Occurrence of plant-parasitic nematodes on enset (*Ensete ventricosum*) in Ethiopia with focus on *Pratylenchus goodeyi* as a key species of the crop

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Received: 24 July 2020; revised: 6 September 2020
Accepted for publication: 7 September 2020

**Summary** – Enset (*Ensete ventricosum*) is an important starch staple crop, cultivated primarily in south and southwestern Ethiopia. Enset is the main crop of a sustainable indigenous African system that ensures food security in a country that is food deficient. Related to the banana family, enset is similarly affected by plant-parasitic nematodes. Plant-parasitic nematodes impose a huge constraint on agriculture. The distribution, population density and incidence of plant-parasitic nematodes of enset was determined during August 2018. A total of 308 fields were sampled from major enset-growing zones of Ethiopia. Eleven plant-parasitic nematode taxa were identified, with *Pratylenchus* (lesion nematode) being the most prominent genus present with a prominence value of 1460. It was present in each sample, with a highest mean population density per growing zone of 16 050 (10 g root)^−1, although densities as high as 25 000 were observed in fields at higher altitudes in Guraghe (2200-3000 m a.s.l.). This lesion nematode is found in abundance in the cooler mountainous regions. Visible damage on the roots and corms was manifested as dark purple lesions. Using a combination of morphometric and molecular data, all populations were identified as *P. goodeyi* and similar to populations from Kenya, Uganda and Spain (Tenerife). Differences in population densities amongst cultivars indicate possible resistance of enset to *P. goodeyi*.

**Keywords** – altitude, food security, lesion nematode, molecular data, morphology, prominence value.

*Ensete ventricosum*, commonly known as enset, is a large perennial herbaceous plant belonging to the Musaceae family, together with banana and plantain. Unlike banana and plantain, however, enset does not produce bunches but instead produces a large underground corm that is harvested. The pseudostem is formed from a bundle of leaf sheaths and large leaves, which may reach up to 10 m high and 2 m diam. (Westphal & Stevels, 1975). Wild enset species are found distributed over sub-Saharan Africa and Asia, but in Ethiopia, where it has been domesticated, it is cultivated as an important food crop grown on approximately 400 000 ha (CSA-Ethiopia, 2016). As a key starch staple food source, enset provides food security for over 20 million people, or at least 20% of the Ethiopian population. Furthermore, it is also used for animal feed, fibre, construction and medicine (Brandt et al., 1997). The crop grows best at cooler, higher altitudes and is found mostly between 1200-3100 m a.s.l., in the south and southwestern areas of the country. Enset-based farming systems represent a traditional and sustainable form of agriculture, which includes a diverse range of crops that are cultivated alongside enset (Cheesman, 1947; Sim-
monds, 1962; Brandt et al., 1997). Enset is a perennial crop that takes, on average, 7 years to mature; however, as with most crops, the period to maturity is likely to be lengthened under challenge from biotic and abiotic threats. Identifying the biotic threats that challenge enset, and consequently addressing them, has received limited attention.

A number of constraints challenge enset production, with bacterial wilt disease caused by *Xanthomonas vasicola* pv. *musacearum* (Xvm) (previously named *X. campestris* pv. *musacearum* (Xcm)) (Studholme et al., 2020) receiving most attention (Addis et al., 2004, 2008; Nakato et al., 2018). The enset root mealy bug (*Cataenococcus ensete*) can cause severe damage to the roots and corm, reducing crop vigour and production (Addis et al., 2010). Also, fungal diseases such as a *Sclerotium* sp. root and corm rot, and *Acremonium* inflorescence spot, causing necrosis on flower bracts and leaves, can affect production, although they appear not to be widespread (Tesera & Quimio, 1994; Quimio & Tesera, 1996). A newly reported leaf streak disease, caused by a new *Badnavirus* species, has also recently been identified (Abraham et al., 2018; Abraham, 2019). Plant-parasitic nematodes, well known as major production constraints to banana and plantain production (Sikora et al., 2018), have received only limited attention on enset (Coyne & Kidane, 2018; Coyne et al., 2018). A few studies have associated various nematode species with the crop, with the lesion nematode *Pratylenchus goodeyi* appearing to be the most prevalent (Peregrine & Bridge, 1992; Tesera & Quimio, 1994; Speijer & Fogain, 1998; Mandefro & Dagne, 2000; Swart et al., 2000; Bogale et al., 2004). The root-knot nematodes *Meloidogyne incognita*, *M. javanica* and *M. ethiopica*, and *Aplolechoides ensete* have also been reported as potential production constraints (Mandefro & Dagne, 2000; Swart et al., 2000).

