Morphological and Mitochondrial Genomic Characterization of Eyeworms (*Thelazia callipaeda*) from Clinical Cases in Central China

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INTRODUCTION

*Thelazia callipaeda* Railliet and Henry, 1910 (Nematoda: Spirurida: Thelaziidae), also known as the oriental eyeworm, is the major etiological agent of human thelaziasis. Cases of thelaziasis have increased in recent years in China. Although this species is of medical importance, the genetics and phylogenetic systematics of *T. callipaeda* are poorly understood. In this study, we first reported three cases of thelaziasis in central China. All clinical isolates were identified as *T. callipaeda* according to morphological characteristics by light microscopy and scanning electron microscopy. Next, complete mitochondrial (mt) genomes for the three *T. callipaeda* isolates from different geographical locations were fully characterized using an Illumina sequencing platform. In addition, all available mt genomes of spirurid nematodes in GenBank were included to reconstruct the phylogeny and to explore the evolutionary histories of the isolates. The genome features of the *T. callipaeda* isolates contained 12 PCGs, 22 transfer RNA genes, two ribosomal RNA genes and a major non-coding region. The mtDNA nucleotide sequences of the *T. callipaeda* isolates from different hosts and different locations were similar. The nad6 gene showed high sequence variability among all isolates, which is worth considering for future population genetic studies of *T. callipaeda*. Phylogenetic analyses based on maximum parsimony and Bayesian inference methods revealed close relationships among Thelaziidae, Onchocercidae, Setariidae, Gongylonematidae, Physalopteridae, Dracunculidae, and Philometridae. The monophyly of the *T. callipaeda* isolates from different hosts and distinct geographical locations was confirmed. The entire mt genomes of *T. callipaeda* presented in this study will serve as a useful dataset for studying the population genetics and phylogenetic relationships of *Thelazia* species.

Keywords: *Thelazia callipaeda*, thelaziasis, mitochondrial genome, next generation sequencing, phylogeny
but most cases occur in eastern and southeastern Asia (Otranto and Dutto, 2008; Nguyen et al., 2012). Human cases are usually associated with poor health and socioeconomic settings, in which heavily affected domestic and wild animals live in close vicinity with humans (Hodžić et al., 2014). China probably has the largest number of cases of thelaziasis in the world with more than 600 cases reported to date (Shen et al., 2006; Wang et al., 2014). In addition, the cases of thelaziasis in China appear to have increased in recent years (Wang et al., 2014). In clinical diagnostics, it is difficult to differentiate thelaziasis from allergic conjunctivitis when some of the adult or larval stages of T. callipaeda are present in the eyes of infected patients (Liu et al., 2013a). Fortunately, the DNA barcoding approach has been useful for Thelazia species (Otranto and Traversa, 2004). Several DNA markers have been applied in the taxonomy and genetic studies of T. callipaeda, such as cytochrome c oxidase subunit 1 gene, cox1 (Otranto et al., 2005b; Nguyen et al., 2012; Wang et al., 2014; Mihalca et al., 2015), and the first internal transcribed spacer of ribosomal DNA, ITS1 (Otranto and Traversa, 2004). However, in comparison with a single gene, the complete mitochondrial (mt) genome can provide more markers for molecular identification, and provide the potential to discover population variants or cryptic species (Ramesh et al., 2012; Zhang et al., 2016). In recent years, several remarkable advances have been made in the different taxonomic levels of spirurid nematodes using complete mt genomes (Yatawara et al., 2010; Park et al., 2011; McNulty et al., 2012). The first complete mtDNA sequence of T. callipaeda has been sequenced and characterized (Liu et al., 2013a). These available mtDNA datasets offer opportunities to compare the diversity of mt genomes among closely related Thelazia taxa. In this study, we first describe three cases of thelaziasis in central China. Additionally, the worms collected from patients were sequenced using next generation sequencing (NGS) technology to perform a mitogenomics comparative analysis of spirurid nematodes and to discuss the implications of the new dataset as a new resource for future genetic studies of T. callipaeda populations.

