Entomopathogenic Bacteria, *Xenorhabdus*: An Alternative Biocontrol Agent for Integrated Management of Root-knot Nematode on Grapevine

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Abstract

Entomopathogenic bacterium, *Xenorhabdus* has a mutualistic relationship with entomopathogenic nematode of the genus *Steinernema* and produces several bio-agent compounds with antimicrobial and nematocidal activities. Root-knot nematodes are considered one of the most important pests facing the cultivation of grapevine worldwide. A micro-plot field trial was conducted in naturally infested soil with *Meloidogyne incognita* to evaluate the potential of two strains of entomopathogenic bacteria namely *Xenorhabdus budapestensis* DSM 16342 (EMA) and *X. szentirmaii* DSM 16338 (EMC) applied separately or integrated with neem cake and/or furadan at half of recommended dose on nematode development and growth improvement of Taify grapevine. Data of nematode populations, number of galls and egg-masses, eggs/g root, plant lengths and weights and number of leaves were recorded four months after application. Results appeared significant differences between treatments and control. The triple application was more effective than dual and single applications in reducing nematode infestation and improving plant growth. Combined application of EMC or EMA with furadan or neem cake increased the efficacy (64.6-68.6%) and improved plant fresh weight (27.4-69.5%). Conclusively, utilization of such bacterial filtrates with either neem cake and/or nematicide could gain a successful approach in integrated nematode management programs.

Keywords: Entomopathogenic bacteria, neem cake, nematicide, *Meloidogyne incognita*, integrated management, Grapevine.
INTRODUCTION

Xenorhabdus, gram-negative bacterium of the family Enterobacteriaceae, is symbiotic bacteria with entomopathogenic nematode (EPN) of the genus Steinernema. This bacterium (EPB) produces many bioactive compounds which demonstrate insecticidal, nematicidal, cytotoxic and anti-microbial activities. These compounds are evolutionary products developed under strict selective pressure, and comprise a potent chemical against a large scale of eukaryotic and prokaryotic organisms. Many EPN-EPB complexes occur, and many antimicrobial peptide profiles could be established. X. budapestensis DSM 16342 (EMA) and X. szentirmaii DSM 16338 (EMC) are the unique sources of highly efficient antimicrobial peptides against plant pathogens. Root-knot nematodes, Meloidogyne spp. were amongst pathogens that cause serious commercial damage to fruit trees including grapevines. Grapevines infested by nematodes ultimately exhibit destroyed roots, leading to the compulsory replacement of the plants. Multiples classical methods and strategies as crop rotations, resistant rootstocks, nematicides and bio-agents are used for the management of phytonematodes infecting grapevine worldwide. The use of entomopathogenic bacterium, Xenorhabdus sp. has been evaluated and employed against root-knot nematodes infecting Taify grapevine. In laboratory and field trials, S. feltiae–X. bovienii complex had suppressive effects on M. incognita infecting tomato plant. Numerous research papers mentioned that for best nematode management on plants, more compatible materials should be applied. Combined applications of the fungus Verticillium chlamydosporium plus Heterorhabditid matalistic bacterium, Photorhabdus luminescens and compost significantly reduced M. incognita infection and improved cucumber plant growth. The area was designed as a randomized complete block (RCB) and replicated five times. Each block included 12 treated plots and untreated check. A plot consisted of one row, 50 cm × 90 cm was practiced. Roots of two-month-old grapevine seedlings var. Taify with two leaves were soaked in bacterial filtrate or LB medium for 15 min before transplanting. Plots were then planted with three seedlings each. An additional volume of 10 mL bacterial suspension or LB medium was added separately into test tubes containing 5-mL of LB liquid medium as an inoculum for 100 mL culture.