Compared to other pathogens, nematodes are, in general, poorly recognised in sub-Saharan Africa (Coyne et al., 2018) and Ethiopia in particular (Abebe et al., 2015). Despite a handful of studies associating nematode species with enset, there has been no concerted effort to establish the pest potential of nematodes on enset. The current study serves to provide a basis for more focused studies towards understanding the pest potential of nematodes on the crop. A comprehensive sampling of nematodes was undertaken in southern Ethiopia to establish the current situation regarding nematode incidence across the region, in relation to commonly cultivated cultivars and the influence of altitude (temperature) on their occurrence, with emphasis on the most prevalent nematode genus, *Pratylenchus*. This study also served to identify ‘hot spots’ where material could be collected for use in trials.

**Materials and methods**

**Survey area**

Enset root and soil samples were collected from the southern part of Ethiopia, from administrative zones where enset is most commonly grown (Sidama, Hadiya, Kembata and Kefa) in August 2018. Based on the Ethiopian administrative structure a total of 308 fields were sampled; 72 fields were selected randomly from each of the four zones and an additional ten fields each from Guraghe and Jimma zones (Fig. 1). In some fields where multiple cultivars were present, samples were collected separately from different cultivars. For each sample, the location, geographical coordinates, altitude and enset cultivar were recorded. Enset thrives best in slightly acidic, well-drained and fertile soils (Brandt et al., 1997). Specific soil characteristics were not assessed for each site; however, we have observed that in each farm enset was grown in soils rich in organic matter. Root and soil samples were removed using a spade by excavating a hole ca 0.5 m distance from the stem, from 3-4 plants of each cultivar per field and placed in plastic bags, labelled and stored in a cooler box for transport to the laboratory. Additional *P. goodeyi* populations were collected from Kenya and Uganda, and others supplied from Canary Islands (courtesy of Javier López-Cepero), which were included in the molecular assessment for comparison with Ethiopian populations.

**Processing of samples**

Soil and root samples were processed separately. Enset roots were carefully washed, cut longitudinally and chopped into ca 0.5 mm-sized pieces and a 10 g sub-sample was used for nematode extraction. For soil samples, a 100 ml sub-sample was extracted after fully mixing the soil for each sample. Nematodes were extracted from both soil and roots using a modified Baermann method over a period of 48 h (Hooper et al., 2005). Nematode suspensions were decanted and nematodes collected on a 38 μm sieve, rinsed into beakers, reduced to 10 ml and densities counted from 1 ml aliquots under a compound microscope. Nematode densities were calculated for each root and soil sample and expressed as the number of ne-
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MORPHOLOGICAL CHARACTERISATION

Nematodes fixed in TAF were processed to anhydrous glycerin for permanent slides, following a modified glycerin-ethanol method (De Grisse, 1969). Morphological features were observed and measurements made using a Leica DM 6000 B compound microscope equipped with Leica Application Suite (LAS) version 4.6.1 fitted with an Andor iXon 885 EMCCD camera. *Pratylenchus* specimens were identified to species level based on available keys (Sher & Allen, 1953; Castillo & Vovlas, 2007).

