Uncovering Nematicidal Natural Products from *Xenorhabdus* Bacteria

Desalegne Abebew, Fatemeh S. Sayedain, Edna Bode, and Helge B. Bode*

**ABSTRACT:** Parasitic nematodes infect different species of animals and plants. Root-knot nematodes are members of the genus *Meloidogyne*, which is distributed worldwide and parasitizes numerous plants, including vegetables, fruits, and crops. To reduce the global burden of nematode infections, only a few chemical therapeutic classes are currently available. The majority of nematicides are prohibited due to their harmful effects on the environment and public health. This study was intended to identify new nematicidal natural products (NPs) from the bacterial genun *Xenorhabdus*, which exists in symbiosis with *Steinernema* nematodes. Cell-free culture supernatants of *Xenorhabdus* bacteria were used for nematicidal bioassay, and high mortality rates for *Caenorhabditis elegans* and *Meloidogyne javanica* were observed. Promoter exchange mutants of biosynthetic gene clusters encoding nonribosomal peptide synthetases (NRPS) or NRPS-polyketide synthase hybrids in *Xenorhabdus* bacteria carrying additionally an hfq deletion produce a single NP class, which have been tested for their bioactivity. Among the NPs tested, fabclavines, rhabdopeptides, and xenocoumacins were highly toxic to nematodes and resulted in mortalities of 95.3, 74.6, and 72.6% to *C. elegans* and 82.0, 90.0, and 85.3% to *M. javanica*, respectively. The findings of such nematicidal NPs can provide templates for uncovering effective and environmentally safe alternatives to commercially available nematicides.

**KEYWORDS:** entomopathogenic bacteria, *Xenorhabdus*, nematicidal natural products, cell-free culture supernatants, *Caenorhabditis elegans*, *Meloidogyne javanica*

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**INTRODUCTION**

Parasitic nematodes infect many species of plants and animals, including humans causing serious diseases that are deleterious to human health and agricultural productivity.\(^1\) Nematodes that parasitize plants are a global problem for agriculture.\(^2,3\) Plant parasitic nematodes represent a significant constraint on global food security. Worldwide, they account for a loss of important agricultural crops, estimated to be about multiple billions of dollars per year.\(^4\) *Meloidogyne incognita, M. javanica*, and *M. arenaria* are the most pathogenic *Meloidogyne* species. Root-knot nematodes are obligate plant parasites that exist in the roots of plants and interact with other plant pathogens to form disease complexes.\(^5\) Mostly root-knot nematodes affect development of host cells and gene expression and create giant cells, which affect absorption of water and nutrients from the soil.\(^6\)

Administration of anthelmintic drugs (e.g., albendazole; ivermectin) is the major means of controlling human and animal nematode infections. However, many anthelmintic drugs are losing their effectiveness because nematode strains with resistance are emerging.\(^7-9\) Different strategies, such as using chemical pesticides, organic fertilizers, resistant host plants, and biological control, have been reported to control root-knot nematodes.\(^7\) However, application of chemical pesticides against nematode pests (e.g., methyl bromide) has declined due to high concerns for environmental welfare and increased demands of organic agriculture.\(^10,11\)

Such problems stress the discovery of new and environmentally friendly nematicides. Soil bacteria can be a source of different biologically active compounds of economic and clinical importance.\(^12\) The genera *Xenorhabdus* and *Photorhabdus* are insect pathogenic bacteria, which exist in symbiosis with *Steinernema* and *Heterorhabditis* nematodes, respectively.\(^13\) After infection, these entomopathogenic bacteria produce a variety of bioactive compounds and hence kill the host insect larvae. These compounds protect the insect cadaver against food competitors, including bacteria and fungi.\(^14\)

Since other nematodes living in the soil are also food competitors, entomopathogenic bacteria are expected to be potential sources of lead molecules for nematicidal chemicals. Hence, we hypothesized that some species of *Xenorhabdus* produce active compound(s) in their NPs that can be toxic to plant parasitic nematodes. Production of specific NPs was achieved by generating Δhfq mutants of the desired strains that have a reduced background of NP production followed by activation of individual biosynthetic gene clusters via a promoter exchange mutant strategy.\(^16\) Accordingly, we tested cell-free culture supernatants of different strains of *Xenorhabdus* for their nematode toxicity. Preliminary work in our laboratory showed that cell-free supernatants of some entomopathogenic bacteria killed second stage juvenile (J2) of *M. javanica* and prevented the egg hatching of this nematode.\(^15\) In this study, the...
nematicidal phenotypes of Xenorhabdus bacteria were characterized, and potent nematicidal NPs were identified, which were exclusively produced through engineering of their corresponding nonribosomal peptide synthetases (NRPS) or NRPS-polyketide synthase (PKS) hybrids encoding biosynthetic gene clusters.

**MATERIALS AND METHODS**

**Culturing Bacterial Strains.** Xenorhabdus bacteria were isolated from their symbiotic nematode species (Table S1), which infected insect larvae, *Galleria mellonella*. Bacterial strains were stored in glycerol suspensions (50% v/v) at −80 °C and were cultured on Luria-Bertani (LB) agar plates (15 g/L agar). Xenorhabdus and *Escherichia coli* strains were grown overnight with shaking at 30 and 37 °C, respectively, in LB broth (10 g/L tryptone, 5 g/L yeast extract, and 5 g/L NaCl at pH 7.0). Cultures were subsequently inoculated (1:100 v/v) in fresh LB media and incubated with a rotary shaker for 2 days. The liquid culture was supplemented, when it is required, with 0.2% L-arabinose and antibiotics in appropriate concentrations (ampicillin 100 μg/mL; kanamycin 50 μg/mL).\(^{17,18}\)