MATERIALS AND METHODS

Ethics Statement
This study was approved by the Life Science Ethics Committee of Zhengzhou University (No. 2016-0096). All worms were collected from eyes of children after their legal guardians provided permission. These worm samples were collected not with the purpose of the present study but were removed from patients to treat the thelaziasis.

Sampling and Data Collection
Three samples of T. callipaeda named HeN-ZZ1, HeN-LY1, and HeN-PDSI were harvested from three patients from different geographical locations of central China (details are shown in the ‘cases report’ part). In addition, all related complete mt genomes available in the GenBank database were included in this study (Table 1). To date (January 1, 2017), GenBank contains 24 mitogenomes of spirurid nematodes representing the following eight families: Dracunculidae (1 mtDNAs), Philometridae (1), Onchocercidae (13), Setariidae (1), Gnathostomatidae (3), Physalopteridae (1), Gongylonematidae (1) and Thelazidae (3). Two Ascaris nematodes, Ascaris lumbricoides (HQ704900) and A. suum (HQ704901), were used as the outgroup.

Morphological Identification
The nematodes collected from the patients were transferred to Petri dishes containing physiological saline (0.9% NaCl). The eyeworms were fixed as previously described (Otranto et al., 2004) and examined by light microscopy (Leica Micro system, German) at different magnifications, as well as under the scanning electron microscopy (SEM). All important morphological characteristics of T. callipaeda were evaluated for adult forms as reported in Otranto et al. (2003).

Extraction of Genomic DNA and Library Preparation
Genomic DNA of each specimen was extracted and purified using EasyPure Genomic DNA Kit (Transgen, China) following the manufacturer’s instructions. The complete mt genome was amplified by long-PCR as two overlapping amplicons from the genomic DNA using primers described by Liu et al. (2013a). Next, these two overlapping products were mixed in approximately equal amounts after determining the concentration of each amplicon. The normalized mixed DNA was fragmented to an average size of 500 bp using Covaris M220 system (Covaris, Woburn, MA, United States). The library was prepared using TruSeq DNA PCR-Free Sample Preparation Kit (Illumina, United States) following the manufacturer’s protocols. After these libraries were purified, the DNA concentration of all samples was quantified and 30 ng of each were pooled together. Finally, the oligonucleotide mix was sequenced on an Illumina HiSeq 2000 at the Genewiz Company (Beijing, China).

Mitogenome Assembly and Annotation
The quality of original sequencing reads was evaluated using the FastQC v.0.11.5 (Andrews, 2010). All ambiguous nucleotides and reads with an average quality value (lower than Q20) were excluded from further analysis. The trimmed sequences were mapped against two reference mt genomes of T. callipaeda worms (NC_018363 and AP0177700) using the CLC Genomic Workbench v.7.0.4 (Qiagen, Germany). The mapped sequences were later subjected to de novo assembly. Contigs with hits to mitochondrial genes or genomes were identified and extracted from the CLC Genomic Workbench. A contig identified as mt genome was manually examined for repeats at the beginning and end of the sequence to establish a circular mtDNA. It was then annotated with MITOS (Bernt et al., 2013) followed by manual validation of the coding regions using the NCBI ORF Finder. The sequin file generated from MITOS was edited and submitted to GenBank with the following accession numbers: KY908318, KY908319, and KY908320.

