Advancing nematode barcoding: A primer cocktail for the cytochrome *c* oxidase subunit I gene from vertebrate parasitic nematodes

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Abstract

Although nematodes are one of the most diverse metazoan phyla, species identification through morphology is difficult. Several genetic markers have been used for their identification, but most do not provide species-level resolution in all groups, and those that do lack primer sets effective across the phylum, precluding high-throughput processing. This study describes a cocktail of three novel primer pairs that overcome this limitation by recovering cytochrome *c* oxidase I (COI) barcodes from diverse nematode lineages parasitic on vertebrates, including members of three orders and eight families. Its effectiveness across a broad range of nematodes enables high-throughput processing.

Keywords: barcoding, identification, nematodes, primers

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Introduction

Roundworms (Nematoda) are known to be among the most physiologically and ecologically diverse of metazoan phyla, occupying habitats from the deep sea to deserts, and from the tropics to polar permafrost (Brown et al. 1949, 1950; De Ley 2006; Dailey 2009; Asbakk et al. 2010; Vanreusel et al. 2010). The phylum includes freeliving, parasitic, mutualistic, opportunistic and symbiotic taxa (Ott et al. 1991; Clarke 2008) and provides a useful model system for the study of human diseases (Fire et al. 1998; Barr 2005; Jadiya et al. 2011) and a tool for ecosystem surveillance (Sambongi et al. 1999; Marcogliese 2005; Ekschmitt & Korthals 2006; Wu et al. 2010; Denver et al. 2011; Hoess et al. 2011; Palm et al. 2011). However, nematodes are also a scourge as many species cause disease in crops, livestock and humans (Hodda & Cook 2009; Manguin et al. 2010). Despite their importance, the taxonomy of nematodes is poorly studied. Species-level identification has traditionally relied on detailed morphological analysis, a task requiring considerable expertise (Coomans 2000) given the morphological conservatism and small size of nematodes (Creer et al. 2010; Powers et al. 2011). Aside from being time-consuming, morphologybased identifications are often problematic because of

Correspondence: Sean W. J. Prosser, Fax: 519-824-5703; E-mail: sprosser@uoguelph.ca high phenotypic plasticity (Coomans 2002; Nadler 2002), the absence of clear diagnostic characters (Wijova *et al.* 2005; Derycke *et al.* 2008) or their restriction to adults in the numerous groups in which larvae are more often encountered (Anderson 2000). Given these constraints, there is recognition that molecular techniques are critical for taxonomic progress (Godfray 2002; Blaxter 2003). Indeed, there are now online databases, such as NemA-TOL (http://nematol.unh.edu/), that are dedicated to organizing and storing ecological and molecular data of nematodes.

Several genetic markers have been used for nematode identification, including small and large subunit ribosomal DNA (SSU and LSU respectively), the internal transcribed spacer (ITS) region of ribosomal DNA and cytochrome c oxidase subunit I (COI) (Blaxter et al. 1998; Floyd et al. 2002; Subbotin et al. 2008; Elsasser et al. 2009; Ferri et al. 2009: Siddal et al. 2012). The ribosomal DNA small subunit (SSU) was the first marker used, and successfully delineated some nematodes but failed to completely explain previous observations based on morphology (Blaxter et al. 1998). As the use of SSU was expanded, it was discovered that the SSU barcode failed to separate many species of nematodes and was better suited for order or family-level discrimination (De Ley et al. 2005). The ribosomal DNA large subunit (LSU) was the second marker used in an attempt to develop a nematode phylogenetic classification system, but