MOLECULAR CHARACTERISATION

The ethanol-preserved nematodes were washed three times in 400 μl of sterile water for 10 min. DNA extraction was done by cutting an individual specimen and transferring the two pieces to an Eppendorf tube containing 20 μl of WLB (50 mM KCl; 10 mM Tris pH 8.3; 2.5 mM MgCl₂; 0.45% NP-40 (Merck Life Sciences); 0.45% Tween-20). The samples were frozen for at least 10 min; 1 μl proteinase K (1.2 mg ml⁻¹) was added and the samples were incubated for 1 h at 65°C and 10 min at 95°C. Finally, the samples were centrifuged for 1 min at max speed 20 (800 g). They were stored at −20°C before running the PCR. The supernatant was taken as a template for PCR reaction; 2 μl was transferred to an Eppendorf tube containing 25 μl master mix (Derycke et al., 2010) and PCR amplification was performed using a Bio-Rad T100™ thermocycler. For the D2-D3 expansion segment of the large sub-unit (LSU) rDNA primers MalF...
The cytochrome c oxidase subunit 1 (COI) gene fragment was amplified using the forward primer JB3 (5′-GGATAGAGCCCRACGTATCTG-3′) (Wiśniewska & Kowalewska, 2015) and 1006R (5′-GGTTCGATTAGTCTTTCGCACCCCT-3′) (Holtermann et al., 2008) were used. The PCR amplification conditions were: initial denaturation of 5 min at 94°C, 35 cycles of (94°C for 1 min; 52°C for 90 s; 68°C for 2 min), and a final extension of 10 min at 68°C. The cytochrome c oxidase subunit 1 (COI) gene fragment was amplified using the forward primer JB3 (5′-TTTTTTGGGCATCCTGAAGTCTAT-3′) (Derrycke et al., 2010) and the reverse primer JB4prat (5′-CCTATTCTCTAAACATAATGAAAATG-3′) adapted from Bowles et al. (1992) with an initial denaturation of 5 min at 94°C, 40 cycles of (94°C for 1 min; 48°C for 30 s; 72°C for 100 s), and a final extension of 10 min at 72°C. The PCR products were separated by electrophoresis on agarose gel stained with GelRed (Biotium) and visualised under UV light. Sanger sequencing of purified PCR fragments was carried out in forward and reverse direction by Macrogen (Europe). Contigs were assembled using Geneious 2019.0.4 (Biomatters; http://www.geneious.com). All contigs were subjected to BLAST searches to check for possible contaminations. The resulting sequences were analysed with other relevant sequences available in GenBank. Multiple alignments of the different DNA sequences were made using MUSCLE with default parameters and followed by manual trimming of the poorly aligned ends using Geneious 2019.0.4. Phylogenetic trees were created by using MrBayes 3.2.6 add-in of Geneious with the GTR + I + G model. The Markov chains for generating phylogenetic trees were set at 1 × 10⁶ generations, four runs, 20% burn-in and sub-sampling frequency of 500 generations (Huelsenbeck & Ronquist, 2001).

Statistical treatment of data

Nematode population densities were calculated for each genus and/or species per field. Nematode count data were log(x + 1) transformed before analysis to minimise variation and conform data to normal distribution (Zuur et al., 2010). The percentage frequency of occurrence was calculated as (FO = (number of sites where a genus was detected/total number of sites) × 100), and prominence values (PV = population density × frequency of occurrence/10) (De Waele & Jordaan, 1988) were also calculated for each nematode genus and/or species (identified from both soil and root samples) across the sampled fields. PV is an indication of the relationship of population density and frequency. The association between nematode density and enset cultivar and the association between nematode density and altitude was analysed using RStudio® and Pearson’s correlation analysis. Using the GIS coordinates for each farm sampled, distribution maps were created for the key nematode species *P. goodeyi*.