Materials and Methods

Bacterium strains

Xenorhabdus strains, X. budapestensis DSM 16342 (EMA) and X. szentirmaii DSM 16338 (EMC) which had been isolated from the entomopathogenic nematodes Steinernema bicornutum and S. rarius, are originated from the Fodor laboratory, Pannonia University, Keszthely, Hungary. The bacterial isolates were routinely grown in the dark on LBTA (Luria Bertani Agar) indicator plates at 25°C (trypton 10 g/L, yeast extract 10 g/L, sodium chloride 10 g/L, agar 15 g/L, and supplementing with bromothymol blue 25 mg/L, and 2,3,5-triphenyltetrazolium chloride 40 mg/L, 1L of distilled water [pH 6.8]). For preparing bacterial filtrate, single black - dark blue colonies of each bacterium was added separately into test tubes containing 5-mL of LB liquid medium as an inoculum for 100 mL culture. 100 mL aliquots of culture in 500 mL Erlenmeyer flasks were shaken overnight at 25°C, and then transferred to flasks containing 400 mL of the same media, shaking at 200 rpm for 5 days. The multiplied bacterial culture was centrifuged (13,000 rpm for 30 min) at 4°C. A supernatant was filtered through 0.22 μm Millipore filter to obtain cell free filtrate. The filtrate was stored at 4°C until required.

Neem cake

It is a by-product of the cold pressing from the neem seeds and kernels. Fresh seeds and kernels were collected from ripe fruits of 10 years old neem (Azadirachta indica A. Juss.) trees growing in Taif region, Saudi Arabia, then cleaned and dried in the shade for one week. Neem cake was obtained by using cold-pressing vegetable oil machine from compressing neem seed and kernel. After proper drying the formulations were crushed and converted into fine powder using grinder and stored in tin containers at 4°C.

Experimental layout and design

A micro-plot field experiment was conducted in a grapevine farm located at Taif region, Saudi Arabia, to determine the influence of entomopathogenic bacteria (EMA and EMC) and/or neem cake integrated with furadan at half of the recommended dose on nematode reproduction and the resulting effect on grapevine plant growth. The experimental area (35 m²) was heavy naturally infested with Meloidogyne incognita. The area was designed as a randomized complete block (RCB) and replicated five times. Each block included 12 treated plots and untreated check. A plot consisted of one row, 50 cm × 90 cm was practiced. Roots of two-month-old grapevine seedlings var. Taify with two leaves were soaked in bacterial filtrate or LB medium for 15 min before transplanting. Plots were then planted with three seedlings each. An additional volume of 10 mL bacterial suspension or LB medium was added separately into test tubes containing 5-mL of LB liquid medium as an inoculum for 100 mL culture.
was introduced to the surface of the soil per plant and allowed to soak in. Two weeks later, neem cake was added around plants at a rate of 2 g/seedling in a single treatment and 1 g/seedling in concomitant applications, incorporated into the soil and then watered to allow decomposition. At the same time furadan as nematicide was applied singly at 0.6 g/seedling and at half dose (0.3 g/seedling) in integrated treatments. Five untreated plots were served as control. Therefore, a total of 13 treatments including a control viz. (1) EMC, (2) EMA, (3) Neem cake @ 2 g/plant, (4) Furadan 10 G @ 0.6 g/plant, (5) EMC+ Neem cake @ 1 g/plant, (6) EMA+ Neem cake @ 1 g/plant, (7) EMC+ Furadan 10 G @ 0.3 g/plant, (8) EMA+ Furadan 10 G @ 0.3 g/plant, (9) Neem cake @ 1 g/plant+ Furadan 10 G @ 0.3 g/plant, (10) EMC+ Neem cake @ 1 g/plant+ Furadan 10 G @ 0.3 g/plant, (11) EMA+ Neem cake @ 1 g/plant+ Furadan 10 G @ 0.3 g/plant, (12) LB medium (negative control), (13) Check (nematode only) were maintained in this experiment. Seedlings were harvested 4 months after planting and roots were washed free from adhering soil with tap water. Lengths and fresh weights of shoot and root, dry weights and number of leaves were measured. From each plot, a composite soil (250 g) was extracted by sieving and decanting method\textsuperscript{17}. At each treatment, Roots were stained in acid fuchsin\textsuperscript{18} and examined for recording the number of galls, egg-masses and nematode in roots under a stereomicroscope at 40–100X magnification. Eggs were collected using sodium hypochloride technique\textsuperscript{19}. The efficacy of treatments on nematode population was calculated with the equation of Henderson and Tilton. Efficacy \% = \frac{[1-(\text{Total nematode population of treated plants after application} \times 250 \text{ g soil})]}{\text{Total nematode population of check plants before application} / 250 \text{ g soil}} \times 100\textsuperscript{20}. Rate of nematode build-up = \frac{Pf}{Pi}, where Pf is the final population and Pi is the initial population. The index for root galling (GI) and egg-mass (EI) were assessed on a 0-5 scale, where 0 = 0 galls or egg-masses and 5 > 100\textsuperscript{21}.