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			Number of specimens studied (successfully			
Order	Family	Genus	sequenced)	Host species	Locality	Collection date
Panagrolaimida	Rhabdiasidae	Rhabdias sp. 1	1 (1)	Smilisca	Colima: Hwy Colima	7 July 2008
Panagrolaimida	Rhahdiacidae	Rhahdias en 7	4 (4)	baudinti Rana sh	-Minatitlan Navani: 5 of Hyyy Bamanca	25 Luna 2009
nnnmnorgnin i					del Oro: Barranqueño bridge	
Panagrolaimida	Rhabdiasidae	Rhabdias sp. 3	5 (5)	Rhinella marina	Colima: Comala	7 July 2008
Panagrolaimida	Rhabdiasidae	Rhabdias lamothei	4 (4)	Leptodeira sp.	Colima: Hyw 98 Minatitlan -Manzanillo	8 July 2008
Rhabditida	Molienidae*	Oswaldocruzia sp.	6) 6	Phrynohyas	Colima: Hyw Colima- Minotitlon	6 July 2008
				Smilisca baudinii	Colima: Hyw 98 Minatitlan	0007 (mf ,
					-Manzanillo	
Rhabditida	Diaphanocephalidae*	Kalicephalus sp.	2 (2)	Leptodeira sp.	Colima: Comala	27 June 2009
				Imantodes sp.	Colima: Ixtlahuacan	24 June 2009
Spirurida	Heterakidae	Strongyluris sp.	1(1)	Trimorphodon	Colima: Hwy 98 Minatitlan	8 July 2008
				biscutatus	-Manzanillo	
Spirurida	Pharyngodonidae	Ozolaimus sp.	5 (0)	Ctenosaura sp.	Colima: Hyw 54 Ixtlahuacan	8 July 2008
Spirurida	Pharyngodonidae	Parapharyngodon sp.	4 (4)	Phrynohyas	Colima: Hyw 98 Minatitlan	8 July 2008
				venulosa	-Manzanillo	
Spirurida	Pharyngodonidae	gen sp. 1	15 (10)	Sceloporus sp.	Jalisco: ND	25 July 2009
Spirurida	Pharyngodonidae	gen sp. 2	2 (2)	Sceloporus formosus	Veracruz: Hyw Xico	28 July 2004
					Viejo- Matlalapa	
Spirurida	Cosmocercidae	Aplectana sp.	18 (17)	Rana pustulosa	Nayarit: S of Hyw Barranca	24 June 2009
				Bufo sp.	del Oro: Barranqueño bridge	
				Leptodeira sp.	Nayarit: Hyw Uzeta-La Gloria	
Commido	On the constrained and	Edmollidae and	11/11)	Dana michilana	Comma Contata Marriati C of Uting Barranoo	000C 0000 L
opuulua	Olicitocerciade	rowyennes sp.		Rana psilonota	del Oro: Barranqueño bridge	24 June 2009 30 June 2010
					Jalisco: Zapopan: Barranca	
					del río Santiago	
Spirurida	Physalopteridae	Physaloptera sp.	7 (7)	Trimorphodon	Michoacan: Hwy 200	5 July 2008
				biscutatus	between La placita and Marnata	
Spirurida	Physalopteridae	gen sp. 1	5 (5)	Sceloporus sp.	Jalisco: ND	25 June 2009
Spirurida	Physalopteridae	gen sp. 2	1 (1)	Imantodes sp.	Colima: Hyw Comala -Minatitlán	25 June 2009
Spirurida	Physalopteridae	Turgida sp.	1 (1)	Didelphis virginiana	Jalisco: Zapopan: Barranca del río Santiago	30 June 2010

requires the amplification of multiple regions to be effective (De Ley *et al.* 2005; Subbotin *et al.* 2008). Similar studies using ITS revealed that a lack of phylum-wide primers combined with difficulties in aligning the extremely variable ITS sequences precluded its use as a universal nematode identification marker amenable to high-throughput platforms (Floyd *et al.* 2002; De Ley *et al.* 2005).

The mitochondrial gene cyctochrome c oxidase subunit I (COI) has also been explored as a potential marker on which to base a nematode phylogenetic classification system (Floyd et al. 2002; Elsasser et al. 2009). In addition to being a mitochondrial gene, COI is translated into an evolutionarily conserved protein and thus has some advantages over SSU, LSU and ITS. However, COI is not immune to the inherent problems associated with nematode barcoding. While the 5' region of COI has been shown to separate nematodes into proper species (Derycke et al. 2010), a phylum-wide primer set has yet to be developed (De Ley et al. 2005). In this study, we report the development of a primer cocktail which enables the recovery of COI barcodes from a broad range of nematode parasites of vertebrates in a high-throughput manner and delivers species-level resolution.

Materials and methods

Specimen collection

Ninety-five adult nematodes collected in Mexico from various reptilian, amphibian and mammalian hosts were analysed (Table 1). Each specimen was collected in duplicate (i.e. from the same habitat within the same host), with one stored in 95% ethanol for DNA extraction and the other cleared on a glass slide with undiluted glycerine to enable identification to family, genus or species level using morphological characteristics (Table 1).

