subsequent expression of T-DNA genes results in the overproduction of auxins and cytokinins, which eventually leads to abnormal gall formation in the host plant. DNA genes then produce tumor-specific compounds called opines, which serve as nutrients for R. vitis.3)

Crown gall infection takes place not only in vineyards but also in nurseries. With nursery production, symptoms develop at the site of wounds made by disbudding, at the base of rooted cuttings, and at grafts; however, in many cases, the infected plants remain symptomless until frost or other physical damage initiates the disease.5) Since nurseries
are often located in places where the risk of frost is low, infected vines most likely do not develop obvious symptoms. Therefore, \textit{R. viti}s is often transmitted through the vegetative propagation of infected asymptomatic grapevines. When mother vines at a nursery are infected, the pathogen can be spread very quickly through the production and dissemination of nursery stocks.

The most serious problems we face are that there are no effective control methods against grapevine crown gall. The nonpathogenic \textit{R. rhizogenes} [= \textit{A. radiobacter} biovar 2] strain K84 has been used successfully to control crown gall in many plant species.\cite{9,10} An agrocin produced by K84 (agrocin 84), which is thought to be the primary factor in the control,\cite{11} targets a tRNA synthetase in tumorigenic \textit{Rhzobium} strains.\cite{12} However, K84 does not prevent the initial infection of grapevine by \textit{R. viti}s (Ti),\cite{3,8,13–16} Although several laboratories have attempted to develop other biological control agents for grapevine crown gall and reported some potential bacterial strains, they have not made a successful development until now.

Previously, we reported that some nonpathogenic \textit{R. viti}s strains, which were isolated from grapevine nursery stock in Japan, inhibited tumor formation in some plant species including grapevine.\cite{13–16} In this review article, we focus on nonpathogenic \textit{R. viti}s strain “ARK-1”, which was identified in our lab and strongly inhibited tumor formation in grapevine in fields, as a new antagonistic strain.

2. The crown gall problem casts a dark shadow over vineyards

Grapes are an important agricultural product because they have a variety of uses, such as table grapes, wine production, juice production, or for drying into raisins. In particular, there is a great demand for wine production around the world, including the recent increase in popularity in the Japanese wine industry. A global examples that can be relevant to Japan is the State of Virginia, located in the mid-Atlantic region of the US. Although the history of wine grape production in Virginia started in 1860’s when settlers from Europe brought grapevines to the New World, the modern Virginia wine industry did not start until the mid-1980’s due to many disease and insect problems.\cite{19} Currently, Virginia ranks fifth in wine grape production in the US, with 1,416 hectares of wine grape production (mainly \textit{Vitis vinifera}, and inter- or intra-specific hybrids, and a very small production of \textit{V. labrusca}) and over 260 wineries.\cite{20} After only three decades, the estimated annual economic income from the Virginia wine industry has reached $1.3 billion (approximately 156 billion Japanese yen) in 2015 (new stats). With the Eastern part of Virginia facing the Atlantic Ocean and the Western part including the Appalachian Mountains, Virginia has a diverse range of climates.\cite{21} Severe winter weather, as well as recent trends in extreme temperature fluctuations during late winter and early spring, tends to cause damage to grapevine trunks, which allows the entry of \textit{R. viti}s.\cite{3}

Therefore, vine growers in Virginia, as well as in the other northern temperate climates, including Nagano and other northern prefectures in Japan, have been suffering from damage from crown gall on grapevines. In particular, the winters of 2013 and 2014 caused significant winter injury to grapevines in Virginia and many states north of Virginia, and many growers lost a considerable amount of grapevines due to crown gall. The level of damage was higher with less winter-hardy cultivars, such as Merlot and Tannat. For example, a 5-year-old Merlot vineyard located in the northern Virginia lost 100% of vines due to a combination of winter injury and crown gall (Nita, personal communication). Merlot is one of the most commonly grown wine grape cultivars in Japan.\cite{22} The unpredictable weather patterns, lack of management strategies, and destructive nature of this disease have made crown gall one of the most difficult diseases for northern temperate climate viticulture.