**Construction of C. elegans.** Culturing of wild type (WT) Bristol N2 strain of *C. elegans* was performed under standard culturing conditions on a nematode growth medium (NGM) agar plate (3 g/L NaCl, 2.5 g/L peptone, and 1.2% agar). After autoclaving, the following ingredients were added as sterile filtered solutions: 1 mL of 1 M CaCl₂, 1 mL of 1 M MgSO₄, 25 mL of 1 M K₃PO₄, and 1 mL of cholesterol (5 mg/mL in EtOH). *C. elegans* is usually grown in the laboratory using *E. coli* OP50 strain as a food source. Overnight culture of *E. coli* OP50 (100 μL per plate) was spread on the NGM agar plate and grown at 37 °C for 12 h. *C. elegans* was transferred from one plate to another using a worm picker or a sterilized scalpel or spatula, and it was cultivated on the bacterial lawn for 3 days at 25 °C.\(^{19}\)

**Cultivation of *M. javanica*.** A culture of plant parasitic nematode *M. javanica* was maintained on the tomato plants in the greenhouse for 3 months. Tomato roots with egg masses were washed, and hand-picked egg sacs of *M. javanica* were placed on a nylon screen immersed in shallow water in glass Petri dishes, and hatched second stage juveniles (J2) were disinfected daily with streptomycin sulfate (0.1%) for 15 min with the vermiculite from Eurobiol. The general plasmids used in this work were purchased from New England Biolabs or Thermo Fisher Scientific. DNA primers were purchased from MWG Operon. The general plasmids used in this work were prepared in the LB liquid medium using appropriate antibiotics (kanamycin 50 μg/mL and ampicillin 100 μg/mL).\(^{18}\)

**DNA Techniques and DNA Manipulation.** Techniques for plasmid DNA preparation, restriction digestion, transformation, and DNA gel electrophoresis were adapted from standard protocols.\(^{17}\) Isolation of genomic DNA was carried out based on the manufacturer's instructions (QIAGEN). PCR amplifications were carried out on thermocyclers (SensoQuest). Restriction enzymes and DNA polymerases (Taq, Q5, and Phusion) were purchased from New England Biolabs or Thermo Fisher Scientific. DNA primers were purchased from MWG Operon. General plasmids used in this work are listed in Table S2. The PCR primers used in this study are shown in Table S4. All plasmids generated in this study (Table S4) were constructed using Hot Fusion Cloning.\(^{22}\)

**Construction of Deletion Mutants.** Deletion mutants in *X. duchattiae* DSM 17909 and *X. budapestensis* DSM 16342 were created using the primers listed in Table S4. The *hfp* gene of these two strains was deleted to abolish or reduce production of NPs.\(^{17,18}\) The *hfp* gene was deleted by amplifying about 1 kb fragments upstream and downstream of the respective genes. The amplified fragments were cloned into the *E. coli* expression assembly and then transformed into *E. coli* S17-1 λpir. Conjugation of the plasmid in *Xenorhabdus* and the generation of double crossover mutants via counter selection were done following established protocols.\(^{22}\) Verification of deletion clones was confirmed via PCR with the verification primers listed in Table S4.

**Generation of Promoter Exchange Mutants.** Promoter exchange mutants in *Xenorhabdus* were created using the primers listed in Table S4.\(^{17,18}\) These strains were generated for the production of specific NPs using the exchange of the natural promoter against an arabinose-inducible P₅₃₅₉ promoter. Promoter exchange mutants of *Xenorhabdus* were constructed following standard protocols.\(^{17}\) Briefly, the pCEP plasmid carrying the first 600–800 bp of a gene of interest of *Xenorhabdus* was constructed by Hot Fusion and transformed into *E. coli* S17-1 λpir. Cell suspensions of transformed *E. coli* were plated on selected LB agar plates containing kanamycin 50 μg/mL. Verification of positive clones was carried out via colony PCR using verification primers (Table S3). *Xenorhabdus* strains were conjugated with positive clones of *E. coli* S17-1 λpir harboring the respective promoter exchange plasmid as indicated previously. For both strains, overnight cultures were prepared in the LB liquid medium using appropriate antibiotics (kanamycin 50 μg/mL; ampicillin 100 μg/mL). The next day, both strains were grown in 5 mL of LB medium to an OD₆₀₀ of 0.6–0.8. The cells were harvested using 1 mL from each strain and washed using a fresh LB medium. The cells of *E. coli* S17-1 λpir were resuspended in 100 μL of LB, and *Xenorhabdus* bacteria were mixed on the LB agar plate in a ratio of 1:3 and incubated at 37 °C for 3 h and transferred to 30 °C until the next day. After 1 day, the cells were resuspended in 2 mL of LB for plating 100 μL of cell suspension on selective LB agar plates supplemented with kanamycin 50 μg/mL and ampicillin 100 μg/mL for further antibiotic resistance selection. The cells were incubated at 37 °C for 72 h. Screening of clones was carried out genetically by PCR using verification primers (Tables S3 and S4).

