Biocontrol Potential of Four Indigenous Entomopathogenic Nematodes From Kenya

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Abstract
Globally, vegetable production including tomato is constrained by pests and diseases. The tomato leafminer, *Tuta absoluta* (Meyrick.), is a major pest of tomato, mainly managed using chemical pesticides. There is a need for integrated pest management (IPM), approaches that are human and environmentally friendly. The study aimed at molecular identification and virulence evaluation of four entomopathogenic nematodes (EPNs) against *T. absoluta* larvae. The DNA was extracted, sequenced, and phylogeny analyzed. The EPNs virulence was evaluated using the insect baiting technique at different nematode concentrations (0-Control, 100, 150, 200, and 250 infective juveniles/ml of distilled water). Larval mortality data were collected for five days. Molecular identification resulted in nucleotide sequence lengths of 877-895 base pairs (bp). All the isolates were found to be *Steinernema* species of EPNs. The % similarity of the isolates and their close relatives ranged between 82-100%. The isolates closely matched *Steinernema* spp. AY230184 (Sri Lanka); AY230186 (Kenya), JN651414 (Ethiopia), and MW151701 (Kenya). The sequences were deposited to Genbank as *Steinernema* sp. Isolate Kalro 75 (MW150871), *Steinernema* sp. Isolate Kalro S86 (MW150872), *Steinernema* sp. isolate Kalro97 (MW150873) and *Steinernema* sp. Isolate KalroR52 (MW150874). All the EPN isolates were virulent against *T. absoluta*. Isolate Kalro R52 and Kalro S86 recorded the highest mortality of 76±7.5%; 72±15.0% respectively in the 48 h. There was a significant difference (P < 0.001), between all the isolates and the Control. The four EPNs were virulent and have bio-control potential against, *T. absoluta*. Research on their virulence against *T. absoluta* in the field is recommended.

Keywords: pesticides, IPM, entomopathogenic nematodes, *Steinernema* sp., virulence, bio-control

1.0 Introduction
The use of chemical pesticides in agriculture has increased over the past decade globally. Annually an estimated 2 million tonnes of pesticides are used worldwide (Sharma et al., 2019; Choudhury & Saha, 2020). Globally, tomato production is constrained by pests and diseases. Among major tomato pests is the tomato leafminer *Tuta absoluta* Meyrick., which is mainly managed using chemical pesticides that are harmful to humans and the environment (Terzidis et al., 2014; Ochilo et al., 2019). Continuous release of new pesticides formulations, lure farmers into applying them hoping for better results (Blair et al., 2014; Singh et al., 2017; Tri et al., 2017; Sharma et al., 2019; WHO & FAO, 2019). However, food, health, and environmental safety concerns have lead to a shift to integrated pest management (IPM) strategies. Such strategies are the deployment of biological pest control agents, including entomopathogenic nematodes (EPNs) (Sanda & Sunusi, 2016; Belien, 2018; Singh et al., 2017; Perez-Alvarez et al., 2019; Surendra, 2019).

Soil inhibiting EPNs in the family Steinernematidae and Heterorhabditidae, are known to provide effective management of insect pests. They kill insect pests in 24-72 hours aided by symbiotic bacteria associated with them. They have gained importance due to safety on animals, do not affect non-target organisms, safe to the environment, ease of mass production, application, a commercial formulation of foliar applications, and exemption from registration rigors (Gozel & Kasap, 2015; Abate et al., 2020; Yuksel & Canhilal, 2019; Saleh et al., 2020). The EPNs as bio-control agents have been successful in the control of mainly soil-borne pests, though their use against above-ground pests has also been reported. Insect pests like leafminers, form mines/galleries on plants as they feed.
The mines/galleries favor nematode infective juveniles (IJs), survival, and protection from adverse environmental conditions (U. Gozel & C. Gozel, 2016; Lacey & Georgis, 2012; Ndereyimana et al., 2019). The search for indigenous EPN strains and species coupled with virulence evaluation on agricultural pests is crucial in the use of EPN as pest bio-control agents (Batista et al., 2014; Acharya et al., 2020; Ngugi et al., 2020). This study aimed at the molecular identification and virulence evaluation of four indigenous EPNs on *Tuta absoluta* (Mayrick) in Kenya.

2. Materials and Methods

2.1 Entomopathogenic Nematode Isolation

The four entomopathogenic nematodes (EPNs) isolates used in this study were Laboratory cultures maintained at Kenya Agricultural and Livestock Research Organisation-Horticulture Research Institute (HRI)-Thika. The isolates had been isolated from soils in different parts of Kenya in 1997 and 2006. The isolate Karlo S86 was from Coast, R52 from Central highlands while Karlo75 and 97 were from the Rift valley (Waturu et al., 1998; Mwaniki, 2008).

Soil samples (top-soils 0-30 cm deep) from each location were put in well-labeled sterile plastic bags and stored at 4-7 °C in the EPN laboratory at KALRO Mwea. The isolates were later transferred to KALRO-HRI-Thika. The nematodes were multiplied and maintained according to methods described by Bedding and Akhurst (1975); White (1927), using the Greater wax moth (*Galleria mellonella*) pre pupa larvae.