3. Screening tests for biological control agents

3.1. Co-inoculation of plant stems with pathogens/nonpathogens. Results of a field survey of potential \textit{R. viti}s (Ti) infections of mother plants and nursery stocks of grapevine in Okayama Prefecture, Japan, revealed that there were numerous nonpathogenic strains isolated on selective medium.\cite{13} Most of these strains were identified as nonpathogenic \textit{R. viti}s using pathogenicity tests, bacteriological characteristics, specific multiplex polymerase chain reaction (PCR), and 16S rDNA sequencing analysis.\cite{13,16}

Screening tests for biological control agents using nonpathogenic \textit{R. viti}s by a needle-prick method were carried out.\cite{13,14,16} with a total of 306 nonpathogenic strains to determine their activities against a pathogenic strain. Cell suspensions of the pathogenic and nonpathogenic strains were prepared
and mixed in various combinations at cell ratios of 1:1. The inoculation consisted of a drop of the mixed cell suspension placed into a needle-prick wound on the stem of a tomato or grapevine seedling. We found that using tomato seedlings was appropriate in this screening test because tomato seedlings grow very fast from seeds and tumors formed easily on the stems by inoculation with Ti (tumorigenic) strains.\(^{13}\)

In a test with a 1:1 cell ratio of pathogen/nonpathogen onto stems of tomato and grapevine, some strains greatly suppressed tumor incidence and the tumor size in stems compared to stems inoculated with only a pathogenic strain. Then, nonpathogenic \textit{R. vitis} strains ARK-1, ARK-2, ARK-3, and VAR03-1 were selected as biological control agents against grapevine crown gall.\(^{16}\)

There are at least five genotypes in \textit{R. vitis} (Ti) based on multilocus sequencing analysis of housekeeping genes \textit{pyrG}, \textit{recA}, and \textit{rpoD}\(^{23}\). On grapevine seedlings, a 1:1 cell ratio of each strain ARK-1, ARK-2, ARK-3, or VAR03-1 to the mixture of seven different \textit{R. vitis} (Ti) strains belonging to five different genotypes, which were isolated from Japan, Australia, and Greece, significantly suppressed tumor incidence in stems. Among these, strain ARK-1 was found to be the strongest among the strains tested for inhibiting tumor formation. ARK-1 reduced tumor formation by an average of 90.6% of the positive control, which was the occurrence of tumor formation after inoculating with the pathogenic strain alone (Fig. 3, 4). So far, there have been no reports other than ours that any antagonistic strains strongly inhibited tumor formation on tomato and grapevine, as ARK-1 did using the needle prick inoculation at a 1:1 cell ratio. ARK-1 is a promising new agent to control grapevine crown gall.

Nonpathogenic \textit{R. rhizogenes} strain K84, which was known as a nonantagonistic strain for control grapevine crown gall, was not able to inhibit tumor formation on grapevine shoots (Fig. 4)\(^{2,13–16}\). In addition, a nonpathogenic \textit{R. vitis} strain VAR06-30 was not able to inhibit tumor formation (Fig. 4) and was not an antagonist for grapevine crown gall.\(^{27}\) We are interested to clarify the mechanism of control of ARK-1 and why ARK-1 is overwhelmingly powerful in suppressing tumor formation compared with K84 and VAR06-30, and we discuss it in a later section.

### 3.2. Biological control by a root dipping method

\textit{R. vitis} (Ti) usually infects plants from the soil into wounded roots. Pruned grapevine roots were soaked for 1 h in a cell suspension (10^8 cells/ml) of nonpathogenic strains ARK-1, ARK-2, ARK-3, or VAR03-1. After soaking, treated grapevine seedlings were planted in pots with soil infested with a mixture of seven \textit{R. vitis} (Ti) strains at a final concentration of 5 x 10^7 cells/g soil and grown in greenhouses.\(^{14–16}\) We carried out four experiments. Tumors formed on roots and a stem were studied. Treatment of grapevine roots with strains ARK-1, ARK-2, ARK-3, or VAR03-1 significantly reduced the percentage of grapevine seedlings with tumors caused by the Ti strains.\(^{14–16}\) Treatment using ARK-1 showed the highest control effect and reduced the incidence of galled plants by an average 85.2% of those on nontreated grapevine seedlings.\(^{16}\) Thus, we verified that pre-inoculation of grapevine roots with ARK-1 was effective in reducing the percentage of plants with galls. These positive results encouraged us to conduct field trials on the assumption of the practical agricultural condition.

### 4. Field trials for practical use

Field trials are an essential part of the development of new agricultural technology. Field trials are especially important in developing biological control agents. Even though good results may be produced in laboratory experiments, field trials often do not show the expected results. Thus, we conducted field trials to verify the control effect of strain ARK-1 by root dipping method with nursery stock of grapevine in several vineyards in Japan.