1http://www.ncbi.nlm.nih.gov/softw/gorf.html
### Phylogenetic Analysis

Nucleotide sequences of the 12 protein-coding genes (PCGs) were separately aligned using MEGA v.6.06 (Tamura et al., 2013). The sequences of rns, rnl, and mt-tRNA genes were aligned in MAFFT v.7 (Katoh and Standley, 2013). The program DAMBE v5.2 (Xia and Xie, 2001) was used to measure the nucleotide substitution saturation using the method of Xia et al. (2003) as the substitution saturation masked the phylogenetic signal. Phylograms were constructed using the maximum parsimony (MP) and Bayesian inference (BI) methods, respectively. The MP analyses were performed in PAUP*4b10 (Swofford, 2003) using heuristic searches with TBR branch swapping and 10,000 random addition sequences. Confidence in each node was assessed by boot-strapping (1000 pseudo-replicates, heuristic search of 20 random addition replicates with TBR option). BI analysis was performed in MrBayes v.3.2 (Ronquist et al., 2012) with 10, 000, 000 generations and sampling trees every 100 generations. The best-fit nucleotide substitution model for each data partition was selected by jModelTest 2 (Darriba et al., 2012) under the Akaike Information Criterion (AIC). Stationarity was assessed using a convergence diagnostic. An average standard deviation of the split frequencies (ASDSF) < 0.01 was used as the criteria of convergence between both runs. The consensus tree was drawn after removing the first 20 000 trees (20%) as the burn-in phase. In addition, the data used in the phylogenetic analyses were partitioned under three schemes: partitioned among PCGs (12 partitions, nad5, cox3, cyt b, nad4L, nad4, atp6, nad2, nad1, nad3, cox1, cox2, and nad6), partitioned by rRNA genes (2 partitions, rnl and rns), and partitioned by PCGs and rRNA (14 partitions).

### RESULTS

#### Cases Descriptions and Parasite Identification

The first case was a 6-year-old boy from Lushan county of Pingdingshan city, Henan province (33°36′ N, 112°25′ E) in central China. In September 2014, the boy was admitted to the First Affiliated Hospital of Zhengzhou University with a 7-day history of creeping sensation, itching, and increased secretions in his right eye, but without any systemic or visual symptoms. His medical history was notable only for an unknown insect flying into his right eye approximately 2 months earlier. On ophthalmological examination, several thread-like worms were seen moving in the nasal upper eyelid of the eyes above the conjunctiva. The worms nested in the subconjunctiva tissue were completely cleared with intraocular forceps. Four worms were removed, 1 in the left eye and 3 in the right eye. The second case was a 2.5-year-old boy who lived in the suburbs of Yangzhou city, Jiangsu province (32°14′ N, 119°06′ E). He was admitted to the First Affiliated Hospital of Tongji Medical College, Huazhong University of Science and Technology with a 5-day history of redness and itching in the right eye. On ophthalmological examination, several thread-like worms were seen moving in the nasal upper eyelid of the eyes above the conjunctiva. The worms were removed with intraocular forceps. Four worms were removed, 1 in the left eye and 3 in the right eye. The third case was a 2-year-old boy from Lushan county of Pingdingshan city, Henan province (33°36′ N, 112°25′ E). He was admitted to the First Affiliated Hospital of Zhengzhou University with a 5-day history of redness and itching in the right eye. On ophthalmological examination, several thread-like worms were seen moving in the nasal upper eyelid of the eyes above the conjunctiva. The worms were removed with intraocular forceps. Four worms were removed, 1 in the left eye and 3 in the right eye. The fourth case was a 3-year-old girl from Lushan county of Pingdingshan city, Henan province (33°36′ N, 112°25′ E). She was admitted to the First Affiliated Hospital of Zhengzhou University with a 7-day history of redness and itching in both eyes. On ophthalmological examination, several thread-like worms were seen moving in the nasal upper eyelid of both eyes above the conjunctiva. The worms were removed with intraocular forceps. Four worms were removed, 1 in the left eye and 3 in the right eye.
of Zhengzhou city, Henan province (34°40' N, 113°12' E). He visited his doctor in August 2015 with an approximately 2 weeks history of itching, conjunctival hyperemia and increased secretions in eyes. Ophthalmological examination found six slender worms on the conjunctival sac of the left eye. No additional abnormal manifestations were noted on physical examination. The symptoms disappeared after the worms were removed with forceps. The third case was a 1.5-year-old child (isolated from dog, No. NC_018363) and Japan (No. AP017700) were added, the similarities of the mt genomes declined to 98.06% (Guangdong and HeN-LY1 isolates) to 99.53% (HeN-ZZ1 and Japan isolates). Comparing the protein-coding gene regions (Figure 2), T. callipaeda displayed high levels of sequence conservation ranging from 96.05% of identical nucleotides in atp6 to 98.74% in nad4L. The amino acid sequence similarities ranged from 84.31% (nad6) to 98.73% (cox1). The most divergent amino acid sequence was nad6 between the HeN-PDS1 and Japan isolates. The mtDNA nucleotide sequences from the T. callipaeda isolates from different locations were conspicuously similar. The full-length mt genomes were identical in 98.22% nucleotides of the 13 668 bp long alignment of the three isolates. When published T. callipaeda mt sequences from Guangdong of China (isolated from dog, No. NC_018363) and Japan (No. AP017700) were added, the similarities of the mt genomes declined to 98.89%. Pairwise genetic distances between individual pairs of these isolates measured as a percentage of identical nucleotides ranged from 98.06% (Guangdong and HeN-LY1 isolates) to 99.53% (HeN-ZZ1 and Japan isolates). Comparing the protein-coding gene regions (Figure 2), T. callipaeda displayed high levels of sequence conservation ranging from 96.05% of identical nucleotides in atp6 to 98.74% in nad4L. The amino acid sequence similarities ranged from 84.31% (nad6) to 98.73% (cox1). The most divergent amino acid sequence was nad6 between the HeN-PDS1 and Japan isolates. The mt large and small subunit rDNAs (rrnL, rrnS) shared 89.95 and 96.71% identical nucleotides, respectively.