**Results**

A total of 308 enset field samples were collected from six administrative zones. Eleven plant-parasitic nematode genera were identified: *Pratylenchus*, *Meloidogyne*, *Helicotylenchus*, *Scutellonema*, *Tylenchorhynchus*, *Rotylenchulus*, *Aphelenchoides*, *Cephalenchus*, *Pratylenchoides*, *Trophurus* and *Hoplolaimus* (Table 1). The genera *Pratylenchus*, *Meloidogyne* and *Aphelenchoides* were recovered from roots (Table 2). With regard to frequency of occurrence (FO%), *Pratylenchus* and *Helicotylenchus* species were present in 100 and 52% of the soil samples, respectively, followed by *Tylenchorhynchus* (16%), *Scutellonema* (10%) and *Meloidogyne* (13%). *Pratylenchus*, *Meloidogyne* and *Aphelenchoides* species occurred in 100, 8 and 4% of root samples, respectively (Table 2). *Pratylenchus* was the most prominent nematode tax across the enset-growing areas with a prominence value of 1460, followed by *Meloidogyne* and *Aphelenchoides* with PVs of 20 and 4, respectively. *Pratylenchus* spp. densities were highest in the highlands of Gurage, where mean densities of 16 050 and 12 217 (10 g root)⁻¹ were observed in Meskan and Ezha woredas/disticts, respectively, although densities as high as 25 000 (10 g root)⁻¹ were recorded from individual fields. The elevation of these areas ranged between 2200 and 3000 m a.s.l. (Fig. 2). Roots from these locations appeared dry and, when split longitudinally, extensive black or purple necrotic cortical tissue was evident (Fig. 3A), which was also observed on the corms (Fig. 3B). Roots from locations infected with *Meloidogyne* presented visible galling damage (Fig. 3C) but no obvious damage was associated with *Aphelenchoides* species.

This study found *P. goodeyi* to be present in every farm sampled and thus widely distributed, but alongside a range of species associated with enset and the cooler climate at the highest altitudes (>2200 m a.s.l.). There was a positive correlation (r = 0.11, P = 0.08) between altitude and population densities of *P. goodeyi* (Fig. 4). The correlation of *P. goodeyi* root density with cultivar showed that densities varied from 20 (‘Bedo’) to 4600 (‘Birdo’) (10 g root)⁻¹, but no statistically significant differences in
### Table 1. Frequency of occurrence, population density and prominence value of major plant-parasitic nematode genera recovered from enset soil in Ethiopia.

| Zone     | District* and elevation (m a.s.l.) | Pratylenchus | Meloidogyne | Helicotylenchus | Scutellonema | Tylenchorhynchus | Rotylenchulus | Aphelenchoides | Cephalodorus | Pratylenchoides | Pratylenchoides | Trophurus | Hoplolaimus |
|----------|------------------------------------|--------------|-------------|----------------|--------------|----------------|---------------|----------------|-------------|----------------|----------------|-----------|-------------|
| Sidama   | Dale 1700-1800                     | +            | +           | +              | +            | +              | +             | +              | +           | +              | +              | +         |            |
|          | Arbegona > 2600                    | +            | +           | +              | +            |                | +             |                | +           | +              | +              | +         |            |
|          | Hula > 2600                        | +            | +           | +              | +            |                |               |                | +           | +              | +              | +         |            |
| Hadiya   | Misha 2300-2600                    | +            | +           | +              | +            | +              | +             |                | +           | +              | +              | +         |            |
|          | Lemo 2300-2600                     | +            | +           | +              | +            | +              | +             |                | +           | +              | +              | +         |            |
|          | Duna 2300-2600                     | +            | +           | +              | +            |                |               |                | +           | +              | +              | +         |            |
| Kembata  | Angacha 2000-2500                  | +            | +           | +              | +            | +              |               |                | +           | +              | +              | +         |            |
|          | Doyo Gena 2200-2700                | +            | +           | +              | +            |                |               |                | +           | +              | +              | +         |            |
|          | Kedida Gamella 2000-2200           | +            | +           | +              | +            |                |               |                | +           | +              | +              | +         |            |
| Keffa    | Chenna 1700-2100                   | +            | +           | +              | +            |                |               |                | +           | +              | +              | +         |            |
|          | Decha 1700-2100                    | +            | +           | +              | +            |                |               |                | +           | +              | +              | +         |            |
|          | Gimbo 1700-2100                    | +            | +           | +              | +            |                |               |                | +           | +              | +              | +         |            |
| FO %     | 100                                 | 13           | 52          | 10             | 16           | 9              | 1             | 3              | 3           | 5              | 5              | 10        |            |
| PD       | 84                                  | 26           | 45           | 34             | 21           | 35             | 10            | 17             | 110         | 47             | 58             |           |            |
| PV       | 84                                  | 9            | 32           | 11             | 8            | 11             | 1             | 3              | 19          | 11             | 13             |           |            |