**Statistical analysis**

Data was subjected to one-way analysis of variance (ANOVA)\textsuperscript{22} followed by Duncan’s multiple range test (P < 0.05) using COSTATE software package and treatment means were compared with the control plants infested with nematodes, according to the Dunnet’s test at P < 0.05 (ns (p < 0.12), * (p < 0.033), ** (p < 0.002) *** (p < 0.001), GraphPad Prism version 7.0). The experiment was performed once.

**RESULTS**

The influence of Taify grapevine seedlings treatment under micro-plot conditions with bacterial filtrate (EMC and EMA) alone or in combination with neem cake and/or nematicide compared to control treatment were studied. Results indicated that all treatments of bacterial filtrates, neem cake and furadan significantly reduced nematode infestation (Fig. 1 and Table 1). Treatments with three integrated components showed the least numbers of J\textsubscript{s} population/250 g soil, nematode in root, galls, egg-masses and eggs/g root. Treatment with filtrate of EMC combined with neem cake and furadan significantly revealed the highest effect on the numbers of J\textsubscript{s} population in soil [F (12, 52) = 389.8; \textit{p}<0.05] (Fig. 1A), nematode in roots [F = 700.2] (Fig. 1B), galls [F = 933.9] (Fig. 1C) and egg-masses [F = 456] (Fig. 1D) when compared with other treatments and control. EMA+ neem+ furadan treatment ranked second to the previous treatment in the same nematode parameters, then application of furadan with EMC or EMA and neem cake plus bacterial filtrates. Single treatments also performed intermediate suppression in the total nematode populations in soil and root, galls and egg-masses. Treatment with furadan, resulted the minimum counts of all nematode infestation criteria followed by EMC treatment, whereas, treatment with neem alone resulted the lowest effect on nematode development (Fig. 1). The population of nematode in both soil and root were suppressed in all treated seedlings to be ranging between 805.6 and 2851.8 in comparison with the control that reached up to 4864.4 (Table 1). The highest efficacies on nematode population were observed when EMC or EMA filtrate was applied in combined form with neem cake and/or furadan as compared to separate allocations. Both treatments of EMC+ neem+ furadan and EMA+ neem+ furadan resulted in the maximum efficacy percentage of 83.9% and 80.7% as well as reduction percentage reached to 83.4% and 80.2%, respectively. Dual application of EMC or EMA with furadan or neem cake occupied the remarkable efficacy that averaged to 76.1%, 73.6% and 66.5%, 64.6% and reduction that valued to 76.4%, 74.3% and 62.5%, 59.7%, respectively. On the other hand, the lowest efficacy treatment was neem cake alone that reached to 54.2% with reduction percentage of 41.4%. While both EMC and EMA applied singly have 57.9% and 55.7% efficacy with reasonable reduction that averaged 48.2% and 43.9%, respectively (Table 1). Data in Table 1 also clarify that all treatments decreased the rate of nematode build-up ranging between 0.35 to 0.99, root galling index (2 to 4), egg-mass index (1.2 to 3.8) and number of eggs/g root (29.8 to 795).

