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Short Note

The need for standardisation: exemplified by a description of the diversity, community structure and ecological indices of soil nematodes.

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Nematodes; Standardisation

1 Abstract

2 Molecular approaches are offering a supplement to, or even the possibility of replacing 3 morphological identification of soil fauna, because of advantages for throughput, coverage and 4 objectivity. We determined ecological indices of nematode community data from four sets of 5 duplicate soil cores, based on morphological identification of nematodes after elutriation from 200g 6 soil and high throughput sequencing (HTS) targeting nematodes both after being elutriated from 7 soils and DNA extracted directly from 10g soil. HTS (at genus and species level) increased the f 8 taxonomic resolution compared to morphology (at family level). DNA extracted from elutriated 9 nematodes identified more nematode taxa than when extracted from soil, due to an enrichment in 10 nematode sequences. Each method also gave a different ecological footprint for the nematode 11 community. Standardisation to previously determined indices based on morphological identification 12 is needed in order to provide more meaningful information about soil quality and for ecological 13 monitoring.

14

15 **1. Introduction**

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17 The study of soil and aquatic micro- and meso-fauna is being transformed by the use of molecular 18 methods (Creer et al., 2010). Not only are the developing molecular methods complementing and 19 even superseding the traditional morphological approaches, they are also developing faster than 20 standard protocols. Philippot et al (2012) highlighted the fact that methodological differences 21 between laboratories, of even the same protocol, are not trivial and hamper comparisons between 22 studies. They urged soil biologists to expand the list of standardised protocols listed by the 23 International Organisation for Standardisation (ISO). This was taken a little further by Römbke et al 24 (2016) who pointed out that when biodiversity data, for example, are being used in a legal context 25 they have to be comparable and lack of standardisation can limit the justification of specific 26 protection measures.

27 Nematodes are important indicators for soil monitoring (Chen et al., 2010) and there is a large 28 body of existing information based on morphological identification, which has led to well established 29 ecological indices based on nematode traits (Ferris et al., 2001). Morphological identification, 30 though, is often only to the family or trophic group (Porazinska et al., 2009) leaving ecological 31 analyses potentially ambiguous or superficial (Yeates and Bongers, 1999). The level of 32 characterisation of the nematode community is also problematical for DNA based methods, as 33 reliable sequence annotation relies on having curated sequences from vouchered specimens which 34 are not always available. There is a fundamental choice to extract DNA directly from soil or to firstly 35 elutriate nematodes and then extract DNA from those nematodes (here 'elutriation' covers 36 nematode extraction from soil, and 'extraction' refers to DNA). Advantages and disadvantages can 37 be argued for either approach. Elutriating nematodes before extracting DNA will enrich nematodes 38 and diminish other fauna, but takes longer and not all nematodes might be elutriated equally 39 efficiently (Persmark et al., 1992). Directly extracting DNA circumvents issues associated with 40 elutriation and saves time, but relatively small amounts of soil are usually extracted (i.e. <10g rather 41 than the >200g recommended as optimal by Wiesel et al., 2015). 42 It is important to be able to relate molecular results to the previous body of work using 43 morphological identification, and to have a good understanding of the limitations inherent with each 44 method (Porazinska et al., 2010; Stone et al., 2016; Quist et al., 2016). Currently only the extraction 45 and morphological identification of soil nematodes is covered by an ISO standard (ISO 23611-4). 46 Given the growing interest in biological soil monitoring (Aalders et al., 2009; Turbé et al., 2010; 47 Pulleman et al 2012, Faber et al 2013; Tsiafouli et al 2015; Griffiths et al, 2016), we considered that a 48 reminder of the importance of standardisation for the introduction of the developing molecular 49 methods was timely and relevant. We undertook an initial systematic comparison of nematode 50 community structure and diversity, derived from morphological identification and molecular 51 identification based on DNA extracted either directly from soil or from elutriated nematodes.