From 2009 to 2013, nine field trials designed as randomized or systematic controlled trials of biological control for grapevine crown gall were carried out at three different experimental fields (A to C) in Okayama, Japan. All field trials used nursery stock of grapevine (\textit{Vitis vinifera} × \textit{V. labrusca} cv. Pione, or \textit{V. vinifera} cv. Neo Muscat). The experimental fields were artificially infected with seven typical strains of \textit{R. vitis} (Ti) belonging to five different genotypes.\(^{25}\) Roots of plants were soaked for 1 h in a cell suspension of strain ARK-1 (10^9 cells/ml) or water, and were then planted in each plot. The arrangement of each plot was random or systematic within each field. Tumor formation on roots and stems of plants was investigated after 6–9 months.

The disease incidences in the nine field trials were subjected to meta-analysis according to a random effects model using the “DerSimonian–Laird method”, because the field trials were performed with different plots sizes, numbers of plots, and plants, farms, and years. Meta-analysis is a set of statistical procedures for synthesizing research results from a number of different studies, and the DerSimonian–
Fig. 2. The typical symptoms of grapevine crown gall: (a) Tumors commonly form on the trunks of grapevines; (b) Tumors commonly form on the shoots of grapevines; (c) Tumors even develop on 1-year-old seedlings; and (d) Infested grapevines often produce inferior growth, are inclined to drop leaves, and disease may cause partial or complete grapevine death.

Fig. 3. Effect of strain ARK-1 on tumor formation. (a) A stem of grapevine was inoculated with mixtures of the R. vitis (Ti) strain as only pathogen. White arrows indicate tumors forming at the inoculation wound site (b) A stem of grapevine was inoculated with mixtures of R. vitis (Ti) strain and ARK-1 at the same time, but at a 1 : 1 cell ratio. White arrows indicate no tumors formation on inoculation wound site. The photograph was taken approximately 3 months after inoculation.
Laird method can incorporate variations among studies. An estimate of a statistical effect, such as the difference in disease severity for plants with or without treatment, is collected from each study along with a measure of the variance of the estimate of the effect. The effect size of antagonist treatment was calculated as an integrated risk ratio. The risk ratio was defined as Risk ratio = (proportion of plants with tumors in antagonist treatment)/(proportion of plants with tumors in water treatment).

As shown in Fig. 5, the integrated relative risk was 0.18 (95% confidence interval: 0.10–0.32, \( p < 0.001 \)), indicating that ARK-1 treatment significantly reduced the disease incidence. The integrated relative risk value of 0.18 indicated that the incidence of crown gall disease during treatment with ARK-1 was decreased to 18% of that without ARK-1, demonstrating that the control effect was very high in the fields. Thus, the integrated relative risk value of 0.18 indicated that ARK-1 very useful in the field. The heterogeneity between each study was tested in a meta-analysis of the inhibitory effects of ARK-1. The I-squared, tau-squared, and \( p \)-values were 0, 0, and 0.97, respectively, indicating that heterogeneity did not exist between field trials and that the integrated relative risk was mainly derived from the inhibiting effect of ARK-1. There are no reports of a biological control agent that is better than ARK-1 for inhibiting tumor formation on the grapevines in the field.

Meta-analysis of the seven field trials showed strong evidence that ARK-1 was effective in controlling grapevine crown gall by application in the field. The results indicated that soaking for 1 hour in a cell suspension of \( 10^8 \) cells/ml of ARK-1 was suitable for practical use. The effectiveness of ARK in controlling crown gall at a low cell density was advantageous in developing ARK-1 as a biopesticide. At present, we verified that \( 5 \times 10^7 \) cells/ml of ARK-1 gave a positive effect with a relative risk value 0.15 (95% confidence interval: 0.03–0.27, \( p < 0.05 \)) on inhibition of gall formation in some field trials.

![Fig. 4. Effect of co-inoculation with nonpathogenic *Rhizobium* strains and Ti strain at a 1:1 cell ratio on grapevine seedlings. Data are the means from three experiments. Error bars indicate 95% confidential intervals. Rhombi labeled with different letters indicate a significant difference from the other bars (\( P < 0.05 \), Ryan’s multiple test).](image)

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![Fig. 5. Integrated evaluation based on a meta-analysis of the effect of nonpathogenic *R. vitis* strains ARK-1 on grapevine crown gall after soaking plant roots in bacterial cell suspensions in field trials. The center and width of the diamond shape demonstrate the value of the integrated relative risk and 95% confidence interval, respectively.](image)