Primer design

Cytochrome *c* oxidase subunit I (COI) sequences were obtained from 56 mitochondrial genome sequences from nematodes in GenBank (Table 2) and aligned using online EBI CLUSTALW2 software (Larkin *et al.* 2007). A lepidopteran COI sequence was included in the alignment as a reference for locating the standard primer binding sites (Folmer *et al.* 1994) for COI barcoding (Hebert *et al.* 2003a,b). The forward and reverse primer binding sites were excised from the 56 sequences and

 Table 2
 Nematode COI sequences used to design cocktail primers

GenBank Accession	Species	GenBank Accession	Species
NC_008231	Agamermis sp. BH-2006	AJ556134	Necator americanus
FJ483518	Ancylostoma caninum	NC_003416	Necator americanus
NC_003415	Ancylostoma duodenale	GQ888716	Oesophagostomum dentatum
GQ398121	Angiostrongylus cantonensis	FM161883	Oesophagostomum quadrispinulatum
GQ398122	Angiostrongylus costaricensis	NC_001861	Onchocerca volvulus
NC_007934	Anisakis simplex	FN313571	Radopholus similis
NC_001327	Ascaris suum	NC_008640	Romanomermis culicivorax
NC_004298	Brugia malayi	NC_008693	Romanomermis iyengari
FJ483517	Bunostomum phlebotomum	EF175763	Romanomermis nielseni
NC_009885	Caenorhabditis briggsae	GU138699	Setaria digitata
EU407789	Caenorhabditis briggsae	NC_005941	Steinernema carpocapsae
EU407793	Caenorhabditis briggsae	DQ520860	Strelkovimermis spiculatus
EU407804	Caenorhabditis elegans	NC_008047	Strelkovimermis spiculatus
NC_001328	Caenorhabditis elegans	AJ558163	Strongyloides stercoralis
EU407805	Caenorhabditis elegans	GQ888717	Strongylus vulgaris
EU407780	Caenorhabditis sp.	GQ888718	Syngamus trachea
GQ888721	Chabertia ovina	GQ888720	Teladorsagia circumcincta
HM773029	Chandlerella quiscali	DQ520858	Thaumamermis cosgrovei
NC_004806	Cooperia oncophora	NC_008046	Thaumamermis cosgrovei
GQ888712	Cylicocyclus insignis	AM411108	Toxocara canis
NC_005305	Dirofilaria immitis	AM411622	Toxocara cati
EU281143	Enterobius vermicularis	AM412316	Toxocara malaysiensis
NC_010383	Haemonchus contortus	FJ664617	Toxocara vitulorum
NC_008534	Heterorhabditis bacteriophora	GU386314	Trichinella spiralis
NC_008828	Hexamermis agrotis	NC_002681	Trichinella spiralis
GQ888722	Mecistocirrus digitatus	GQ888719	Trichostrongylus axei
GQ888714	Metastrongylus pudendotectus	GQ888711	Trichostrongylus vitrinus
GQ888715	Metastrongylus salmi	NC_005928	Xiphinema americanum

4 S. W. J. PROSSER ET AL.

Table 3 Primers used in this study. M13 tails are in lowercase bold

Primer	Sequence $(5' \rightarrow 3')$	Reference
NemF1_t1	tgtaaaacgacggccagtCRACWGTWAATCAYAARAATATTGG	This study
NemF2_t1	tgtaaaacgacggccagtARAGATCTAATCATAAAGATATYGG	This study
NemF3_t1	tgtaaaacgacggccagtARAGTTCTAATCATAARGATATTGG	This study
NemR1_t1	caggaaacagctatgactAAACTTCWGGRTGACCAAAAAATCA	This study
NemR2 t1	caggaaacagctatgactAWACYTCWGGRTGMCCAAAAAAYCA	This study
NemR3 t1	caggaaacagctatgactAAACCTCWGGATGACCAAAAAATCA	This study
LCO1490_t1	tgtaaaacgacggccagtGGTCAACAAATCATAAAGATATTGG	Folmer et al. 1994
HCO2198 t1	caggaaacagctatgacTAAACTTCAGGGTGACCAAAAAATCA	Folmer et al. 1994
M13F	TGTAAAACGACGGCCAGT	Messing 1993
M13R	CAGGAAACAGCTATGAC	Messing 1993