The effect of treatments on growth criteria of the grapevine seedlings including length and weight of shoot and root and number of leaves was recorded (Fig. 2 and Table 2). Results indicated that immersing seedling root in bacterial filtrates of EMC and EMA then treated with neem cake and furadan significantly...
provided best result in all growth parameters compared to other treatments and control (Fig. 2). Based up on lengths of harvested plants, treatment of seedlings with EMC filtrate prior to cultivation then amended with neem cake plus furadan resulted in significant largest plants relative to other treatments and control as measured by their shoot length (F= 60.1) (Fig. 2A) and root length (F= 23.5) (Fig. 2B). EMA+ neem cake+ furadan treatment ranked second in the same criteria, followed by combined application of EMC or EMA plus neem cake, then EMC alone treatment. When seedlings treated with EMA or neem cake plus furadan or neem cake alone, not significant difference was noted in shoot length (Fig. 2A). Separate application of furadan gave the smallest plants when compared to other treatments and nematode alone. The same trend was recorded in root length (Fig. 2B). Data also showed that the greatest shoot and root fresh weights were obtained with the seedlings exposed to EMC or EMA concomitant with neem cake and furadan with significant difference between them and other treatments as well as control. Statistically, there was no significant difference in shoot weight (F= 79.3) among treatments (EMC with EMA+ neem cake), (EMA with EMC+ furadan) and (neem cake with neem cake+ furadan) (Fig. 2C). Regarding root fresh weight, except for single furadan treatment, there was a

![Fig. 1. Influence of treatments with bacterial filtrates, neem cake and furadan in single or concomitant applications on the infection of Taify grapevine seedlings with Meloidogyne incognita under micro-plot field conditions. Error bars represent SD.](image-url)
Table 1. Suppressive effect of entomopathogenic bacteria alone or combined with neem cake and/or furadan on the development of *Meloidogyne incognita* infested Taify grapevine seedlings under micro-plot field conditions

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Initial J.5/250 g soil</th>
<th>Final nematode population</th>
<th>Efficacy%</th>
<th>Reduction%</th>
<th>Rate of build-up</th>
<th>Root gall index</th>
<th>Egg-mass index</th>
<th>No. of Eggs/g root</th>
</tr>
</thead>
<tbody>
<tr>
<td>EMC</td>
<td>2770</td>
<td>2520.4 d</td>
<td>57.9</td>
<td>48.2</td>
<td>0.91</td>
<td>4 c</td>
<td>3 b</td>
<td>401.4 d</td>
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<td>EMA</td>
<td>2850</td>
<td>2728.8 c</td>
<td>55.7</td>
<td>43.9</td>
<td>0.96</td>
<td>4 c</td>
<td>3 b</td>
<td>494.4 c</td>
</tr>
<tr>
<td>Neem</td>
<td>2880</td>
<td>2851.8 c</td>
<td>54.2</td>
<td>41.4</td>
<td>0.99</td>
<td>4 c</td>
<td>3.8 a</td>
<td>795 b</td>
</tr>
<tr>
<td>Furadan</td>
<td>2640</td>
<td>2377 d</td>
<td>58.4</td>
<td>51.1</td>
<td>0.90</td>
<td>4 c</td>
<td>3 b</td>
<td>297.4 e</td>
</tr>
<tr>
<td>EMC + Neem</td>
<td>2520</td>
<td>1826 e</td>
<td>66.5</td>
<td>62.5</td>
<td>0.72</td>
<td>3.8 c</td>
<td>3 b</td>
<td>203.4 f</td>
</tr>
<tr>
<td>EMA + Neem</td>
<td>2560</td>
<td>1961.8 e</td>
<td>64.6</td>
<td>59.7</td>
<td>0.77</td>
<td>4 c</td>
<td>3 b</td>
<td>282.6 e</td>
</tr>
<tr>
<td>EMC + Furadan</td>
<td>2220</td>
<td>1149.2 g</td>
<td>76.1</td>
<td>76.4</td>
<td>0.52</td>
<td>2 d</td>
<td>3 b</td>
<td>129.8 gh</td>
</tr>
<tr>
<td>EMA + Furadan</td>
<td>2190</td>
<td>1249.2 g</td>
<td>73.6</td>
<td>74.3</td>
<td>0.57</td>
<td>3 d</td>
<td>3 b</td>
<td>157.6 fg</td>
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<tr>
<td>Neem + Furadan</td>
<td>2430</td>
<td>1648.2 f</td>
<td>68.6</td>
<td>66.1</td>
<td>0.68</td>
<td>3 d</td>
<td>3 b</td>
<td>181.6 fg</td>
</tr>
<tr>
<td>EMC + Neem + Furadan</td>
<td>2320</td>
<td>805.6 h</td>
<td>83.9</td>
<td>83.4</td>
<td>0.35</td>
<td>2 e</td>
<td>1.2 d</td>
<td>29.8 i</td>
</tr>
<tr>
<td>EMA + Neem + Furadan</td>
<td>2310</td>
<td>965 h</td>
<td>80.7</td>
<td>80.2</td>
<td>0.42</td>
<td>2.8 d</td>
<td>2 c</td>
<td>71.8 hi</td>
</tr>
<tr>
<td>LB</td>
<td>2340</td>
<td>4635.4 b</td>
<td>8.4</td>
<td>4.7</td>
<td>1.98</td>
<td>4.6 b</td>
<td>4 a</td>
<td>1043 a</td>
</tr>
<tr>
<td>Check (Nematode only)</td>
<td>2250</td>
<td>4864.4 a</td>
<td>—</td>
<td>—</td>
<td>2.16</td>
<td>5 a</td>
<td>4 a</td>
<td>1070.4 a</td>
</tr>
</tbody>
</table>