2.2 Insect Cultures

The *Galleria mellonella* and *Tuta absoluta* cultures were established from the insect life stages obtained from beekeepers and tomato farmers respectively. The *G. mellonella* were raised in the laboratory on an artificial diet; 45 g bee wax, 95 g brewer’s yeast, 307 g maize flour, and 225 g honey (Waturu, 1998). The culture of *T. absoluta* was maintained on tomato crop in a screen house at KALRO-HRI-Thika.

2.3 Entomopathogenic Nematodes DNA extraction, PCR, and Sequencing

Fresh female EPNs of each isolate were extracted from a five-day-old infected *G. mellonella* cadavers. The cadavers were dissected in Ringer’s solution and female nematodes were picked. The DNA of the EPN isolates was extracted according to Caoili et al. (2017) protocol. The resultant DNA was quantified and purity was assessed through spectrophotometry. The Polymerase Chain Reaction (PCR) was conducted according to Hominick et al. (1997). The TW81-F (5′-GTTTCCAGTAGGTAACCTGC-3′) and AB28-R (5′-ATATGCTTAAAGTTTGCAGG-3′) primers were used for the ITS rDNA amplification. The conditions for the PCR were 35 cycles per minute at 94 °C for 5 min, 94 °C for 1 min, 55 °C for 1 min, 72 °C for 2 min, and 72 °C for 5 min. Electrophoresis was conducted on 1.5% agarose gel at 160 volts for 1hr. The DNA was sequenced by Macrogen Inc. in Korea.

2.4 Sequence and Phylogenetic Analysis

The nucleotide sequence editing was done according to Hall, (1999), using Bio Edit v 7.0.5, and aligned using the MUSCLE alignment tool (Edgar, 2004; Madeira et al., 2019). The sequences were subjected to a similarity search from the NCBI database using the Basic Local Alignment Search Tool (BLASTn) (https://blast.ncbi.nlm.nih.gov). Phylogenetic (Neighbour-Joining, Distance method) tree was reconstructed from 12 best BLASTn results using SeaView version 4 bioinformatic program (Edgar, 2004; Gouy et al., 2010). The Coronavirus-2 (MW732681), was used as the out-group taxa. The bootstrap test was set at 1000 replications to determine branch support (Nguyen & Hunt, 2007).

2.5 Virulence of the Entomopathogenic Nematode Isolates

The *Galleria mellonella* pre-pupa larvae were infected with the four EPN isolates. The odorless and spongy *G. mellonella* cadavers were randomly selected, and infective juveniles (IJ), obtained according to White, (1927), procedure. The IJs were used in the evaluation of virulence of the four indigenous EPN isolates against *Tuta absoluta*. Nematode doses (0, 100, 150, 200, and 250IJs/) in ml of distilled water were used in the experiment. The experiment comprised five treatments in five replicates for each EPN isolate all in a completely randomized design (CRD). In the Control (0), nematodes IJs excluded in the one (1) ml of distilled water used. Data on insect mortality was collected daily for 5 days.

2.6 Data Analysis

The larval mortality data were subjected to analysis of variance (ANOVA) using Statistical software, Genstat 15th edition. Means were separated using Fisher’s protected least significant difference test at a 5% significance level. Results were presented graphically and in tables.
3. Results

3.1 Sequence and Phylogenetic Analysis

The lengths of the rDNA nucleotide sequences obtained were between 877-895 bp as shown in Table 1. The base composition on average among the isolates was Adenine (A) 29%; Cytosine (C) 23%; Guanine (G) 21.7% and Thymine (T) 26.1%. Individual EPN isolate base composition is summarised in Table 1.

Table 1. Entomopathogenic nematode sequence length

| Nematode isolates | Sequence length | % Nucleotide bases |
|-------------------|----------------|-------------------|
|                   |                | A     | C     | G     | T     |
| Isolate Karlo75   | 877bp          | 25    | 20.8  | 26.5  | 27.8  |
| Isolate KarloS86  | 881bp          | 30.8  | 25.2  | 17.8  | 26.2  |
| Isolate KarloR52  | 815bp          | 32.4  | 25.3  | 17.8  | 24.5  |
| Isolate Karlo97   | 895bp          | 28.3  | 21.5  | 24.6  | 25.7  |

Note. A = Adenine, C = Cytosine, G = Guanine, T = Thymine.

Phylogeny study of the four isolates alone resulted in a tree with two major clades each comprising of two closely related EPN isolates. The isolate Kalro75 clustered with Kalro97 while Kalro52 and S86 grouped (Figure 1).