phenograms for the two primer binding sites were generated using EBI CLUSTALW2. Both trees revealed three clusters (not shown) and the consensus sequence for each cluster was used to design a primer cocktail consisting of one primer for each cluster (i.e. three forward and three reverse primers). The three primer sequences in each cocktail were tailed with modified M13 sequences (Messing 1993) as described in Ivanova *et al.* (2007). The three forward and three reverse primers were mixed in a 1:1:1 ratio to make the final forward (C_NemF1_t1: NemF1_t1 + NemF2_t1 + NemF3_t1) and reverse (C_NemR1_t1: NemR1_t1 + NemR2_t1 + NemR3_t1) cocktails (Table 3).

DNA extraction, PCR amplification and sequencing

Total DNA was extracted from whole nematodes using standard glass fibre methods (Ivanova et al. 2006). After purification, 2 µL of DNA was added to a PCR reaction consisting of 6.25 μ L of 10% D-(+)-trehalose dihydrate (Fluka Analytical), 2.00 µL of Hyclone ultra-pure water (Thermo Scientific), 1.25 µL of 10X PlatinumTaq buffer (Invitrogen), 0.625 µL of 50 mM MgCl₂ (Invitrogen), 0.125 μ L of each primer or primer cocktail, 0.0625 μ L of 10 mM dNTP (KAPA Biosystems) and 0.060 μ L of 5 U/ μ L PlatinumTaq DNA Polymerase (Invitrogen) for a total reaction volume of 12.5 μ L. Thermal cycling conditions were 94 °C for 1 min, five cycles at 94 °C for 40 s, 45 °C for 40 s, 72 °C for 1 min, followed by 35 cycles at 94 °C for 40 s, 51 °C for 40 s, 72 °C for 1 min and a final extension at 72 °C for 5 min. The resulting amplicons were visualized on a 2% agarose E-gel® 96 precast gel (Invitrogen) and bidirectionally sequenced using M13F and M13R as sequencing primers (Table 3).

Cycle sequencing was performed using a modified BigDye 3.1 Terminator (Applied Biosystems) protocol (Hajibabaei *et al.* 2005). Cycle sequencing conditions were 96 °C for 1 min followed by 35 cycles at 96 °C for 10 s, 55 °C for 5 s, 60 °C for 2.5 min and a final extension at 60 °C for 5 min. Sequencing was performed on an ABI

Table 4 PCR success rates of nematode cocktail primers $(C_NemF1_t1 + C_NemR1_t1)$ compared with Folmer primers $(LCO1490_t1 + HCO2198_t1)$

Primers	Number of PCR positives	Success Rate
C_NemF1_t1 + C_NemR1_t1	85/95	89.5%
LCO1490_t1 + HCO2198_t1	83/95	87.4%

Table 5 Sequencing success rates and trace quality scores ofPCR products generated with nematode cocktail primers $(C_NemF1_t1 + C_NemR1_t1)$ or Folmer primers $(LCO1490_t1 + HCO2198_t1)$. Sequencing success rates werecalculated by dividing the number of recovered sequences (afterediting) by the total number of sequenced samples (i.e. 95)

Primers	Average PHRED Score	Success Rate (661 bp only)	Success Rate (any sequence over 100 bp)
C_NemF1_t1 + C_NemR1_t1	49	88.4%	88.4%
LCO1490_t1 + HCO2198_t1	44	65.2%	75.7%

3730XL capillary sequencer (Applied Biosystems). Traces were assembled and edited using CodonCode v. 3.0.1 (CodonCode Corporation, Dedham, Massachusetts). Trace quality scores were calculated using KB Basecaller (ABI software) and trace statistics were calculated using Sequence Scanner (Applied Biosystems). Sequences have been deposited in BOLD (www.boldsystems.org) under sample ID's MXHEL359–MXHEL453 within the project entitled: Parasitic nematodes from Mexican vertebrates (NEMNP) and in GenBank under accession numbers KC130665 - KC130748.