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**Table:**

<table>
<thead>
<tr>
<th>Field trials</th>
<th>Plants with tumors</th>
<th>Plants with tumors</th>
<th>Total plants</th>
<th>Risk ratio</th>
<th>95% confidence interval</th>
<th>Weight (%)</th>
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<td>2009-A</td>
<td>1</td>
<td>30</td>
<td>8</td>
<td>0.13</td>
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<td>7.8%</td>
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<tr>
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<td>1</td>
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<td>14</td>
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<td>4</td>
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<tr>
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<td>36</td>
<td>7</td>
<td>0.14</td>
<td>[0.02; 1.10]</td>
<td>7.6%</td>
</tr>
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<td>2011-A</td>
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<tr>
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<td>14</td>
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<td>15.9%</td>
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<td>14</td>
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<td>[0.10; 0.81]</td>
<td>29.6%</td>
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<tr>
<td>2012-C</td>
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<td>4</td>
<td>0.25</td>
<td>[0.03; 2.11]</td>
<td>7.0%</td>
</tr>
<tr>
<td>2013-A</td>
<td>0</td>
<td>38</td>
<td>2</td>
<td>0.21</td>
<td>[0.01; 4.24]</td>
<td>3.5%</td>
</tr>
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</table>

Random effects model 282 284 0.18 [0.10; 0.32] 100%
In addition to grapevines, we have evidence that ARK-1 can effectively control apple, Japanese pear, peach, rose, and tomato crown gall caused by R. radiobacter (Ti) and R. rhizogenes (Ti) strains, based on four fields and three greenhouse trials.\textsuperscript{18} Therefore, ARK-1 effectively protects six different species (grapevine, apple, Japanese pear, peach, rose, and tomato) of host plants against three different Rhizobium Ti strain species (R. radiobacter (Ti), R. rhizogenes (Ti), and R. vitis (Ti)).

5. Population dynamics of ARK-1 on roots of grapevine

In order to determine the rate of ARK-1 survival on grapevine roots in the soil, an antibiotic-resistant mutant of ARK-1, ARK-1sc, was developed. ARK-1sc was a streptomycin (St)-copper sulfate (CuSO\textsubscript{4})-resistant mutant (St-CuSO\textsubscript{4}-mutant) obtained by growing strain ARK-1 on a special culture medium with St and CuSO\textsubscript{4} named “St-CuSO\textsubscript{4} Potato Sucrose Agar (PSA) medium”.\textsuperscript{25,26} The ARK-1sc strain was inoculated onto roots using the root dipping method, and at 1-, 6-, 18-, 24-, 30-, and 36-month after inoculation, the colony forming units (CFU) of ARK-1sc were counted on St-CuSO\textsubscript{4}-PSA media. As shown in Fig. 6, 12 months after inoculation with strain ARK-1sc, the bacterial population inside roots was 2 × 10\textsuperscript{6} CFU/g (fresh weight) of the root. Colonization by ARK-1sc inside roots remained at 5 × 10\textsuperscript{4} CFU/g of the root for up to 24 months, and then dropped to 6 × 10\textsuperscript{3} CFU/g of root after 36 months. On the other hand, at 12 months after inoculation with strain ARK-1sc, the bacterial population on the root surfaces was 2 × 10\textsuperscript{4} CFU/g of the root. Colonization by ARK-1sc on root surfaces decreased to 2 × 10\textsuperscript{2} CFU/g of root after 24 months, and after 30 months, ARK-1sc was not detected on root surfaces.

Thus, the result for ARK-1sc indicated that strain ARK-1 not only established populations in the rhizosphere of grapevine but also persisted inside roots for up to 3 years. On the other hand, colonization of ARK-1sc on root surfaces decreased to the detection limit after 30 months. These comparisons of ARK-1sc survival inside roots and on roots surfaces demonstrated that the bacterial population on root surfaces was always lower than that inside the roots. ARK-1 was originally isolated from internal tissues of grapevine,\textsuperscript{13} indicating that ARK-1 is an endophytic bacterium. This phenomenon needs further investigation because the ability of ARK-1 to colonize grapevine roots might affect the persistence of the control of crown gall by ARK-1. It is also important to determine in which grapevine tissue ARK-1 colonizes. This can be investigated by visualizing bacterial colonization patterns with the use of a luminous protein (e.g., green fluorescent protein).