*Each treatment was represented by five replicates (Plots), each with three plants. Numbers in each column followed by the same letter are not significantly different (P < 0.05 using Duncan's Multiple Range Test in COSTATE statistical program). *Final nematode population = Number of J.5/250 g soil + Number of nematode in root. *Reduction % = Total nematode population of check plants - Total nematode population of treated plants/ Total nematode population of check plants x 100.
significant difference between the nematode treatment and other treatments ($F = 39.7$) (Fig. 2D). Data presented in Table 2 showed that the tallest plants (60.2 cm) were observed from the combined application of EMC with neem cake and furadan with percentage of increase averaged to 90.5%, followed by EMA+ neem cake+ furadan treatment (82.9%), then dual applications of neem cake with EMC (79.7%) or EMA (71.5%). However, single application of EMC recorded 67.7% increase in plant length. The shortest seedling height was measured from the LB medium (negative control) treatment (32.4 cm, 2.5%) and furadan treatment (34 cm, 7.6%). Likewise, integrated treatment of EMC or EMA with neem cake plus furadan surpassed the other tested treatments in increasing percentages of increase in plant fresh and dry weights with values of 89% and 88.5% or 76.2% and 76.9%, respectively (Table 2). There was no statistically significant difference between EMC and EMA+ neem cake treatments in improving the previous plant growth measurements. For number of leaves, the results mirrored those from plant length and weight, maximum number of leaves was observed by the application of EMC (83.3%) or EMA (56.4%) with neem cake and furadan. However, number of leaves was minimum in case of single application of furadan (9%) and LB medium (5.1%) treatments.
Table 2. Influence of entomopathogenic bacteria alone or integrated with neem cake and/or furadan on growth parameters of Taify grapevine seedlings infested with *M. incognita* under micro-plot field conditions