Since previous studies (Elsasser *et al.* 2009; Derycke *et al.* 2010) reported varying success in barcode recovery with a commonly used primer set (LCO1490 and HCO2198, Folmer *et al.* 1994), we compared the success of sequence recovery with M13-tailed versions of LCO1490

and HCO2198 (Table 3) and our new cocktail. All PCR reagents were identical between the two primer sets, and the same DNA templates were employed. For each primer

set, all 95 nematode samples were sequenced, even if an amplicon was not visible on the E-gel. Sequences were aligned using EBI CLUSTALW2, imported into MEGA5



Fig. 1 Neighbour-joining tree of COI barcode sequences generated by the nematode cocktail primers. A divergence of 2% or greater is indicative of a separate operational taxonomic unit. Codes following names of taxa refer to GenBank accession numbers.

(Tamura *et al.* 2011), and a neighbour-joining algorithm (NJ) was used to generate a phenogram.

Results

PCR success rates (Table 4), as measured by the presence or absence of a visible amplicon on the E-gel, were very similar with the Folmer primers (87%) and the new primer cocktail (89%) (Fisher's exact test, P = 0.8212). However, there was a marked difference in sequence quality and recovery (Table 5). The traces produced by the primer cocktail (n = 188) had a mean PHRED score of 49 (SD = 12), whereas those produced by the Folmer primers (n = 181) had a mean PHRED score of 44 (SD = 12) (Student's t-test, P = 0.0001). However, any traces with PHRED quality scores between 40 and 50 are usually equally interpretable (personal observation). More importantly, full-length barcodes (661 bp) were recovered from 88% of the specimens with the new primer cocktail, but from just 65% of reactions which employed the Folmer primers (Fisher's exact test. P = 0.0001). DNA barcodes were obtained from a total of 84 specimens with 62 yielding full-length sequences with both primer sets, while 12 were only recovered by the cocktail, and another 10 were fully recovered by the cocktail but only partially (~500 bp) by the Folmer primers. Every sequence generated by the cocktail allowed the assignment of its source specimen to an operational taxonomic unit that agreed with its morphological identification (Martínez-Salazar 2008; Velarde-Aguilar, personal observation) (Fig. 1). Individuals from all genera were successfully barcoded except Ozolaimus (Table 1); its failure may reflect poor DNA preservation since we observed that ethanol partially evaporated from the vial that kept these specimens, and sequences were successfully recovered from members of closely related genera.

Discussion

The Nematoda may be the most species-rich phylum of animals, with approximately 27 000 described species (Hugot *et al.* 2001; Hodda 2011), but taxonomic knowledge must progress significantly to validate this hypothesis. The ribosomal DNA small subunit (SSU) (Blaxter *et al.* 1998), large subunit (LSU) (Subbotin *et al.* 2008) and internal transcribed spacer (ITS) region (Floyd *et al.* 2002; De Ley *et al.* 2005) have all been used as a tool for species discrimination, but the lack of phylum-wide primers or their failure to delineate closely allied species in certain nematode groups limit their utility in the analysis of nematode diversity (De Ley *et al.* 2005). The COI gene has also been explored as a potential marker for species identification (Floyd *et al.* 2002; Elsasser *et al.* 2009) because of its effectiveness in other major animal phyla (e.g. Campagna et al. 2010; Clare et al. 2011; Kumar et al. 2012; Weigt et al. 2012). The barcode region of COI has delivered species-level resolution in certain nematode lineages (Derycke et al. 2010), but sequence recovery has proven difficult (De Ley et al. 2005). The primer cocktail developed in this study appears to overcome this difficulty as it recovered full-length barcode sequences from nematodes belonging to three orders and eight families (Fig. 1), while 25% of the PCR products from Folmer primers contained co-amplified sequences, perhaps reflecting poor binding with the target COI gene. Moreover, the sequences recovered from our cocktail were able to differentiate congeneric species, such as the four species of Rhabdias, each from a different host and showing consistent morphological differences as detected by Martínez-Salazar (2008) (Fig. 1; Table 1). Although we examined various taxa of nematodes parasitic of vertebrates, further testing is required to validate the effectiveness of our primer set across the phylum. We examined representatives from three of the six currently recognized nematode orders parasitic on vertebrates (Hodda 2011), all representatives of the Class Chromadorea. However, it is possible that our primer cocktail is effective across a large diversity of nematodes because the primers were designed based on members of the Class Dorylaimea and other orders of Chromadorea of medical and veterinary importance. An obvious next step will involve testing barcode recovery from representatives of other orders of parasitic and free-living nematodes.