* Each district has 24 sampled fields.

**FO %** = Frequency of occurrence (FO %), i.e., number of fields where a genus is detected/total number of fields sampled \( \times 100 \).

**Population density (PD)** = Mean number of individuals of a genus over the sampled fields where the genus was detected; densities per 100 ml soil.

**Prominence value (PV)** = Mean population density \( \times \) (Frequency of occurrence)\(^{1/2} \) \( \times 10^{-1} \).

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Table 2. Frequency of occurrence, population density and prominence value of major plant-parasitic nematode taxa recovered from enset roots in Ethiopia.

| Zone   | District* and elevation (m a.s.l.) | Pratylenchus | Meloidogyne | Aphelenchoides |
|--------|-----------------------------------|--------------|-------------|----------------|
| Sidama | Dale 1700-1800                     | +            | +           |                |
|        | Arbegona >2600                     | +            | +           |                |
|        | Hula >2600                         | +            | +           |                |
| Hadiya | Misha 2300-2600                    | +            | +           | +              |
|        | Lemo 2300-2600                     | +            | +           |                |
|        | Duna 2300-2600                     | +            | +           |                |
| Kembata| Angacha 2000-2500                   | +            | +           |                |
|        | Doyo Gena 2200-2700                | +            | +           |                |
|        | Kedida Gamella 2000-2200           | +            | +           |                |
| Keffa  | Chenna 1700-2100                    | +            | +           |                |
|        | Decha 1700-2100                     | +            | +           |                |
|        | Gimbo 1700-2100                     | +            | +           |                |

| FO (%) | 100 | 8  | 4  |
| PD     | 1460| 69 | 22 |
| PV     | 1460| 20 | 4  |

* Each district has 24 sampled fields.

**FO %** = Frequency of occurrence (FO %), i.e., number of fields where a genus is detected/total number of fields sampled × 100.

Population density (PD) = Mean number of individuals of a genus over the sampled fields where the genus was detected; densities per 100 ml soil.

Prominence value (PV) = Mean population density × (Frequency of occurrence)$^{1/2}$ × 10$^{-1}$.

Fig. 2. *Pratylenchus goodeyi* densities per 10 g root across enset-growing zones of Ethiopia.

densities amongst the cultivars were observed. The number of samples for each cultivar also differed, reflecting farmer and/or geographical preferences for different cultivars.

*Pratylenchus* was the most frequently occurring nematode genera in soil samples with mean soil density of 84 nematodes (100 ml soil)$^{-1}$ followed by *Helicotylenchus* (45) and *Scutellonema* (34). The genera *Pratylenchus*
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A

B

C

Fig. 3. A: Longitudinal section of enset roots showing lesions caused by the lesion nematode; B: Purple lesions caused by the lesion nematode on enset corm; C: Galling on enset roots caused by root-knot nematodes, *Meloidogyne* spp.