**Fermentation and Cell-Free Culture Supernatant Preparation.** *Xenorhabdus* bacteria listed in Table S1 were used to harvest cell-free culture supernatants for testing their nematicidal activity. They were cultured for 2 days at 200 rpm on a rotary shaker in LB broth at 30 °C. The cultures were cultivated in 1 L Erlenmeyer flasks containing 100 mL + 0.2% L-arabinose and inoculated with a 24 h pre-culture (0.1%, v/v). For the pre-culture, appropriate antibiotics were added to the LB medium when necessary at the following concentrations: kanamycin 50 μg/mL and ampicillin 100 μg/mL. The cell-free culture supernatants were prepared by centrifugation at 4000 rpm for 30 min in 50 mL Falcon tubes and filtered through a 0.2 μm filter. The supernatants were heat-treated at 90 °C for 10 min so that protein toxins denature and their effect can be separated from the NPs. Culture supernatants of *E. coli* OP50 and *Xenorhabdus* strains (WT and Δhfq) were used as controls. Additionally, freeze-dried supernatants of bacterial strains were prepared. Production of specific NPs from each promoter exchange strain was analyzed using HPLC-MS or MALDI-MS before being used for nematicidal bioactivity testing. For HPLC-MS analysis, strains were cultured in 5 mL of LB liquid medium with 0.2% L-arabinose and 2% Amberlite XAD-16 resin. After 72 h, XAD-16 beads were separated and extracted with 5 mL of MeOH for 1 h. The cell-free supernatant of the strains was used for MALDI-MS analysis.

**Nematicidal Activity Test against *C. elegans*.** Solid assay was adapted from the plate assay described by Tan et al.\(^{23}\) NGM agar plates (35 mm in diameter) were seeded with an overnight culture of bacteria (100 μL). The plates were incubated at 30 °C (*Xenorhabdus* strain) and 37 °C (*E. coli* OP50) for 48 and 24 h, respectively, to enable growth of the bacteria. L4 stages of *C. elegans* (up to 50 larvae per plate) were added onto each NGM. Lids of the plates were covered with parafilm. Incubation of plates was done at 25 °C, and death of the nematodes was analyzed every 24 h. Worms were considered dead after being unresponsive upon tapping the plate under a microscope.

The nematicidal activity against *C. elegans* was determined in a 24-well microtiter plate by a slightly modified method,\(^{24–26}\) where the cell-free culture supernatant of different bacterial strains was added for testing. Nematodes grown on the NGM seeded with an *E. coli* OP50 lawn of cells were washed from the plates with M9 buffer (3 g of KH₂PO₄, 6 g of Na₂HPO₄, 5 g of NaCl, and, after autoclaving, the addition of 1 mL of 1 M MgSO₄). Finally, a nematode suspension was filtered through a sieve with pores of 40 μm. In this assay, 80–100 L4 stage *C. elegans* were added in to a well of the 24-well microtiter plate containing 300 μL of cell-free culture supernatant from a bacterial strain to be tested. The plates were incubated at 25 °C in the dark, and the viability of the worms was recorded under a stereomicroscope at 40× magnification every 24 h for 3 days. The cell-free culture supernatant of *E. coli* OP50 was used as a negative control. The killing assay was conducted in triplicate. The nematodes were classified as dead when no movement was observed under a stereomicroscope and when their muscles were not visible. The number of dead nematodes was counted after each time interval.
bodies were straightened. Mortality of the nematodes was calculated as the ratio of dead nematodes compared to the total number of tested nematodes.

**Nematicidal Activity Test against M. javanica.** To evaluate the nematicidal effect on second stage juveniles (J2) of root-knot nematodes (M. javanica), 1.5 mL of sterile distilled water was added to each 15 mL Falcon tube containing the freeze-dried supernatant, resulting in a 10-fold concentration of the original supernatant. Then, the solutions were filtered through the 0.2 μm filter and were tested against J2 M. javanica. About 100 J2s of root-knot nematodes were added in each well of a 24-well plate containing 0.50 mL of fresh supernatant of bacterial strains (0.45 mL of supernatant +0.03 mL of nematode suspension +0.02 mL of streptomycin sulfate 0.1%). The number of dead J2 was recorded after 24 and 48 h with the stereomicroscope. Juveniles without movement were considered dead, and they were touched with a fine needle to confirm their death. The experiment was conducted based on a completely randomized design with three replications.20

**Microscopy.** Stereomicroscopy was used for counting live and dead worms during the nematode killing assay through a magnification of 40×. Detection of green fluorescent protein (GFP)-labeled Xenorhabdus in the gut of C. elegans was conducted using fluorescent microscopy. C. elegans was stained with Nile-Red stain as the control following an established protocol.27

**Statistical Analysis.** Analysis of variance (ANOVA) for all the obtained data was performed using the SAS (v. 9.1) software. Furthermore, the LSD test was employed for significant differences among treatments at P < 0.05.28

## RESULTS AND DISCUSSION

**Nematicidal Activity of Lawn of Cells of Xenorhabdus Bacteria against C. elegans.** C. elegans is a free-living nematode, which typically grows on NGM agar plates containing E. coli OP50, which represents the standard laboratory food for C. elegans.15,29 Based on this information, C. elegans was grown on the lawn of cells of X. budapestensis, X. szentirmaii, X. doucetiae, and X. nematophila. As a result, we observed that the lawn of cells of Xenorhabdus bacteria killed C. elegans while grazing on it (Figure 1A). This initiated us to identify the cause of death of this free-living nematode. To verify whether C. elegans grazes Xenorhabdus bacteria into its intestine, we used GFP-labeled Xenorhabdus bacteria as a lawn of cells. In this experiment, we noticed that the GFP-labeled Xenorhabdus bacteria caused infection and distributed throughout the entire length of the nematode intestine. A mass of cells of the bacteria caused engorgement at the pharynx, near the mouth of C. elegans (Figure 1B). Most of C. elegans were found dead within 2–3 days of the experiment. Hence, Xenorhabdus bacteria either affect normal physiological function of the intestine of the nematodes or produce nematicidal NPs in the gut of the nematodes to kill them within a short period of time. Other studies reported that C. elegans was antagonized through the colonization of the intestine by different human pathogens such as Salmonella typhimurium and Pseudomonas aeruginosa.30 Similarly, we showed that GFP-labeled X. nematophila bacteria disseminated through the intestine of C. elegans, which resulted in death of the worms over the span of a few days, although the precise mechanism of killing remained unknown.