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References

- Anderson RC (2000) Nematode Parasites of Vertebrates: Their Development and Transmission, 2nd edn. CABI Publishing, Oxford, UK.
- Asbakk K, Aars J, Derocher AE et al. (2010) Serosurvey for Trichinella in polar bears (Ursus maritimus) from Svalbard and the Barents Sea. Veterinary Parasitology, 172, 256–263.
- Barr MM (2005) Caenorhabditis elegans as a model to study renal development and disease: sexy cilia. Journal of the American Society of Nephrology, 16, 305–312.
- Blaxter M (2003) Counting angels with DNA. Nature, 421, 122–124.
- Blaxter ML, De Ley P, Garey JR et al. (1998) A molecular evolutionary framework for the phylum Nematoda. *Nature*, **392**, 71–75.

PRIMERS FOR NEMATODE DNA BARCODING 7

- Brown M, Cronk B, DeSinner F, Green JE, Gibbons JE, Kuitunen-Ekbaum E (1949) A note on trichinosis in animals of the Canadian Northwest Territories. *Canadian Journal of Public Health*, 40, 20–21.
- Brown M, Green JE, Boag TJ, Kuitunen-Ekbaum E (1950) Parasitic infections in the Eskimos at Igloolik, N.W.T. Canadian Journal of Public Health, 41, 508–512.
- Campagna L, Lijtmaer DA, Kerr KCR et al. (2010) DNA barcodes provide new evidence of a recent radiation in the genus Sporophila (Aves: Passeriformes). Molecular Ecology Resources, 10, 449–458.
- Clare EL, Lim BK, Fenton MB, Hebert PDN (2011) Neotropical bats: estimating species diversity with DNA barcodes. *PLoS ONE*, **6**, e22648.
- Clarke DJ (2008) *Photorhabdus*: a model for the analysis of pathogenicity and mutualism. *Cellular Microbiology*, **10**, 2159–2167.
- Coomans A (2000) Nematode systematics: past, present and future. Nematology, 2, 3–7.
- Coomans A (2002) Present status and future of nematode systematics. *Nematology*, 4, 573–582.
- Creer S, Fonseca VG, Porazinska DI *et al.* (2010) Ultrasequencing of the meiofaunal biosphere: practice, pitfalls and promises. *Molecular Ecology*, **19**(Suppl. 1), 4–20.
- Dailey MD (2009) A new species of *Parafilaroides* (Nematoda: Filaroididae) in three species of fur seals (Carnivora: Otariidae) from the southern hemisphere. *Journal of Parasitology*, **95**, 156–159.
- De Ley P (2006) A quick tour of nematode diversity and the backbone of nematode phylogeny. WormBook: the online review of C. elegans biology, 25, 1–8.
- De Ley P, De Ley IT, Morris K et al. (2005) An integrated approach to fast and informative morphological vouchering of nematodes for applications in molecular barcoding. *Philosophical Transactions of the Royal Society B-Biological Sciences*, **360**, 1945–1958.
- Denver DR, Clark KA, Raboin MJ (2011) Reproductive mode evolution in nematodes: insights from molecular phylogenies and recently discovered species. *Molecular Phylogenetics and Evolution*, 61, 584–592.
- Derycke S, Fonseca G, Vierstraete A, Van-Fleuteren J, Vincx M, Moens T (2008) Disentangling taxonomy within the *Rhabditis (Pellioditis) marina* (Nematoda, Rhabditidae) species complex using molecular and morphological tools. *Zoological Journal of the Linnaean Society*, **152**, 1–15.
- Derycke S, Vanaverbeke J, Rigaux A, Backeljau T, Moens T (2010) Exploring the use of cytochrome oxidase *c* subunit 1 (COI) for DNA barcoding of free-living marine nematodes. *PLoS ONE*, 5, e13716.
- Ekschmitt K, Korthals GW (2006) Nematodes as sentinels of heavy metals and organic toxicants in the soil. *Journal of Nematology*, **38**, 13–19.
- Elsasser SC, Floyd R, Hebert PDN, Schulte-Hostedde AI (2009) Species identification of North American guinea worms (Nematoda: *Dracunculus*) with DNA barcoding. *Molecular Ecology Resources*, **9**, 707–712.
- Ferri E, Barbuto M, Bain O et al. (2009) Integrated taxonomy: traditional approach and DNA barcoding for the identification of filarioid worms and related parasites (Nematoda). Frontiers in Zoology, 6, 1–12.
- Fire A, Xu S, Montgomery MK, Kostas SA, Driver SE, Mello CC (1998) Potent and specific genetic interference by double-stranded RNA in *Caenorhabditis elegans*. Nature, 391, 806–811.
- Floyd R, Abebe E, Papert A, Blaxter M (2002) Molecular barcodes for soil nematode identification. *Molecular Ecology*, **11**, 839–850.
- Folmer O, Black M, Hoeh W, Lutz R, Vrijenhoek R (1994) DNA primers for amplification of mitochondrial cytochrome *c* oxidase subunit I from diverse metazoan invertebrates. *Molecular Marine Biology and Biotechnology*, **3**, 294–299.