*Morphological and molecular characterisation of Pratylenchus*

Using a combination of morphometric and molecular data with *Pratylenchus* specimens, *P. goodeyi* was the only species of the genus identified. Morphologically, the Ethiopian populations displayed typical diagnostic characteristics of *P. goodeyi*, including four lip annuli, four inconspicuous lateral field lines, stylet 16-18 μm long with pronounced, anteriorly flattened stylet knobs, large spermatheca filled with sperm, tail conoid, ventrally concave with dorsal contour sinuate just prior to tail tip, which matched the characterisation described by Castillo & Vovlas (2007). Seven *Pratylenchus* specimens (five females and two males) were measured: female; L = 0.56 mm; a = 32.88; b = 4.12; c = 17.93; V = 73.42; stylet = 16.44 μm; male: L = 0.55 mm; a = 29.48; b = 4.36; c = 23.1; T = 55.52; stylet = 16.1 μm.

*Pratylenchus goodeyi* populations from Ethiopia, Kenya, Uganda and the Canary Islands were used for molecular analysis. Eighty-one D2-D3 of 28S rDNA (GenBank accession numbers of selected sequences: MT569985, MT569991-94) and 101 mtDNA COI sequences with a maximum intraspecific variability of, respectively, 3 (0.5%) and 16 (4.1%) nucleotides were obtained. The D2-D3 phylogenetic tree (based on 652 bp long alignment with 116 sequences) revealed that all obtained sequences are in a maximally supported clade together with virtually identical *P. goodeyi* sequences from GenBank (0-3 bp difference), but without internal resolution (Fig. 5). For COI, sequences were obtained with premature stop codons that appeared difficult to align with other *Pratylenchus COI* sequences. This indicates that the used primers appeared not to have targeted the genuine COI region, but mitochondrial fragments into the nuclear genome (nuclear mitochondrial pseudogenes). Subsequent attempts with several other primers (Folmer et al., 1994; Kanzaki & Futai, 2002; Derycke et al., 2010) did not alleviate this pseudogene problem, i.e., always the same pseudogene was obtained. Nevertheless, the resulting phylogenetic tree (based on 360 bp long alignment with 102 sequences) clearly clustered all our sequences with a very similar (0-16 bp different) COI reference of *P. goodeyi* (unpublished sequence in the framework of the study of Janssen et al. (2017a)), which confirms the species identity. However, the internal resolution in this *P. goodeyi* clade was limited and without relation to host or location, impeding statements related to intraspecific relations (separated analyses of the pseudogene sequences). Despite the evidence of nuclear pseudogenes that com-

*choides, Hoplolaimus* and *Trophurus*, which occurred in fewer sites, had densities of 110, 58 and 47 nematodes (100 ml soil$^{-1}$) (Table 1).
complicated this study, it was evident that all investigated *Pratylenchus* species are unmistakably *P. goodeyi*, which is confirmed by morphology, D2-D3 sequences and a putative species-specific pseudogene (Fig. 5).

**Discussion**

The present study shows that although a range of plant-parasitic nematode species are associated with enset in the major producing zones in south and south western Ethiopia, *P. goodeyi* dominates strongly and is the most prominent species. Our study also represents the most extensive assessment of nematodes on enset to date, and the latest since Addis et al. (2006) with 98 farms sampled in 2004 and Bogale et al. (2004) who assessed 49 farms in 1999. The predominance of *P. goodeyi* in the previous studies and in our study identifies this nematode as probably the greatest nematode threat to enset. Root-knot nematodes (*Meloidogyne* spp.) were recovered from a few root samples, with relatively lower PV scores and densities; galling damage was observed on enset roots in the current study, which indicates it is becoming more problematic as this is the first time this appears to have been observed. Previously *Meloidogyne* spp. was found in 9% of 98 enset samples (Addis et al., 2006) and 60% in a smaller study (Bogale et al., 2004), which shows some variability in the recovery of these nematodes between studies. The current comprehensive study therefore demonstrates the incidence of *Meloidogyne* spp. associated with the crop across the region and supports the growing concern of this pest becoming more serious on crops across sub-Saharan Africa (Coyne et al., 2018). *Aphelenchoides* spp. were isolated from the roots of enset, reflecting previous studies, although no discoloured leaves were observed, which has previously been associated with *A. ensete* infection (Swart et al., 2000; Bogale et al., 2004; Addis et al., 2006). Although *Aphelenchoides* spp. have been associated with damage to enset, this does not appear to be prominent (PV = 4). No *Helicotylenchus multicinctus* were recorded from the roots, even though this nematode is common on banana in Ethiopia and was recorded from 5% of enset roots by Addis et al. (2006). Neither was any *Radopholus similis* recorded on enset, in line with previous studies (Bogale et al., 2004; Addis et al., 2006), even though it was present on banana in the previous studies (Addis et al., 2006). Enset therefore, may not be a suitable host for *H. multicinctus* or *R. similis*, unlike banana. However, environmental factors may not be suitable for *R. similis*, which is known to be thermophilic and present at warmer, lower altitudes than enset is generally cultivated (Kashaija et al., 1994).