**Nematicidal Activity of Cell-Free Culture Supernatants of Xenorhabdus Bacteria against C. elegans.** Most of Xenorhabdus bacteria showed strong nematicidal activity against the L4 stage of C. elegans during the microtiter plate nematicidal assay. Cell-free culture supernatants of WT of X. budapestensis, X. szentirmaii, and X. doucetiae grown in the LB medium resulted in mortalities of 91.0, 90.3, and 77.0% for C. elegans, respectively, after 48 h of the experiment (Figure S1; Table 1). HPLC-MS data analysis was conducted for the cell-free culture supernatant of nematicidal Xenorhabdus bacteria (e.g., X. budapestensis WT) and non-nematicidal E. coli OP50 (control). The profile of their base peak chromatograms (Figure S3) agrees with their nematicidal activities (Figure S2). Even if there was no significant variation among them (P > 0.05), cell-free supernatants of WT of X. budapestensis and X. szentirmaii had the greatest nematicidal effect followed by X. doucetiae (Table 1). Our findings agree with earlier results that have shown that cell-free culture supernatants of Xenorhabdus and Photorhabdus possess nematicidal activity against nematodes.15,32–37

**Characterization of Nematicidal Phenotypes of Xenorhabdus Bacteria.** We showed that cell-free culture supernatants of WT of Xenorhabdus bacteria killed C. elegans (Table 1, Figure 2A, and Figure S1), which could be due to either protein toxins or NPs. To differentiate the real cause of death for the nematode, cell-free culture supernatants were heated to inactivate protein toxins in cell-free culture supernatants of X. szentirmaii, X. budapestensis, X. nematophila, and X. doucetiae. All supernatants kept a high nematicidal activity after heat treatment at 90 °C for 10 min (Figure 3). This result indicated that the nematicidal compounds produced by these bacteria are heat stable as suggested for small molecule NPs and unlike toxic proteins. Even if toxic proteins also have a similar effect,38 our results demonstrated that Xenorhabdus are capable of fast killing C. elegans through production of NPs in their culture supernatants.

The gene encoding Hfq has been shown to influence virulence in some pathogens like Pseudomonas aeruginosa and Salmonella typhimurium and production of NPs in Photorhabdus and Xenorhabdus bacteria.39,40 After Δhfq mutant strains of X. szentirmaii, X. budapestensis, and X. doucetiae were generated, we compared cell-free culture supernatants of Δhfq strains with their corresponding WT for their nematicidal activity (Table 1). Cell-free culture supernatants of WT of Xenorhabdus bacteria

![Figure 1. Nematicidal effect of Xenorhabdus bacteria on the L4 stage of C. elegans. (A) Nematodes transferred onto a lawn of X. budapestensis on NGM were killed within 2–3 days. Pictures were taken using a stereomicroscope. The scale bars are 100 μm. (B) GFP-labeled X. nematophila bacteria (green) were grazed from the NGM agar plate by C. elegans and found disseminated through the gut of the nematode, and the bacteria caused engorgement at the pharynx of the nematode. The nematodes were stained with Nile-Red stain (red) as a contrast, and pictures were taken using fluorescence microscopy. The scale bar is 100 μm.](https://doi.org/10.1021/acs.jafc.1c05454)
show high nematode toxicity against *C. elegans*. In contrast, their corresponding Δhfq strains did not show such toxicity and differed significantly (*P* < 0.05) (Table 1). The absence of nematocidal properties compared to the WT is related to deletion of the *hfq* gene, which is a global regulator of gene expression (production of NPs) through sRNA/mRNA interactions.41

**Table 1. Nematicidal Activity of Cell-Free Culture Supernatants of Xenorhabdus Bacteria against the L4 Stage of *C. elegans*[^a^]**