6. Understanding the mechanism of strain ARK-1

6.1. Need live-cells of ARK-1. In order to understand the mechanism of the biological effects of ARK-1, an inoculation study was conducted with three different forms of ARK-1: a cell suspension of ARK-1 as “live cells”; a cell suspension of ARK-1 autoclaved at 121 °C in 15 minutes as “dead cells”; and a culture filtrate of ARK-1 by filtering with a syringe filter unit (0.2 µm diameter) as “culture filtrate (CF)”. Each treatment consisted of a mixture of ARK-1 cells and an R. vitis (Ti) (tumorigenic) strain cells at 1 : 1 ratios, and these were inoculated onto stems of grapevine seedlings using the needle-prick methods described above.\textsuperscript{13,14,16}

Results showed that the live strain ARK-1 in a cell suspension significantly suppressed tumor incidence in stems compared to the control treatment using Ti strains (protective value = 89.1%).\textsuperscript{16} On the other hand, the dead cells and CF suspension did not reduce tumor incidence (both protective values = 6.4%).\textsuperscript{16} Basically, the mechanism of antagonistic microorganisms as biological control agents for plant disease involves bacteriocins or antibacterial materials. However, the lack of control by dead cells
and CF suspension indicated a different mechanisms for ARK-1 and that ARK-1 may have cellular functions to control crown gall formation.

6.2. Reducing the pathogen population at the wound site. A survival assay of ARK-1 was conducted. To differentiate the inoculated biological control agents from indigenous rhizobacteria, _R. vitis_ strains used in the experiment consisted of three antibiotic-resistant mutant strains, ARK-1sc, VAR06-30sc, and VAT03-9n. ARK-1sc, as described above, is an St-CuSO₄ mutant obtained by growing ARK-1 on St-CuSO₄-PSA medium.²⁵,²⁶ Nonpathogenic _R. vitis_ strain VAR06-30 was a nonantagonistic strain that was not able to inhibit tumor formation (Fig. 4), and a St-CuSO₄ mutant obtained by growing VAR06-30 on St-CuSO₄-PSA medium was called VAR06-30sc.²⁷) _R. vitis_ (Ti) strain VAT03-9 was a tumorigenic strain isolated from grapevine,¹³ and VAT03-9n was a nalidixic acid (nal)-resistant mutant (nal-mutant) obtained by growing on nal-PSA medium.²⁷) The three mutants grew on the St-CuSO₄-PSA medium (ARK-1sc and VAR06-30sc) and the nal-PSA medium (VAT03-9n) at rates similar to the wild-type in PSA medium.²⁷)

Inoculation to grapevine seedlings took place with one of three different cell suspensions: 1) ARK-1sc and VAT03-9n (1 : 1 ratio); and 2) VAR06-30sc and VAT03-9n (1 : 1 ratio). Additional experimental runs consisted of grapevine seedlings inoculated with the each cell suspension of ARK-1sc, VAR06-30sc, or VAT03-9n to demonstrate the population dynamics with single strain inoculation.

Populations of ARK-1sc on the plants resulted in significantly higher number (P < 0.05) than those of VAT03-9n at 7 and 9 days after inoculation (da) (Fig. 7a). On the other hand, populations of VAR06-30sc and VAT03-9n did not significantly differ in numbers up to 9 dai (Fig. 7b). Single inoculation of ARK-1sc, VAR06-30sc, and VAT03-9n did not result in a significant difference in population level even after 9 dai (Fig. 7c). These results suggested that ARK-1sc could not reduce the pathogen population at the wound site on the grapevine plants during the early period after infection, but after 7 dai, suppressed the population of VAT03-9n. The other result was obtained that colonization by ARK-1sc remained roughly constant at 4 × 10⁷ CFU/g of grapevine shoot for up to 88 dai, but colonization by Ti strain decreased to 1 × 10⁶ CFU g⁻¹ of the grapevine shoot after 88 dai.²⁶) In the co-inoculation assay, the populations of Ti strain were approximately 10⁷ CFU/g in grapevine shoots with tumors and 10⁶ CFU/g in shoots without tumors.²⁶) These results demonstrated that ARK-1sc reduced the population of the tumorigenic bacterial population to significantly lower level by an order of ten-fold. Therefore, one of the mechanisms allowing the biological activity of ARK-1 is a significant inhibition of a population development of the Ti strain from 7 dai.

6.3. Suppressing the expression of essential virulence genes in the tumorigenic strain. Transformation of the host DNA by _R. vitis_ would most likely to occur prior to 5 dai.²⁸) The suppression of gall formation with co-inoculation of ARK-1 and the Ti strain occurred even with a lack of suppression of the Ti strain population by ARK-1 until 5 dai (Fig. 7a). This result indicated that the mechanism by ARK-1 to reduce crown gall symptoms might not involve the suppression of the Ti strain population in grapevine plants.