52

53 2. Materials and methods

54

55 From each corner of a square metre grassland plot, we collected two intact soil cores of 5.8cm 56 diameter and 10cm depth (ISO 23611-2) directly adjacent to each other. From one core per corner 57 (n=4) DNA was extracted from a random subsample of 10 g (PowerMax Soil DNA isolation kit (MO 58 BIO Laboratories)) and called 'soil extracted DNA'. The other core per corner (n=4) was used to 59 elutriate the nematodes from 200 g of fresh soil with an Oostenbrink elutriator (ISO 23611-4). 60 Elutriated nematodes were sub-divided and one sample frozen before extracting DNA (Qiagen DNeasy Blood & Tissue Kit), resulting in a so-called "diversity soup" (Yu et al. 2012) and one sample 61 62 fixed for morphological identification (Yoder et al., 2006). DNA extracts were subjected to DNA 63 metabarcoding (Porazinska et al. 2009; and supplementary details). Nematode relative abundance 64 data (Table 1 and Supplementary tables 1, 2,) were arcsin transformed for principal component 65 analysis (PCA) and one-way ANOVA. Diversity was calculated as Shannon and reciprocal Simpson 66 indices. Functional indices were calculated using the nematode indicator joint analysis (NINJA) 67 programme (Sieriebriennikov et al., 2014). 68 69 3. Results 70 71 At the family level the DNA based methods revealed more taxa (20) than the morphological 72 analysis (18), while at higher taxonomic resolution the diversity soup method gave more taxa (34 73 OTU's) than the soil extracted DNA (25 OTU's). Increasing taxonomic resolution significantly 74 increased diversity indices (i.e. Shannon 4.4 versus 6.5) and the diversity soup method revealed 75 greater diversity than the soil extracted DNA (i.e. 1/Simpson 2.0 versus 2.3). From the 76 metabarcoding, 76% of reads from the diversity soup and 7% of reads from soil extracted DNA were

nematode sequences. Maturity Index was greatest for the diversity soup community (2.3, 3.4, 2.3 for

78 morphology, diversity soup and soil extracted DNA, respectively), while Basal Index (50, 13, 9) and

Channel Index (33, 15, 4) were both larger for morphology than either DNA method. The
communities fell in different quadrants on an enrichment index vs structure index plot (Fig. 1).
Principal component analysis revealed a different nematode community composition with each
method and by running the analysis to include or exclude rare taxa we could show that patterns are
driven by differences in relative abundance of the main taxa rather than the presence / absence of
rare taxa.

85

86 4. Discussion

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88 The objective of this study was to determine how dependent the metrics for community analysis 89 are on the methods used. Here we show for the first time that different extraction approaches, 90 even an identical high-throughput sequencing approach that targets either DNA of nematodes after 91 being extracted first or directly from extracted DNA, shows not only different taxonomic community 92 composition but most strikingly suggests a different soil quality. We recognise that this is a limited 93 study both in terms of samples analysed and comparatively low sequence depth obtained by 454 94 pyrosequencing, but the principle was to highlight the crucial need for standardisation in comparing 95 between samples. The pattern of the result would have been the same whether we used 454 96 pyrosequencing for HTS or another sequencing platform (Luo et al., 2012; Mahe et al., 2015). 97 The primers (NF-1 and 18Sr2b, Porazinska et al. 2009) give good coverage of soil nematodes and 98 have been widely used, but are not nematode specific and also amplify other eukaryotes. As far as 99 we are aware that there are no universally perfect primers that target all groups of nematodes in the 100 same way, however, primer issues cannot explain differences between the two molecular methods 101 to compare nematode communities. Biases in the extraction/elutriation methods are the only 102 explanation for the observed differences, which implies that we still have only a limited idea how soil 103 nematode communities really look like.