Godfray HCJ (2002) Challenges for taxonomy. Nature, 417, 17-19.

- Hajibabaei M, DeWaard JR, Ivanova NV et al. (2005) Critical factors for assembling a high volume of DNA barcodes. *Philosophical Transactions* of the Royal Society B-Biological Sciences, 360, 1959–1967.
- Hebert PDN, Cywinska A, Ball SL, DeWaard JR (2003a) Biological identifications through DNA barcodes. Proceedings of the Royal Society of London Series B-Biological Sciences, 270, 313–321.
- Hebert PDN, Ratnasingham S, DeWaard JR (2003b) Barcoding animal life: cytochrome c oxidase subunit 1 divergences among closely related species. Proceedings of the Royal Society of London Series B-Biological Sciences, 270, S96–S99.

Hodda M (2007) Phylum Nematoda In: Linnaeus Tercentenary: Progress in Invertebrate Taxonomy. (eds Zhang Z.-Q., Shear W. A.) Zootaxa 1668, 1–766.

- Hodda M (2011) Phylum Nematoda Cobb 1932. In: Animal biodiversity: An outline of higher-level classification and survey of taxonomic richness. (ed Zhang Z.-Q.) Zootaxa, 3148, 63–95.
- Hodda M, Cook DC (2009) Economic impact from unrestricted spread of potato cyst nematodes in Australia. *Phytopathology*, **99**, 1387–1393.
- Hoess S, Claus E, Von der Ohe PC et al. (2011) Nematode species at risk -A metric to assess pollution in soft sediments of freshwaters. Environment International, 37, 940–949.
- Hugot JP, Baujard P, Morand S (2001) Biodiversity in helminths and nematodes as a field of study: an overview. *Nematology*, **3**, 199–208.
- Ivanova NV, Dewaard JR, Hebert PDN (2006) An inexpensive, automation-friendly protocol for recovering high-quality DNA. *Molecular Ecol*ogy Notes, 6, 998–1002.
- Ivanova NV, Zemlak TS, Hanner RH, Hebert PDN (2007) Universal primer cocktails for fish DNA barcoding. *Molecular Ecology Notes*, 7, 544–548.
- Jadiya P, Khan A, Sammi SR, Kaur S, Mir SS, Nazir A (2011) Anti-Parkinsonian effects of *Bacopa monnieri*: insights from transgenic and pharmacological *Caenorhabditis elegans* models of Parkinson's disease. *Biochemical and Biophysical Research Communications*, **413**, 605–610.
- Kumar NP, Srinivasan R, Jambulingam P (2012) DNA barcoding for identification of sand flies (Diptera: Psychodidae) in India. *Molecular Ecol*ogy Resources, **12**, 414–420.
- Larkin MA, Blackshields G, Brown NP *et al.* (2007) Clustal W and Clustal X version 2.0. *Bioinformatics*, **23**, 2947–2948.
- Manguin S, Bangs MJ, Pothikasikorn J, Chareonviriyaphap T (2010) Review on global co-transmission of human *Plasmodium* species and *Wuchereria bancrofti* by *Anopheles* mosquitoes. *Infection Genetics and Evolution*, 10, 159–177.
- Marcogliese D (2005) Parasites of the superorganism: are they indicators of ecosystem health? International Journal of Parasitology, 35, 705–716.
- Martínez-Salazar EA (2008) Sistemática y biogeografía del género Rhabdias Stiles y Hassall, 1905 (Nematoda: Rhabdiasidae) en México. Tesis DoctoradoPosgrado en Ciencias Biológicas, Universidad Nacional Autónoma de México, 495pp.
- Messing J (1993) M13 cloning vehicles: their contribution to DNA sequencing. *Methods in Molecular Biology*, 23, 9–22.
- Nadler SA (2002) Species delimitation and nematode biodiversity: phylogenies rule. *Nematology*, 4, 615–625.
- Ott JA, Novak R, Schiemer F, Hentschel U, Nebelsick M, Polz M (1991) Tackling the sulfide gradient: a novel strategy involving marine nematodes and chemoautotrophic ectosymbionts. *Marine Ecology*, 12, 261–279.
- Palm HW, Kleinertz S, Ruckert S (2011) Parasite diversity as an indicator of environmental change? An example from tropical grouper (*Epinephelus fuscoguttatus*) mariculture in Indonesia. *Parasitology*, **138**, 1793–1803.
- Powers T, Harris T, Higgins R, Mullin P, Sutton L, Powers K (2011) MO-TUs, morphology, and biodiversity estimation: a case study using nemtodes of the suborder criconematina and a conserved 18S DNA barcode. *Journal of Nematology*, 43, 35–48.
- Sambongi Y, Nagae T, Liu Y *et al.* (1999) Sensing of cadmium and copper ions by externally exposed ADL, ASE, and ASH neurons elicits avoidance response in *Caenorhabditis elegans*. *NeuroReport*, **10**, 753–757.
- Siddal ME, Kivst S, Phillips A, Oceguera-Figueroa A (2012) DNA barcoding of parasitic nematodes: is it Kosher? *Journal of Parasitol*ogy, 98, 692–694.
- Subbotin SA, Ragsdale EJ, Mullens T, Roberts PA, Mundo-Ocampo M, Baldwin JG (2008) A phylogenetic framework for root lesion nematodes of the genus *Pratylenchus* (Nematoda): evidence from 18S and D2 –D3 expansion segments of 28S ribosomal RNA genes and morphological characters. *Molecular Phylogenetics and Evolution*, 48, 491–505.
- Tamura K, Peterson D, Peterson N, Stecher G, Nei M, Kumar S (2011) MEGA5: molecular evolutionary genetics analysis using maximum likelihood, evolutionary distance, and maximum parsimony methods. *Molecular Biology and Evolution*, 28, 2731–2739.