The multiple sequence alignment of ITS rDNA of the EPN isolates compared with GenBank accessions revealed that the isolates belonged to the EPN genus *Steinernema* species. The phylogenetic tree reconstructed resulted in two major clades of *Steinernema* spp. All the four EPN isolates were in the same clade where they clustered with Genbank accessions, *Steinernema* spp. AY230184 (Sri Lanka); AY230186 (Kenya); JN651414 (Ethiopia) and MW151701 (Kenya) (Spiridonov et al., 2004; Tamiru et al., 2013; Ngugi et al., 2021). The % similarity of the isolates and their close relatives ranged between 82-100%. Isolate Kalro75 was closely related to a Kenyan *Steinernema* sp. isolate Kalro (MW151701) with a similarity of 100%. The isolate KalroS86 was closely related to accession *Steinernema* sp. AY230186 from Kenya. The Genbank accession MW732681 (Corona virus-2) was the most divergent species (Figure 2).
Figure 2. Phylogenetic relationships (Neighbour-Joining-method) based on ITS of rDNA sequences of the four nematode isolates (Kalro75, Kalro97, KalroR52, and KalroS86) with 12 Steinernema spp. from Genbank. Numbers on branches are bootstrap values for clades supported above the 70% level.

3.2 Virulence of Entomopathogenic Nematode Isolates

All the nematode isolates were virulent on larvae of *Tuta absoluta* causing death two days (48 h) after infection. No larval mortality was observed in the Control experiment. Isolate KalroR52 was the most virulent at nematode suspension of 150 IJs/ml of distilled water, with % mean larval mortality of 76±7.5. The least virulent isolate was Kalro97 with a mean mortality % of 36±4.0 (Figure 3). In descending order of virulence, the isolates were KalroR52 (76±7.5%), KalroS86 (72±15.0%), Kalro75 (64±7.5%), and Kalro97 (36±4.0%). A significant difference (P < 0.001) between all the four EPN isolates and the Control (Zero nematodes) was observed. In addition, there was a significant difference (P < 0.001) between isolates KalroR52, Kalro75, KalroS86, and Kalro97 (Figure 3).
4. Discussion

Results from molecular identification of the isolates confirmed the four EPNs were *Steinernema* species clustered into two categories of two isolates each (isolate Kalro75 and Kalro97; KalroR52 and S86). This could have resulted from differences in the genetic makeup of each isolate. This was supported by sequence analysis where the nucleotide base pair percentages were very close among the pair of isolates in each cluster. According to Stock (2002) and Noujeim et al. (2016), EPNs exhibit differences within and among species. The variations in species and strains are an important consideration in the use of EPN as a biological pest control agent (Shapiro-Ilan, 2012; Gozel, 2016). The generated phylogenetic tree with isolates sequences and those retrieved from Genbank confirmed relatedness with already described EPN species. There were two major clades within the study EPN isolate clustering with Accessions, *Steinernema* spp. MW130872 (Sri Lanka); MW130873 (Kenya); JN651414 (Ethiopia), and MW151701 (Kenya). This suggested close evolutionary and geographical relatedness among the four isolates with the already identified *Steinernema* spp. It has been reported that climate influences the distribution of EPNs globally. The geographical distribution of EPNs is a biological resource necessary in the utilization of native species for biological pest and disease control (Hominick, 2002; Tarasco, 2014).

All the Four EPN isolates were virulent against *Tuta absoluta* larvae while no death was observed in the Control experiment. Virulence of 95 and 100%, at 150 IJs/ml was reported by Khanum and Javed (2020), from EPNs. *Steinernema pakistanense* NNRC-AS.04 and *S. bifurcatum* NNRC-As.65 respectively against termites (*Coptotermes heimi* Wasmann). The nematode concentration of 150 IJs/ml resulted in insect mortality of > 60% among KalroR52, S86, and 75. This pest mortality was higher than mortality caused by some synthetic pesticides like Malathion with 51.7% mortality 72 h after application against Fall Armyworm (FAW) was reported by Sisay et al. (2019). According to Batalla-Carrera et al. (2010), Jacobson and Martin (2011), and Ngugi et al. (2021), the EPNs *Steinernema carpocapsae*, *S. feltiae*, *Steinernema* spp.-Kalro, and *Heterorhabditis bacteriophora* are lethal to *T. absoluta* larvae. For EPNs to qualify for bio-control, they should be virulent against the target pest (Shapiro-Ilan, 2015; Shehata et al., 2019).

The level of virulence, by insect mortality, differed among the isolates at the same nematode concentration. The isolates were from different parts of Kenya, with Kalro52 that was from Central highlands that are known for tomato production being the most virulent to *T. absoluta*. It’s suggested that probably climatic conditions of these areas influenced the virulence of the EPN isolates against *T. absoluta*. In addition, EPN species host-finding characteristic varies among the *Steinernema* species or strains (Poinar & Grewal, 2012; Nguyen et al., 2018).

5. Conclusions and Recommendations

The molecular analysis of ITS rDNA indicated that the four study isolates were *Steinernema* species of EPNs. The sequences are deposited as Genbank Accessions *Steinernema* sp. isolate Kalro75 (MW150871), *Steinernema* sp. isolate Kalro S86 (MW150872), *Steinernema* sp. isolate Kalro97 (MW150873) and *Steinernema* sp. isolate KalroR52 (MW150874). The four EPN isolates were virulent against *T. absoluta* larvae. Screen-house and field trials to evaluate the virulence of the isolates are recommended towards the development of EPN based bio-control against tomato leafminer, *Tuta absoluta* (Meyrick.).

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