In line with previous studies (Bogale et al., 2004; Addis et al., 2006), *Meloidogyne*, *Heliocotylenchus*, *Scutellonema*, *Tylenchorhynchus* and *Rotylenchulus* were among the plant-parasitic nematodes associated with enset soil samples. Our study also detected species of *Cephalenchus*, *Pratylenchoides*, *Trophurus* and *Hoplo-
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Fig. 5. Bayesian 50% majority-rule consensus tree inferred from Pratylenchus goodeyi COI and D2-D3 (pseudogene) sequences, obtained from enset in southern Ethiopia, and from banana in Kenya and Tenerife.
from enset soil samples, but all were in relatively low densities and do not appear to be causing any major threat to the crop. The incidence and distribution of *P. goodeyi* was prominent in higher altitudes (2000-3000 m a.s.l.), such as the extremely high densities observed in some farms from the Guraghe mountains, with over 20 000 (10 g root)$^{-1}$ recorded and even up to 26 000 (10 g root)$^{-1}$. This by far exceeds the density of *P. goodeyi* (5484 (10 g root)$^{-1}$) previously recorded by Bogale *et al.* (2004) from enset rhizome tissue and the 15 000 recorded by Peregrine & Bridge (1992). A maximum of 5600 nematodes (10 g root)$^{-1}$ was recovered from banana in Cameroon (Bridge *et al.*, 1995) and mean densities of 2500 nematodes from East African Highland banana roots in Uganda (Kashaija *et al.*, 1994). Enset, therefore, appears to be able to tolerate high densities of *P. goodeyi*. There are no reports on how this affects crop growth and production; therefore, it remains speculative as to the level of damage being caused to enset. A case study on banana in Rwanda showed that the highest *P. goodeyi* densities and root necrosis were present in the best performing banana plants, a possible explanation being that the negative impact of the nematode was masked by the fact that the plants were receiving better nutrient inputs (Gaidashova *et al.*, 2009). However, it is assumed that at such high densities as observed during our study, substantial damage is being caused. Roots with high *P. goodeyi* densities were associated with root necrosis and purple lesions, while the outer cortex of corms at times presented severe necrotic lesioning, especially on planting materials (Fig. 3b). When visiting farms, substantial portions of the corm with lesions and rotten areas were observed being removed during the preparation of corm material for food processing, resulting in much reduced corm size and food quantity. The wide range in *P. goodeyi* densities could indicate possible variations in the biology or pathogenicity of geographic populations. Populations of *P. goodeyi* from elsewhere within Africa were also shown to be similar to the Ethiopian populations, indicating a relatively recent distribution of the species within Africa (Bridge *et al.*, 1997). Difference in pathogenicity between geographic populations or ‘pathotypes’ has been speculated, given the contrary evidence of damage observed by *P. goodeyi* on bananas and the uniformity of *P. goodeyi* populations (Speijer & Bosch, 1996; Coyne, 2007). Populations occurring in Tanzania appeared similar to those from other countries (Mgonja *et al.*, 2019), even though some of these populations were recovered from tropical lowland areas, which is atypical for the species. Similarly, populations of *P. goodeyi* are being recovered from other tropical lowland locations (Coyne, 2007; Sikora *et al.*, 2018). As yet, there is no conclusive evidence to demonstrate differences between populations. In the current study, both morphometric and molecular techniques were used to identify the *Pratylenchus* populations. In general, morphological identification of *Pratylenchus* species is difficult due to the low number of diagnostic features, high morphological plasticity and incomplete taxonomic descriptions (Castillo & Vovlas, 2007; Janssen *et al.