8 S. W. J. PROSSER ET AL.

- Vanreusel A, De Groote A, Gollner S, Bright M (2010) Ecology and biogeography of free-living nematodes associated with chemosynthetic environments in the deep sea: a review. *PLoS ONE*, **5**, e12449.
- Weigt LA, Baldwin CC, Driskell A, Smith DG, Ormos A, Reyier EA (2012) Using DNA barcoding to assess Caribbean reef fish biodiversity: expanding taxonomic and geographic coverage. *PLoS ONE*, 7, e41059.
- Wijova M, Moravec F, Horak A, Modry D, Lukes J (2005) Phylogenetic position of *Dracunculus medinensis* and some related nematodes inferred from 18S rRNA. *Parasitology Research*, 96, 133–135.
- Wu HC, Chen PC, Tsay TT (2010) Assessment of nematode community structure as a bioindicator in river monitoring. *Environmental Pollution*, 158, 1741–1747.

S.W.J.P. wrote the initial manuscript. M.G.V.A., V.L.R. and P.D.N.H. edited and contributed to the manuscript. M.G.V.A. and V.L.R. collected and identified all nematode specimens. S.W.J.P. designed nematode primers and performed all molecular laboratory work and sequence editing/interpretation. M.G.V.A. and V.L.R. analysed and interpreted genetic-distance results.

Data Accessibility

DNA sequences: GenBank accessions KC130665 - KC130748 and BOLD (www.boldsystems.org) sample ID's MXHEL 359-MXHEL453.

A spreadsheet with sampling and taxonomic details for each individual, GenBank accession numbers for its DNA sequences, and its BOLD entry uploaded as online supplementary material.

DNA sequence alignment used to design primers, final DNA sequence alignment and phylogenetic data: BOLD (www.boldsystems.org) project 'Parasitic nematodes from Mexican vertebrates' (NEMNP).

Supporting Information

Additional Supporting Information may be found in the online version of this article:

Table S1 Specimen collection and taxonomic details.BOLDprocess ID's and GenBank accession numbers are listed for eachspecimen.