104 An advantage of the diversity soup method is that most of the other soil eukaryotes are removed 105 by elutriation, thus giving a larger number of reads for nematodes than from the soil extracted DNA. 106 As the technology improves and sequence numbers per sample increase, then the simultaneous 107 study of all soil eukaryotes becomes a practical option (de Groot et al., 2016). The greater 108 taxonomic resolution of the DNA methods cannot be matched by morphology, unless it is a 109 painstakingly detailed study which precludes the throughput necessary in contemporary research 110 (Yang et al., 2014), and could be expected to be more informative about community structure than 111 morphology. That nematode community analyses differed between extraction methods, in aspects 112 of diversity, structure and ecological indices, mirrored results from Quist et al. (2016). Other studies 113 have also noted that different sampling methods give individual community results because of their 114 particular biases, so that there is no 'true' biodiversity dataset (Yang et al., 2014). Despite the 115 diversity soup and morphological methods both starting with the same aqueous solution of 116 nematodes, the profound differences in nematode community structure could be attributed to 117 identification skills and/or PCR biases and were partly explained by the relatively small contribution of Tylenchidae and bacterial-feeding nematodes in the diversity soup, as seen in similar comparisons 118 119 (Griffiths et al., 2006; Donn et al., 2011, 2012; Darby et al., 2013). The comparison of soil extracted 120 DNA vs. diversity soup might be affected by sample size, as the 10g soil used for direct extraction is 121 much less than the 200g recommended to reliably reveal a soil nematode community (Wiesel et al., 122 2015). This might explain the lack of larger omnivore and predator nematodes in the soil extracted 123 DNA (such as Aporcelaimellus, Discolaimus, Dorylaimidae, Nygolaimus) (Quist et al., 2017). The 124 calculated functional indices would indicate different soil food web conditions, which is clearly 125 erroneous as we compared the same samples. Therefore method standardisation, including 126 extensive studies using mock communities of known and highly diverse nematode communities, 127 needs to be adopted (as indicated by Darby et al., 2013) in order to be able to compare taxonomic as 128 well as the functional and indicative attributes of soil nematode communities.

129

130	5.	Conc	lusion

132	DNA methods will be increasingly used because of reducing analysis costs, high throughput,
133	greater taxonomic resolution and compatibility with available technical skills. There is a need now to
134	understand the methodological discrepancies (sample size; extraction and PCR biases; primer
135	specificity; read number and taxonomic resolution) identified here and to calibrate the molecular
136	methods to the morphological information. The developing high-throughput molecular methods
137	have to be standardised for ecological and applied indication purposes.
138	
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- 243 Data accessibility
- 244 The sequence data will be uploaded to the European Nucleotide Archive (http://www.ebi.ac.uk/ena)
- 245 on acceptance.
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270	Table 1. The percentage distribution of nematode families determined from a morphological
271	examination of elutriated nematodes (morphology); high throughput sequencing of DNA extracted
272	from elutriated nematodes diversity soup) and DNA directly extracted from soil (soil extract). DNA
273	data have been amalgamated to allow analysis at the same taxonomic resolution as the
274	morphological data. The F-statistic (P) was calculated on arcsin transformed data. Detransformed
275	means are presented. Data also presented on the percentage distribution of nematode feeding
276	types. Means followed by a different letter and in bold are significantly different, $n = 4$.

Nematode family		Method		Р
	Diversity Soup	Morphology	Soil Extract	_
Alaimidae	0.16a	0.00a	1.21b	0.002
Anguinidae	0.04	0.00	0.16	0.244
Aphelenchoididae	0.79a	5.33b	1.21a	0.007
Aporcelaimidae	14.95a	7.43a,b	0.84b	0.045
Cephalobidae	23.58a	45.37b	11.89a	0.007
Diplogasteroidae	0.00	0.00	6.31	0.207
Diphtherophoridae	1.63	0.37	9.31	0.067
Dolichodoridae	0.10	0.18	0.12	0.977
Dorylaimidae	4.43a	0.00b	0.28b	0.003
Microlaimidae	0.72a	0.00a	10.93b	<0.001
Monhysteridae	0.12a	5.56b	0.43a	0.012
Nordidae	0.00	0.12	0.00	0.422
Nygolaimidae	24.17a	0.00b	4.55b	0.005
Paratylenchidae	0.04	0.48	0.00	0.516
Plectidae	13.97	13.55	9.54	0.379
Prismatolaimidae	0.72a	0.00a	31.42b	<0.001
Qudsianematidae	6.46	1.97	0.45	0.134
Rhabditidae	1.60	2.76	0.92	0.342
Tylenchidae	0.08a	13.28b	0.03a	<0.001
Trichodoridae	0.22	0.00	0.24	0.325
Functional groups				
Bacterial Feeders	41.90a	67.67b	80.47b	0.008
Fungal Feeders	2.44	5.88	10.56	0.232
Omnivores	27.23a	10.50b	2.06b	0.025
Plant Feeders	0.52a	15.31b	0.98a	0.001
Predators	24.17a	0.00b	4.55b	0.005