The virulence (vir) genes and T-DNA are located mostly on pTi and chromosomal virulence genes (chev) on chromosomes.⁶,⁷) The plant molecules acetosyringone and α-hydroxycetosyringone induce the expression of the entire vir regulon in _Rhizobium_ as well as the formation of T-DNA intermediate molecules.⁷) These plant molecules occur specifically in the exudates from wounded and metabolically active plant cells, thus, they probably allow _Rhizobium_ to recognize susceptible cells in nature.⁷) T-DNA transfer and processing require products of the vir genes (virA, virB, virC, virD, virE, and virG), which are located outside of the T-DNA coding region.⁶,⁷,²⁹,³⁰) The expressions of virB, virC, virD, and virE are positively regulated at the transcriptional level by plant signal molecules.⁶,⁷,²⁹) Two tumorigenic _Rhizobium_ proteins, VirD2 and VirE2, directly associate with the T-DNA strand; one VirD2 molecule covalently attaches to the 5'-end of the T-DNA strand, and VirE2, a protein that binds single-stranded DNA, cooperatively coats the rest of the T-DNA strand (Fig. 1).⁶,⁷,²⁹)

To provide insights into that mechanism, we investigated the effect of ARK-1 on suppressing the expression of two virulence genes, virD2 and virE2, by the Ti strain; this suppression appeared to be responsible for the biological control by ARK-1.

6.3.1. Suppressive effect of ARK-1 co-inoculation on the expression of the vir genes of the Ti strain. To clarify whether ARK-1 suppresses the expression of vir genes of Ti strain VAT03-9 in planta, the expression of virD2 and virE2 was analyzed in grapevine plants co-inoculated with ARK-1 and
Shoots of grapevine seedlings were inoculated with nonpathogenic strains (ARK-1 or VAR06-30) and Ti strain VAT03-9 at a cell ratio of 1:1, and CF of ARK-1 plus VAT03-9 by the needle-prick method as described above. Shoot samples containing one wound site were collected, and bacterial RNA was extracted. The amount of mRNA expression of virD2 and virE2 of the Ti strain were analyzed using a reverse-transcription quantitative polymerase chain reaction (RT-qPCR).

In plants inoculated with only VAT03-9, the induction of bacterial virD2 and virE2 expression was detected at 1 dai (Fig. 8). No significant difference in the expression levels of virD2 and virE2 were observed upon co-inoculation with VAR06-30 or CF of ARK-1 and VAT03-9 in comparison with inoculation with only VAT03-9 (Fig. 8). On the other hand, virD2 and virE2 expression levels were significantly suppressed upon co-inoculation with ARK-1 and VAT03-9 (Fig. 8). These results suggested that treatment with ARK-1, which inhibited tumor formation, could suppress the expression of these genes, hence, and they could not inhibit tumor formation.

To evaluate the starting and ending time points of the suppression of vir gene expression by ARK-1, virD2 expression was analyzed from 1 hour after inoculation. The results showed that the induction of virD2 expression was significantly suppressed in the plants inoculated with ARK-1 and VAT03-9, while no significant suppression was observed in the plants inoculated with VAR06-30 or CF of ARK-1.

Fig. 7. Population dynamics of the nonpathogenic strains and VAT03-9n after co-inoculation onto grapevine plants. (a) Populations of the nonpathogenic strain ARK-1sc and of the tumorigenic (Ti) strain VAT03-9n after co-inoculation on wounded shoots of grapevine seedlings at a 1:1 cell ratio. (b) Populations of the nonpathogenic strain VAR06-30sc and of the Ti strain VAT03-9n after co-inoculation on wounded shoots of grapevine seedlings at a 1:1 cell ratio. (c) Populations of the nonpathogenic strain ARK-1sc, of strain VAR06-30sc, and of the tumorigenic (Ti) strain VAT03-9n after each inoculation on wounded shoots of grapevine seedlings. Data are means ± standard deviation for five plants. (a) and (b) Significant differences at a given point in time are indicated by * (t-test, *P < 0.05; **P < 0.01; ***P < 0.005). (c) No significant differences at a given point in time are indicated by ns (Tukey’s HSD test, ***P ≥ 0.05). dai, days after inoculation.
inoculation (hai) to 4 dai in grapevine plants co-
inoculated with ARK-1 and VAT03-9 (Fig. 9). The level of virD2 expression after an inoculation with VAT03-9 was significantly higher at 1 dai than at any other time (Fig. 9). The levels of virD2 expression were significantly lower with a co-
inoculation with ARK-1 and VAT03-9 than with a single inoculation of VAT03-9 at 12 hai, 1 dai, and 2 dai (Fig. 9). These results suggested that co-
inoculation with ARK-1 was able to suppress virD2 expression by the Ti strain in grapevine plants during the peak period of virD2 gene expression. However, the effect of ARK-1 became unclear at 3 and 4 dai because virD2 expression in plants inoculated with only the Ti strain (VAT03-9) was also low during this period.