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Plant Length (cm)</th>
<th>Increase%</th>
<th>Plant Fresh weight(g)</th>
<th>Increase%</th>
<th>Plant Dry weight(g)</th>
<th>Increase%</th>
<th>No. of Leaves</th>
<th>Increase%</th>
</tr>
</thead>
<tbody>
<tr>
<td>EMC</td>
<td>53 c</td>
<td>67.7</td>
<td>50.8 c</td>
<td>54.9</td>
<td>30.2 de</td>
<td>45.2</td>
<td>20 cd</td>
<td>28.2</td>
</tr>
<tr>
<td>EMA</td>
<td>45.6 de</td>
<td>44.3</td>
<td>43.8 de</td>
<td>33.5</td>
<td>27.8 fg</td>
<td>33.7</td>
<td>19.4 de</td>
<td>24.4</td>
</tr>
<tr>
<td>Neem</td>
<td>41 f</td>
<td>29.7</td>
<td>40.4 f</td>
<td>23.2</td>
<td>23.4 ij</td>
<td>12.5</td>
<td>17.2 fg</td>
<td>10.3</td>
</tr>
<tr>
<td>Furadan</td>
<td>34 g</td>
<td>7.6</td>
<td>37.4 g</td>
<td>14</td>
<td>22.4 jk</td>
<td>7.7</td>
<td>17 fg</td>
<td>9</td>
</tr>
<tr>
<td>EMC + Neem</td>
<td>56.8 a-c</td>
<td>79.7</td>
<td>55.6 b</td>
<td>69.5</td>
<td>32.8 c</td>
<td>57.7</td>
<td>23 b</td>
<td>47.4</td>
</tr>
<tr>
<td>EMA + Neem</td>
<td>54.2 bc</td>
<td>71.5</td>
<td>51.6 c</td>
<td>57.3</td>
<td>31.2 cd</td>
<td>50</td>
<td>21.2 c</td>
<td>36.4</td>
</tr>
<tr>
<td>EMC + Furadan</td>
<td>47.2 d</td>
<td>49.4</td>
<td>46.2 d</td>
<td>40.9</td>
<td>28.2 ef</td>
<td>35.6</td>
<td>19.6 de</td>
<td>25.6</td>
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<tr>
<td>EMA + Furadan</td>
<td>43.6 d-f</td>
<td>38</td>
<td>42.4 ef</td>
<td>29.3</td>
<td>26 gh</td>
<td>26</td>
<td>18.8 de</td>
<td>20.5</td>
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<td>Neem + Furadan</td>
<td>42 ef</td>
<td>32.9</td>
<td>41.8 ef</td>
<td>27.4</td>
<td>24.8 hi</td>
<td>19.2</td>
<td>18.2 ef</td>
<td>16.7</td>
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<td>EMC + Neem + Furadan</td>
<td>60.2 a</td>
<td>90.5</td>
<td>62 a</td>
<td>89</td>
<td>39.2 a</td>
<td>88.5</td>
<td>28.6 a</td>
<td>83.3</td>
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<tr>
<td>EMA + Neem + Furadan</td>
<td>57.8 ab</td>
<td>82.9</td>
<td>57.8 b</td>
<td>76.2</td>
<td>36.8 b</td>
<td>76.9</td>
<td>24.4 b</td>
<td>56.4</td>
</tr>
<tr>
<td>LB</td>
<td>32.4 g</td>
<td>2.5</td>
<td>33.6 h</td>
<td>2.4</td>
<td>22 jk</td>
<td>5.8</td>
<td>16.4 g</td>
<td>5.1</td>
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<tr>
<td>Check (Nematode only)</td>
<td>31.6 g</td>
<td>—</td>
<td>32.8 h</td>
<td>—</td>
<td>20.8 k</td>
<td>—</td>
<td>15.6 g</td>
<td>—</td>
</tr>
</tbody>
</table>

* Each treatment was represented by five replicates (Plots), each with three plants. Numbers in each column followed by the same letter are not significantly different (P < 0.05 using Duncan's Multiple Range Test in COSTATE statistical program). Increase % = Growth measurement of treated plants - Growth measurement of check plants / Growth measurement of check plants x 100.
**DISCUSSION**