| cell-free culture supernatant | mortality (%) ± SD |
|------------------------------|-------------------|
|                             | 24 h   | 48 h   | 72 h   |
| *X. budapestensis* WT        | 85.3 ± 2.6[^a^] | 91.0 ± 2.6[^a^] | 97.0 ± 1.0[^a^] |
| Δhfq                         | 5.3 ± 1.7[^a^]  | 12.6 ± 2.6[^a^] | 23.3 ± 2.3[^a^] |
| GameXPepptide (1)            | 13.0 ± 1.5[^a]  | 22.0 ± 4.7[^a]  | 34.6 ± 5.0[^a]  |
| Rhabdopeptide (2)            | 29.8 ± 3.3[^a]  | 52.8 ± 6.8[^a]  | 74.6 ± 2.0[^a]  |
| Fabclavine (3)               | 43.6 ± 3.4[^a]  | 75.3 ± 4.3[^a]  | 95.3 ± 1.4[^a]  |
| E. coli OP50, control        | 4.3 ± 1.2[^a]   | 6.3 ± 1.2[^a]   | 6.3 ± 1.2[^a]   |
| *X. szeitimair* WT           | 85.3 ± 2.6[^a]  | 90.3 ± 2.1[^a]  | 98.6 ± 1.3[^a]  |
| Δhfq                         | 8.6 ± 1.4[^a]   | 11.3 ± 1.2[^a]  | 18.6 ± 4.4[^a]  |
| Pyrrolizienamide (4)         | 10.0 ± 2.5[^a]  | 12.8 ± 2.6[^a]  | 25.3 ± 3.9[^a]  |
| Xenobactin (5)               | 12.0 ± 1.5[^a]  | 14.6 ± 0.3[^a]  | 29.3 ± 0.6[^a]  |
| Fabclavine (6)               | 61.6 ± 0.8[^a]  | 84.0 ± 4.5[^a]  | 93.3 ± 0.8[^a]  |
| E. coli OP50, control        | 4.3 ± 1.2[^a]   | 6.3 ± 1.2[^a]   | 6.3 ± 1.2[^a]   |
| *X. douceliae* WT            | 53.3 ± 24.9[^a] | 77.0 ± 1.5[^a]  | 91.3 ± 0.8[^a]  |
| Δhfq                         | 7.3 ± 1.3[^a]   | 11.3 ± 0.6[^a]  | 16.6 ± 0.8[^a]  |
| PAX-peptide (7)              | 10.0 ± 2.3[^a]  | 17.6 ± 1.7[^a]  | 30.0 ± 3.0[^a]  |
| Xenomocoumarin (8)           | 12.3 ± 2.3[^a]  | 45.0 ± 2.6[^a]  | 72.6 ± 2.4[^a]  |
| Xenorhabdin (9)              | 6.3 ± 1.2[^a]   | 23 ± 1.4[^a]    | 38.0 ± 3.0[^a]  |
| Phenylenlactamide (10)       | 3.0 ± 1.7[^a]   | 17.6 ± 2.0[^a]  | 31.3 ± 4.8[^a]  |
| Rhabduscin (11)              | 9.6 ± 1.7[^a]   | 47.3 ± 2.6[^a]  | 61.0 ± 3.0[^a]  |
| E. coli OP50, control        | 4.3 ± 1.2[^a]   | 6.3 ± 1.2[^a]   | 6.3 ± 1.2[^a]   |
| 0 – 10                       | (8) + LB       | 4.6 ± 0.6[^a]   | 13.0 ± 2.0[^a]  | 24.6 ± 2.6[^a]  |
| 11 – 20                      | (9) + LB       | 3.6 ± 0.3[^a]   | 9.3 ± 0.8[^a]   | 12.0 ± 1.5[^a]  |
| 21 – 30                      | (11) + LB      | 5.3 ± 1.4[^a]   | 9.6 ± 0.8[^a]   | 14.0 ± 1.7[^a]  |
| 31 – 40                      | (8) + (9)      | 20.3 ± 3.1[^a]  | 38.3 ± 4.9[^a]  | 45.0 ± 5.5[^a]  |
| 41 – 50                      | (8) + (11)     | 22.0 ± 1.7[^a]  | 38.6 ± 5.2[^a]  | 55.0 ± 1.7[^a]  |
| 61 – 70                      | (9) + (11)     | 29.6 ± 2.3[^a]  | 34.6 ± 2.7[^a]  | 38.0 ± 2.5[^a]  |
| 81 – 90                      | (9) + (11)     | 23.3 ± 0.3[^a]  | 3.6 ± 0.8[^a]   | 7.0 ± 0.5[^a]   |
| 91 – 100                     | LB, control    | 2.33 ± 0.3[^a]  | 3.6 ± 0.8[^a]   | 7.0 ± 0.5[^a]   |

[^a^] Mean values represent the mean of triplicates. Means in each column indicated by the same letter are not significantly different at *P* < 0.05 according to the LSD test. Comparison of the mean values is conducted for each strain separately using their mean values at each day of the experiment. Supernatants of WT, their corresponding Δhfq strains, and promoter exchange strains (Δhfq-pCEP-NP; induced with 0.2% arabinose) were used. Cell-free culture supernatants of *E. coli* OP50 and LB liquid media were used as the control. The experiments were conducted in triplicate, and mean values of the mortality are indicated here. Bioactivities are shown for none (white) to the highest activity (red).

Identification and Exclusive Production of Nematicidal Natural Products from *Xenorhabdus* Bacteria. We observed potent nematocidal activity from cell-free culture supernatants of WT of different *Xenorhabdus* bacteria. On the other hand, there was almost no nematocidal activity from their corresponding Δhfq mutants (Table 1). From these two different phenotypes of a strain, it is possible to suggest that the nematocidal NP is produced by *hfq* dependent NRPS and NRPS-PKS hybrid biosynthetic gene clusters (BGCs). We first generated *hfq* mutant strains, which did not produce any NP known from the WT strains. Next, exclusive production of an NP was achieved using the easyPAC[^18^] (easy Promoter Activation for Compound Identification) in which the native promoter was exchanged against an arabinose-inducible PBAD promoter in the *hfq* mutant strain (Figures 4A and S4–S6).

Accordingly, we generated 11 different easyPAC strains for which nematocidal bioactivity was analyzed using the cell-free culture supernatants against the L4 stage of *C. elegans*, including controls of LB, *E. coli* OP50, Δhfq, and the corresponding WT.
In X. budapestensis−Δhfq, GameXPeptides (1), rhabdopeptides (2), and fabclavines (3) were exclusively produced using activation of the PBAD promoter. Similarly, in X. szentirmaii−Δhfq, pyrrolizixenamide (4), xenobactin (5), and fabclavines (6) were individually produced. PAX-peptide (7), xenocoumacin (XCN2) (8), xenorhabdin (9), phenylethylamide (10), and rhabduscin (11) were also produced from X. dauceiae−Δhfq following the same procedure. The production status of these NPs was verified using HPLC-MS and/or MALDI-MS data analysis (Figure 4A; Figures S4–S6). We observed that activation of some BGCs could also result in production of an NP class having multiple derivatives (Table S5).

Out of 11 NPs that we tested, almost all resulted in more than 50% mortality of C. elegans. However, fabclavines (3; 6), rhabdopeptides (2), and xenocoumacin (8) appeared to be the most nematicidal NPs showing mortality greater than 70% with fabclavines (3) in X. budapestensis strain being the most toxic with 95.3% mortality against C. elegans (Table 1).