6.3.2. Suppressive effect of ARK-1 co-inoculation on vir gene expression of Ti strain in culture media with or without acetosyringone. The kinetics of virD2 and virE2 expression were analyzed in culture media with or without acetosyringone to examine whether the suppression of the vir genes was indeed caused by ARK-1. Four cell suspensions (ARK-1 plus VAT03-9, VAR06-30 plus VAT03-9, and CF of ARK-1 plus VAT03-9) at cell ratios of 1:1 were inoculated onto the stems of grapevine plants after wounding. Stem samples were harvested at 1 dai. Data are means ± standard deviation for samples corresponding to individual plants. Bars labeled with different letters indicate a significant difference from the other bars (P < 0.01, Tukey’s HSD test). CF, culture filtrate.

Fig. 8. Relative expression levels of the virD2 and virE2 genes in bacterial cells of the R. vitis Ti strain in grapevine plants measured using reverse-transcription quantitative polymerase chain reaction (RT-qPCR). Values (%) represent the relative expression levels of the vir genes at 1 day after inoculation (1 dai) compared with the value when only Ti strain VAT03-9 was inoculated (“only VAT03-9”), which had a value of 100%. Each cell suspension or mixture (ARK-1 plus VAT03-9, VAR06-30 plus VAT03-9, CF of ARK-1 plus VAT03-9, and only VAT03-9) at cell ratios of 1:1 was inoculated onto the stems of grapevine plants after wounding. Stem samples were harvested at 1 dai. Data are means ± standard deviation for samples corresponding to individual plants. Bars labeled with different letters indicate a significant difference from the other bars (P < 0.01, Tukey’s HSD test). CF, culture filtrate.

Fig. 9. Changes in the relative expression levels of the virD2 gene in grapevine plants measured using the reverse-transcription quantitative polymerase chain reaction (RT-qPCR). Values (%) represent the relative expression levels of the vir genes at 1 day after inoculation (1 dai) compared with the value when only Ti strain VAT03-9 was inoculated (“only VAT03-9”), which had a value of 100%. A mixed cell suspension of the nonpathogenic strain ARK-1 and of the Ti strain VAT03-9 at cell ratios of 1:1 or a single cell suspension of VAT03-9 were inoculated onto the stems of wounded grapevine plants. Stem samples were harvested at times ranging from 1 hour after inoculation (hai) to 4 dai. Data are means ± standard deviation for samples corresponding to individual plants. Significant differences at a given point in time between the ARK-1 plus Ti mixture and only the Ti strain are indicated by * (t-test, *P < 0.05; **P < 0.01; ***P ≥ 0.05); bars for only the Ti strain that are labeled with different letters differ significantly from other bars for only the Ti strain (Tukey’s HSD test, P < 0.05).
performed for the detection of the was extracted after incubation, and RT-qPCR was inoculated with one of the mixed suspensions PS-AS medium (PS medium with acetosyringone), were prepared as described above. PS medium or with a VAT03-9 cell suspension. Bacterial RNA were prepared as described above. PS medium or PS-AS medium (PS medium with acetosyringone), was inoculated with one of the mixed suspensions or with a VAT03-9 cell suspension. Bacterial RNA was extracted after incubation, and RT-qPCR was performed for the detection of the *virD2* and *virE2* genes and relative quantification of the mRNA concentrations.

In the PS-AS medium, *virD2* and *virE2* expression levels in VAT03-9 cells were significantly reduced by co-incubation with ARK-1 (Fig. 10); however, the expression level was not significantly affected by co-incubation with VAR06-30 or CF of ARK-1 (Fig. 10). These results suggested that the presence of ARK-1 can suppress *virD2* and *virE2* expression by the Ti strain in a culture medium that contains acetosyringone.