Assembly of the mt Genome
A total of 961 424, 961 094, and 970 104 sequence reads with a total of 162 140 093, 166 174 524, and 162 301 339 bases were mapped to full mitochondrial genomes for the T. callipaeda isolates of HeN-LY1, HeN-PDS1, and HeN-ZZ1, respectively (Supplementary Table S1). All three novel mt genomes were fully annotated and each encoded an identical set of 12 PCGs, 22 tRNA genes, and 2 rRNA genes. All genes were encoded in an identical order and direction, and all PCGs annotated across the three representatives of T. callipaeda were of the same length (Figure 2). Several genes overlapped at the boundaries, such as nad6/trnR (9 bp), trnL1/cox3 (3 bp), trnL2/trnN (11 bp), nad5/trnF (32 bp), cox2/trnH (10 bp), rrnL/nad3 (8 bp), trnS1/nad2 (3 bp), and trnT/nad4 (6 bp) (Supplementary Table S2). All PCGs had ATG, TTT, ATA, and TAT termination codons and other was the typical crescent shape of the shorter right spicule. (Mc) SEM of anterior region of the male. (Md) SEM of caudal end of male. (Me) SEM of caudal end of male, five pairs of postcloacal papillae.

FIGURE 1 | Photomicrographs and scanning electron micrographs (SEMs) of female and male Thelazia callipaeda. (Fa) Light micrograph of female anterior head and buccal capsule. (Fb) Light micrograph of immature eggs or germ cells filling the uterine tubules of female. (Fc) SEM of the tip of the female tail and the female anus. (Fd) SEM of the transversally striated cuticle on the anterior of female. (Fe) SEM of the anterior region of female, vulva with a short vulvar flap. (Ma) Light micrograph of male anterior head. (Mb) Light micrograph of the male caudal end and detail of the shorter right spicule. (Mc) SEM of anterior region of the male. (Md) SEM of caudal end of male. (Me) SEM of caudal end of male, five pairs of postcloacal papillae.