Previously, fabclavines were described as antibacterial and antifungal with broad spectrum properties, and recent studies confirmed that fabclavines have potent feeding-deterrent activity against deadly mosquito vectors. Fabclavines also displayed antibacterial efficacy against a multidrug-resistant E. faecalis. Additionally, it was demonstrated that zeamine, identified in Serratia plymuthica, and fabclavines show similarity in their structures containing a polyamine moiety. It was hypothesized that zeamine cytotoxicity involves disruption of the plasma membrane to facilitate solubilization of nematode cuticles. Hence, fabclavines might have a similar mechanism of action, but this is a subject for future in-depth investigation.

### Table 2. Nematicidal Effect of Xenorhabdus Bacteria against M. javanica (J2)

| Cell-free culture supernatant | Mortality (%) ± SD |
|------------------------------|-------------------|
|                              | 24 h              | 48 h              |
| X. szentirmaii (WT)          |                   |                   |
| X. szentirmaii (Δpttase)     | 94.0 ± 1.0⁣       | 93.3 ± 1.1⁣       |
| X. szentirmaii (Δhfq-ΔisinAB) | 80.3 ± 1.5⁣      | 0.0 ± 0.0⁣        |
| X. szentirmaii (Δhfq-ΔisinAB, pCEP-ianA, rhabduscin) + | 80.6 ± 2.0⁣      | 0.0 ± 0.0⁣        |
| X. szentirmaii (Δhfq-ΔisinAB, pCEP, fabclavine) + | 85.3 ± 4.0⁣      | 0.0 ± 0.0⁣        |
| X. xentirmaii (Δhfq-ΔisinAB-pCEP, fabclavine) + | 86.0 ± 5.5⁣      | 0.0 ± 0.0⁣        |
| X. xentirmaii (Δhfq-ΔisinAB-pCEP, rhabdopeptide) + | 86.6 ± 2.5⁣      | 0.0 ± 0.0⁣        |
| X. xentirmaii (Δhfq-ΔisinAB-pCEP, fabclavine) - | 93.0 ± 1.5⁣      | 90.0 ± 7.0⁣       |
| X. nematophila (WT)          |                   |                   |
| X. nematophila (Δhfq-ΔisinAB) | 89.0 ± 3.0⁣      | 86.6 ± 0.5⁣       |
| X. nematophila (Δhfq-ΔisinAB, pCEP, xenorhabdin) - | 71.6 ± 1.5⁣      | 0.0 ± 0.0⁣        |
| X. nematophila (Δhfq-ΔisinAB-pCEP, xenorhabdin) + | 80.0 ± 4.0⁣      | 0.0 ± 0.0⁣        |
| X. nematophila (Δhfq-ΔisinAB-pCEP, xenocoumacin) + | 80.6 ± 6.1⁣      | 0.0 ± 0.0⁣        |
| X. nematophila (Δhfq-ΔisinAB-pCEP, xenocoumacin) - | 86.6 ± 1.5⁣      | 0.0 ± 0.0⁣        |
| X. nematophila (Δhfq-ΔisinAB-pCEP, xenocoumacin) + | 92.6 ± 2.0⁣      | 85.3 ± 3.0⁣       |
| Control (water, LB)          | 0.0 ± 0.0⁣        | 0.0 ± 0.0⁣        |

*Mean values in each column indicated by the same letter are not significantly different at P < 0.05 according to the LSD test. + represents induced natural product in strain. - represents not induced natural product in strain. * represents mean of mortality in fresh supernatants. ‡ represents mean of mortality in freeze-dried supernatants. Fresh and freeze-dried cell-free supernatants of WT and their corresponding promoter exchange strains (Δhfq-pCEP-NP; induced with 0.2% arabinose) were used. Sterile water and LB liquid media were used as the control. The experiments were conducted in triplicate, and mean values of the mortality are indicated here. Bioactivities are shown for none (white) to the highest activity (red), see Table 1 for the color code.

Figure 3. Effect of heat inactivation of the cell-free culture supernatant on nematicidal activity. Heating of the cell-free culture supernatant does not affect the nematicidal activity of different WT of Xenorhabdus bacteria against C. elegans (L4). Supernatants were heated at 90 °C for 10 min to inactivate protein toxins. The survival rate of C. elegans was calculated after 48 h of the experiment. Heated cell-free culture supernatants have nearly the same nematicidal activity as the non-heated ones, this indicates the presence of nematode toxic NPs in heated cell-free culture supernatants. ** indicates the absence of significant difference (P > 0.05) between the nematicidal activity of heated and nonheated supernatants.
However, most of nematicidal drug classes impair the neuromuscular system of nematodes by interacting with ion channels and receptors on neurons and muscles.\textsuperscript{47} It was reported that rhabdopeptides contribute to insect killing acting as insect specific virulence factors\textsuperscript{48} and displayed positive effect against protozoal parasites.\textsuperscript{49} In our work, we observed nematicidal activity of rhabdopeptides against \textit{C. elegans}.