### 7. Discussion

Newly introduced nonpathogenic and antagonistic *R. vitis* strain ARK-1 is endophytic in grapevine and strongly inhibits tumor formation in grapevine. We also applied ARK-1 as a biological control of crown gall disease, and demonstrated that ARK-1 was able to control grapevine crown gall in field trials. Figure 11 shows our hypothesis of the mechanism of the effect of ARK-1 based on our research results until now. ARK-1 produced a zone of inhibition against tumorigenic *Rhizobium* spp. in *in vitro* assays; however, further examinations revealed that antibiosis was not the main mechanism of biological control. ARK-1 also reduced the pathogen population at the wound site; however, it seems that the suppression of virulence gene expression is the key mechanism to reduce gall production.26) Confirmation of suppression of two *vir* genes (*virD2* and *virE2*) using the RT-qPCR was conducted in both *in vitro* and *in planta* experiments. The products of these genes, VirD2 and VirE2 proteins, are directly associated with the transformation of plant cells by *Rhizobium* strains of *R. vitis*, resulting in gall formation. The suppression occurred prior to the reduction in the pathogen population.10,17) This mechanism is unique to ARK-1, because it is different from previously reported mechanisms of biological control of plant crown gall disease. Two known mechanisms of biological control of crown gall are antibacterial compounds produced by nonpathogenic strains11),13),31),32) and a unique mechanism associated with the quorum-sensing and caseinolytic protease (clp) systems of nonpathogenic *R. vitis* strain F2/5.33) We will investigate whether ARK-1 could suppress the expression of *vir* genes other than *virD2* and *virE2* in grapevine in the near future.

The suppression of *vir* genes by ARK-1 occurred a few days after inoculation; however, the pathogenic population did not change until 7 dai. This result suggested that the suppression of *vir* gene expression probably inhibited the transformation of the host plant cells, thereby preventing the Ti strain from acquiring and catabolizing opines, which are amino acids produced as a result of transformation. Without the advantage of the opines as a food source, a growth of the Ti strain might suffer and result in a
lower population than the ARK-1 population in grapevines. This hypothesis should be tested in future research.

In conclusion, ARK-1 suppressed the expression of the virulence genes of the Ti strain at the wound site and suppressed the development of crown gall disease via what appears to be a previously unreported mechanism. Even with these findings, several important questions remain to be addressed. More field validations of ARK-1 in different vineyard conditions in other countries around the world is necessary to verify the repeatability of biological control of crown gall by ARK-1 treatment. Testing in different environmental conditions will also aid our understanding of the mechanism of suppression provided by ARK-1. Our research projects may contribute to controlling crown gall in not only grapevines but also diverse crop species throughout the world in the near future. The ultimate goal in conducting this study is to evaluate strain ARK-1 as a biopesticide for the control crown gall disease. At present, we are developing a new biopesticide made from ARK-1 and getting positive results that the new biopesticide treatment is effective in controlling crown gall in grapevine and other plant species in field trials. We hope that a new biopesticide made from ARK-1 obtained from our studies will contribute to managing plant crown gall disease in agriculture around the world.

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References


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Profile

Akira Kawaguchi was born in Okayama, Japan, in 1976. He graduated from the Faculty of Agricultural Science, Kyushu University in 2000. He was promoted to researcher in 2002 at the Agricultural Experimental Station, Okayama Agricultural General Center, and he obtained his Ph.D. in 2007 from Okayama University. He was promoted to senior researcher in 2009 at Agricultural Experimental Station, Okayama Agricultural General Center. He was posted in 2010 at Agricultural Research Institute, Okayama Prefectural Technology Centre for Agriculture, Forestry and Fisheries. He was promoted to chief in 2015 at the Department of Agriculture, Forestry and Fisheries, Okayama Prefectural Government. He was promoted to researcher at the Western Region Agricultural Research Center, National Agricultural and Food Research Organization (NARO). He has studied biological control using a new antagonistic microorganism to manage the soil-borne grapevine disease “crown gall”, which occurs all over the world. His research achievements include the discovery of a new antagonistic bacterial strain named ARK-1, evaluation of the positive effect of ARK-1 to control crown gall on not only grapevine but also other plant species in field trials, population dynamics of ARK-1 in grapevine, some mechanisms of the effect of ARK-1, and so on. His final goal is to develop the new biopesticide to control crown gall and clarify the control mechanism of ARK-1 in the near future. For his achievements, he received 12th JSPS Prize and 12th Japan Academy Medal in 2016.