*, 2017a). DNA-based identification tools are therefore important for *Pratylenchus* species (Waeyenberge *et al.*, 2000), but also a strong link between morphology and DNA sequences is of crucial importance to prevent sequence-based misidentifications (Janssen *et al.*, 2017b). However, the morphological characterisation of the Ethiopian *P. goodeyi* all corresponded closely to the documented characteristics (Sher & Allen, 1953). *Pratylenchus goodeyi* is also one of the few *Pratylenchus* species that can be relatively easily identified based on morphology alone. The molecular assessment of *P. goodeyi* populations, based on the D2-D3 and COI region, and including specimens from countries other than Ethiopia, did not reveal informative differences. As expected, intraspecific resolution of the D2-D3 region is limited for *Pratylenchus* (Janssen *et al.*, 2017a). For COI a higher resolution can be expected; however, sequences that are most likely nuclear pseudogenes have complicated our analyses. Pseudogenes have been detected in several eu-karyotes and impede the usefulness and dependability of DNA (Leite, 2012). Nonetheless, for nematode taxonomic and phylogenetic studies, pseudogenes are not well recognised as being problematic. Furthermore, the COI region of several *Pratylenchus* has been sequenced (Janssen *et al.*, 2017a, b), but the pseudogene problem only appears to be present in *P. goodeyi*, in all globally distributed populations. How specifically *P. goodeyi* differs in this respect remains to be investigated. Although there was no significant difference in *P. goodeyi* population densities among the cultivars, variations in levels of infection across cultivars from the current study show that possible differences in resistance exist in enset against *P. goodeyi*. The assessment of 111 cultivars using Random Amplified Polymorphic DNA (RAPD), demonstrated that each cultivar had unique DNA (Birmeta *et al.*, 2002). However, given the difference in infection levels between geographic and altitudinal locations, this needs proper assessment through controlled inoculation studies. Differences in nematode densities between 71 en-
set cultivars, sampled from 98 farms, showed possible differences in resistance to *P. goodeyi* (Bogale et al., 2004) but this again requires verification.

During our survey, we perceived that very few farmers were aware of, or had any knowledge of, nematode pests. To some extent, they were aware of the bacterial wilt problem on enset and other foliar diseases but not of nematodes. As in the case of many smallholder farmers and agricultural agents even in sub-Saharan Africa, there remains a huge gap in the awareness of nematodes as pests and their management, even though nematodes are regarded as economically important pests of most crops in the region (Coyne *et al*., 2018). With such a high frequency of occurrence of *P. goodeyi* on enset in Ethiopia, and with such high densities recorded, it is assumed that this nematode is causing damage to crop growth and production. With a lack of information on the damage potential of this nematode to enset, this survey will provide a basis for identifying hotspots for nematode material for use in assessing the efficacy of the nematode on enset, potential on-farm assessment and interaction of nematodes with other organisms.

**Acknowledgements**

The authors thank the Norwegian Agency for Development Cooperation (Norad) for funding this study which is part of the project “Controlling disease in sweet potato and enset in South Sudan and Ethiopia to improve productivity and livelihoods under changing climatic conditions using modern technologies” under the NORHED program (Agreement No. ETH-13/0017).

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