**Toxicity of Cell-Free Culture Supernatants of Xenorhabdus Bacteria against \textit{M. javanica}.** The most active

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**Figure 4.** Exclusive production of natural products using promoter exchange via the easyPACId approach. (A) HPLC-MS data analysis of WT and Δhfq mutants of \textit{X. budapestensis} and NPs detected after a promoter exchange in the Δhfq mutant. Extracted ion chromatograms (EICs) of derivatives of GameXPeptide (GXP) (1a, 1b, and 1c) and rhabdopeptide (Xbud_02752) (2a, 2b, and 2c) are indicated to show their exclusive production. (B) Structures of all NPs involved in the study.
nematicidal compounds that we identified were also tested for their activity against root-knot nematodes (Figure 2B). Although nematicidal activity was shown in all fresh supernatants of bacterial mutants, only freeze-dried supernatants of fabclavines (6), rhabdopeptides (2), and xenocoumaclin (8) resulted in 82.0, 90.0, and 85.3% mortalities of J2 of *M. javanica*, respectively, after 48 h (Table 2). A study reported that ammonium produced by *Xenorhabdus* causes nematicidal activity on J2 of *M. incognita*.34 Indole and 3,5-dihydroxy-4-isopropylstilbene (IPS), from the culture filtrate of *P. luminescens* MD, were shown to have nematicidal activity. IPS caused mortality of *C. elegans* but had no effect on J2 of *M. incognita*, while indole was lethal to *M. incognita*.50 Extracts of *P. luminescens* CH35 showed nematicidal activity on *M. incognita* but, however, had weak effect on *C. elegans*.50 Hence, entomopathogenic bacteria have great potential to produce different nematicidal NPs.

Three types of rhabdopeptides from *X. budapestensis* SN84 have been recently isolated that indicated nematicidal properties on *M. incognita*.51 Our results were in agreement with these results since we observed a similar nematicidal activity of cell-free culture supernatants of rhabdopeptides 2 producing strains. During detailed HPLC-MS analysis of an induced promoter exchange strain of *X. budapestensis*-Δhfq-Prad-Xbud-02752, we detected different derivatives of rhabdopeptides 2, which could be isolated and purified in the future to study their structure–activity relationship (Figure 4A).

In conclusion, *Xenorhabdus* bacteria produced a variety of nematicidal NPs. We identified the responsible NPs from the liquid culture of different strains of *Xenorhabdus* bacteria by applying the easyPACId approach of exclusive production of NPs, which was achieved through engineering of the corresponding NRPS and NRPS-PKS encoding BGCs. Herein, we enabled the strains to produce exclusively the active compounds. In addition, such microbially produced NPs are in principle degradable and ecofriendly for agricultural application. This makes them potentially useful for the biocontrol of nematodes in crop and vegetable production. We are still missing greenhouse data and toxicity data of the biocontrol of nematodes in crop and vegetable production.

In principle degradable and ecofriendly for agricultural applications, BGC, biosynthetic gene cluster; bp, base pairs; BPC, base peak chromatogram; DNA, deoxyribonucleic acid; EIC, extracted ion chromatogram; EtOH, ethanol; fcl, fabclavine; GFP, green fluorescent protein; Gxp, GameXPeptide; h, hour; Hfg, host factor of the RNA bacteriophage Qß; HPLC, high-performance liquid chromatography; IJ, infective juvenile; Kb, kilobase; LB, lysogeny broth; MALDI, matrix-assisted laser desorption/ionization; MeOH, methanol; min, minute; MS, mass spectrometry; m/z, ratio of mass to charge; NGM, nematode growth medium; NP, natural product; NRPS, nonribosomal peptide synthetase; OD<sub>600</sub>, optical density at a wavelength of 600 nm; Prad, l-arabinose-inducible promoter; pCEP, cluster expression plasmid; PCR, polymerase chain reaction; PKS, polyketide synthase; PPTase, phosphopantetheine transferase; Rdp, rhabdopeptide; WT, wild type; XCN, xenocoumaclin

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# ABBREVIATIONS

BGC, biosynthetic gene cluster; bp, base pairs; BPC, base peak chromatogram; DNA, deoxyribonucleic acid; EIC, extracted ion chromatogram; EtOH, ethanol; fcl, fabclavine; GFP, green fluorescent protein; Gxp, GameXPeptide; h, hour; Hfg, host factor of the RNA bacteriophage Qß; HPLC, high-performance liquid chromatography; IJ, infective juvenile; Kb, kilobase; LB, lysogeny broth; MALDI, matrix-assisted laser desorption/ionization; MeOH, methanol; min, minute; MS, mass spectrometry; m/z, ratio of mass to charge; NGM, nematode growth medium; NP, natural product; NRPS, nonribosomal peptide synthetase; OD<sub>600</sub>, optical density at a wavelength of 600 nm; Prad, l-arabinose-inducible promoter; pCEP, cluster expression plasmid; PCR, polymerase chain reaction; PKS, polyketide synthase; PPTase, phosphopantetheine transferase; Rdp, rhabdopeptide; WT, wild type; XCN, xenocoumaclin

# REFERENCES

(1) Jasmer, D. P.; Goversse, A.; Smant, G. Parasitic nematode interactions with mammals and plants. *Annu. Rev. Phytopathol.* 2003, 41, 245–270.

(2) Bulgheresi, S. All the microbiology nematodes can teach us. *FEMS Microbiol. Ecol.* 2016, 92, No. fix007.

(3) Chen, J.; Li, Q. X.; Song, B. Chemical Nematicides: Recent Research Progress and Outlook. *J. Agric. Food Chem.* 2020, 68, 12175–12188.

(4) Chitwood, D. J. Research on plant-parasitic nematode biology conducted by the United States Department of Agriculture-Agricultural Research Service. *Pest Manage. Sci.* 2003, 59, 748–753.
In Manual of Techniques in Insect Pathology, a novel source for chaetoglobosin A, is a destructive parasite of the cereal crop. Z. Zhou, Q.; Kaiser, M.; Klemmt, P.; Bode, H. B. Simple and very effective entomopathogenic, nematophilic bacteria of the family Enterobacteriaceae. Entomopathogenic Nematodes: A Review of Their Use in Crop Protection. J. Agron. Crop Sci. 2006, 192, 269–300.