The use of certain natural or synthetic materials that have already been involved in integrated management of such pathogenic nematodes is a desire trend. Of these materials, there are bacterial and plant products. Utilization of such compounds singly or mixing has earned much more benefits in excreting nematode development and diminishing grapevine damage under greenhouse or outdoor conditions. However, results of the present micro-plot field investigation initiate the novel phenomenon in suppressing root-knot nematode associated with improving grapevine growth parameters using entomopathogenic bacterial filtrate separately or mixing with either neem cake and/or furadan as a nematicide at the minimum rates to avoid environmental pollution. Apparently, results from the present experiment indicated that triple concomitant applications included bacterial filtrates gave the maximum reduction of nematode population, root galling and egg-mass formation as well increasing grapevine growth over the control and the other treatments. Meanwhile, dual applications of bacteria with furadan at half of recommended dose ranked second in reducing previous nematode parameters, although they not acted as a good growth promoters for grapevine seedlings. On the other hand, plants receiving bacterial filtrates then amended with neem cake improved plant growth and gave reasonable reductions in nematode criteria. Exposing grapevine roots to bacterial filtrates before treating with neem cake may possibly undergo physiological changes stimulating a certain degree of resistance in plants against nematode penetration and development. These results agree with those of, who mentioned that *Xenorhabdus* sp. filtrate suppressed *M. incognita* penetration into groundnut roots. Several researchers recorded that entomopathogenic symbiotic bacteria, *Xenorhabdus* and *Photorhabdus* are environmentally benign and produce some of the active compounds include xenorhabdins and xenocoumacins, bacteriocins, proteinaceous chitinases and non-protein indoles and stilbene derivatives. These metabolites have shown different bioactivities against pests and pathogens including nematodes. Recently, seven compounds were isolated from *X. budapestensis* SN84 and tested for their nematicidal properties against *J. s* of *M. incognita*. Among tested compounds, Rhabdopetide J, 2 showed strong inhibitory activity. The toxicity and repellency effects of cell-free bacterial suspensions of *Xenorhabdus* on the second-stage juveniles of *M. incognita* were almost entirely due to ammonium. The present results are also agreed with those of who reported that combined application of *Pasteuria penetrans* and neem extract maximized shoot length and weight of babchi plant and minimized number of juveniles per root system. Neem cake plus *Glomus fasciculatum* increased the plant growth of tomato and reduced *Meloidogyne incognita* reproduction and root-galling. The metabolites released during the decomposition of neem including azadirachtin, carotenoids, phenolic compounds, triterpenoids, salannin, limonoids and steroids and ketones stimulated and change the physiology of plant cells to release abnormal compounds which repel the nematodes from the uninfected cells and tissues of plant. In the present study, among single applications, furadan at full recommended dose was the uppermost treatment achieving the highest nematode suppressive rates, while a phytotoxic effect may occur since it gave the least values of growth characters. Here we investigated entomopathogenic bacterial filtrate as a possible sustainable adjuvant for use with neem cake and nematicide. Neem cake was possibly increase activity of antagonistic microorganisms by releasing mineral elements into soil, increasing osmotic potential of soil solutions and thereby nematode control was enhanced and plant growth was improved. Concomitant application of bacterial filtrate with neem cake plus furadan decreased rate of nematode build-up 6-fold, whereas, double application included furadan (4-fold) and neem cake (3-fold). Obviously, the present findings indicated that the bacterial filtrate applied either singly or integrated with neem cake and/or furadan at a half of recommended dose was the best applications in improving growth of grapevine and suppressing *M. incognita* development and reproduction in the naturally infested soil. Although several investigations recorded the nematicidal activity of entomopathogenic symbiotic bacterial filtrates singly in laboratory and greenhouse but this is the first report that EMC and EMA could fit well to the principles of integrated nematode management, thanks to their safety to environment, humans and animals and absence of nematode resistance. The results also support our hypothesis that bacterial filtrates can act additively or synergistically with other agricultural inputs in sustainable management programs of nematode.

**CONCLUSION**

It can be concluded from the present investigation that the use of EMC or EMA, neem cake and nematicide in integrations represents a promising novel approach for the integrated management of root-knot nematode infecting Taify grapevine and enhances the growth of plant.

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DATA AVAILABILITY
All datasets generated or analyzed during this study are included in the manuscript.

ETHICS STATEMENT
This article does not contain any studies with human participants or animals performed by any of the authors.

REFERENCES