Entomopathogenic nematodes show strong nematicidal activities against the root-knot nematode, Meloidogyne javanica. Journal of Agricultural and Food Chemistry pubs.acs.org/JAFC

(25) Aballay, A.; Yonge, P.; Ausubel, F. M. Salmonella typhimurium proliferates and establishes a persistent infection in the intestine of the nematode, Caenorhabditis elegans. Curr. Opin. Chem. Biol. 2019, 45, 671–680.

(26) Bode, E.; Heinrich, A. K.; Hirschmann, M.; Abebew, D.; Shi, Y.-H.; Touray, M.; Gulsen, S. H.; Erincik, O.; Wenski, S. L.; Rutter, J. W.; Ozdemir, T.; Galimov, E. R.; Quintaneiro, L. M.; Rosa, L.; Thomas, G. M.; Cabreiro, F.; Barnes, C. P. Detecting Changes in the Caenorhabditis elegans Intestinal Environment Using an Engineered Bacterial Biosensor. ACS Synth. Biol. 2019, 8, 2620–2628.

(27) Mishra, S.; DiGennaro, P. Root-knot nematodes demonstrate temporal variation in host penetration. J. Nematol. 2020, 52, 1–8.

(28) Mishra, S.; DiGennaro, P. Root-knot nematodes demonstrate temporal variation in host penetration. J. Nematol. 2020, 52, 1–8.

(29) Stock, S.; Simonyi, S.; Keller, N.; Eckstein, S.; Frank, D.; Fuhrmann, J. E. U.; Matilla, M. A.; Salmond, G. P. C. The broad-spectrum antibiotic, zeamine, kills the nematode worm, X. szentirmaii, as well as inverted repeats without ligase. Angew. Chem., Int. Ed. 2019, 58, 1537–1541.

(30) Aballay, A.; Yonge, P.; Ausubel, F. M. Salmonella typhimurium proliferates and establishes a persistent infection in the intestine of the nematode, Caenorhabditis elegans. Curr. Opin. Chem. Biol. 2019, 45, 671–680.

(31) Kaya, H. K.; Patricia, Stock, S. Techniques in insect nematology. In Manual of Techniques in Insect Pathology, 2nd ed.; Eliever 2009; pp 281–324.

(32) Wadhwa, T. A.; Bode, H. B. Promoter Activation in Engineered Bacterial Biosensor. Proc. Natl. Acad. Sci. U. S. A. 2003, 100, 2760–2765.

(33) Wang, G.; Huang, X.; Li, S.; Huang, J.; Wei, X.; Li, Y.; Xu, Y. The A-D, o-Pyrone Bearing a 4-O-Methyl-β-d-glucopyranoside Moiety from the Spider-Associated Ascomycete Akanthomyces novoguineensis. Molecules 2017, 22, 1202.

(34) Cimen, H.; Touray, M.; Gulsen, S. H.; Erincik, O.; Wenski, S. L.; Bode, H. B.; Shapiro-Ilan, D.; Hazir, S. Antifungal activity of the nematode, X. szentirmaii, on different fungal phytopathogens and identification of the antifungal compounds from X. szentirmaii. Appl. Microbiol. Biotechnol. 2021, 105, 5517–5528.
(44) Kajla, M. K.; Barrett-Wilt, G. A.; Paskewitz, S. M. Bacteria: A novel source for potent mosquito feeding-deterrents. Sci. Adv. 2019, 5, 6141.

(45) Donmez Ozkan, H.; Cimen, H.; Ulug, D.; Wenski, S.; Yigit Ozer, S.; Telli, M.; Aydin, N.; Bode, H. B.; Hazir, S. Nematode-Associated Bacteria: Production of Antimicrobial Agent as a Presumptive Nominee for Curing Endodontic Infections Caused by Enterococcus faecalis. Front. Microbiol. 2019, 10, 2672.

(46) Wu, J.; Zhang, H.-B.; Xu, J.-L.; Cox, R. J.; Simpson, T. J.; Zhang, L.-H. 13C labeling reveals multiple amination reactions in the biosynthesis of a novel polyketide polyamine antibiotic zeamine from Dickeya zeae. Chem. Commun. 2010, 46, 333–335.

(47) Wolstenholme, A. J. Ion channels and receptor as targets for the control of parasitic nematodes. Int. J. Parasitol.: Drugs Drug Resist. 2011, 1, 2–13.

(48) Reimer, D.; Cowles, K. N.; Proschak, A.; Nollmann, F. I.; Dowling, A. J.; Kaiser, M.; ffrench-Constant, R.; Goodrich-Blair, H.; Bode, H. B. Rhabdopeptides as insect-specific virulence factors from entomopathogenic bacteria. ChemBioChem 2013, 14, 1991–1997.

(49) Zhao, L.; Cai, X.; Kaiser, M.; Bode, H. B. Methionine-Containing Rhabdopeptide/Xenortide-like Peptides from Heterologous Expression of the Biosynthetic Gene Cluster kj12ABC in Escherichia coli. J. Nat. Prod. 2018, 81, 2292–2295.

(50) Orozco, R. A.; Molnár, I.; Bode, H.; Stock, S. P. Bioprospecting for secondary metabolites in the entomopathogenic bacterium Photorhabdus luminescens subsp. sonorensis. J. Invertebr. Pathol. 2016, 141, 45–52.

(51) Bi, Y.; Gao, C.; Yu, Z. Rhabdopeptides from Xenorhabdus budapestensis SN84 and Their Nematicidal Activities against Meloidogyne incognita. J. Agric. Food Chem. 2018, 66, 3833–3839.