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The mtDNA nucleotide sequences from the T. callipaeda isolates from different locations were conspicuously similar. The full-length mt genomes were identical in 98.22% nucleotides of the 13 668 bp long alignment of the three isolates. When published T. callipaeda mt sequences from Guangdong of China (isolated from dog, No. NC_018363) and Japan (No. AP017700) were added, the similarities of the mt genomes declined to 98.89%. Pairwise genetic distances between individual pairs of these isolates measured as a percentage of identical nucleotides ranged from 98.06% (Guangdong and HeN-LY1 isolates) to 99.53% (HeN-ZZ1 and Japan isolates). Comparing the protein-coding gene regions (Figure 2), T. callipaeda displayed high levels of sequence conservation ranging from 96.05% of identical nucleotides in atp6 to 98.74% in nad4L. The amino acid sequence similarities ranged from 84.31% (nad6) to 98.73% (cox1). The most divergent amino acid sequence was nad6 between the HeN-PDS1 and Japan isolates. The mt large and small subunit rDNAs (rrnL, rrnS) shared 89.95 and 96.71% identical nucleotides, respectively.
Phylogeny Reconstruction

The test of substitution saturation showed that the observed indexes of substitution saturation (Iss) for all alignments were significantly lower than the corresponding critical index substitution saturation (Iss.c), indicating that there was little saturation in the present sequences (Table 2). The likelihood models identified by the jModelTest (AIC) suggested that the GTR model was most suitable for included data partitions (Supplementary Table S3). A total of six analyses by two phylogenetic inference methods (MP and BI) under three partition schemes (partitioned among PCGs, partitioned by rRNA genes, and PCGs + rRNA) were completed in this study. Generally, the trees generated from different methods with different partition schemes shared similar topologies (Figure 3). The earliest diversifications among these remaining families gave rise to Dracunculidae and Philometridae and then to Physalopteridae. In addition, Dracunculidae and Philometridae composed clade with high support value (posterior probabilities = 1.0). The next diversification event would have separated Thelaziidae, Gongylonematidae, Setariidae, and Onchocercidae, in which the Thelaziidae and Gongylonematidae were basal. The Onchocercidae as a monophyletic group (pp = 1.0) was the last diverged. However, the monophyly of family Thelaziidae was unresolved: the T. callipaeda isolates from different geographical locations generated a single group (pp = 1.0), and the other species Spirocerca lupi plus Gongylonematidae made up a separate clade.

DISCUSSION

Thelaziasis can occur when infected flies of the genus Phortica feed on the ocular secretions, tears, and conjunctiva of hosts.

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**Table 2** Results of conserved sites (Cs), variable sites (Vs), parsim-info sites (Ps), singleton sites (Ss) and test of substitution saturation of aligned sequences of spirurid nematodes.

| Partition | Final length | Cs | Vs | Ps | Ss | Iss* | Iss.c | #P-value |
|-----------|--------------|----|----|----|----|------|-------|----------|
| atp6      | 603          | 68 | 535| 453| 78 | 0.5074| 0.6998| 0.0000   |
| cox1      | 1656         | 645|1011| 845|166 | 0.3036| 0.7820| 0.0000   |
| cox2      | 711          | 183| 528| 442| 83 | 0.3929| 0.7396| 0.0000   |
| cox3      | 783          | 193| 590| 516| 74 | 0.4288| 0.7478| 0.0000   |
| cytB      | 1095         | 381| 714| 596|118 | 0.3527| 0.7652| 0.0000   |
| nad1      | 909          | 284| 625| 495|130 | 0.3838| 0.7537| 0.0000   |
| nad2      | 867          | 142| 725| 564|161 | 0.4853| 0.7451| 0.0000   |
| nad3      | 345          | 79 | 266| 225| 41 | 0.5216| 0.6615| 0.0000   |
| nad4      | 1233         | 352| 881| 700|181 | 0.3961| 0.7721| 0.0000   |
| nad4L     | 243          | 26 | 217| 179| 38 | 0.5893| 0.6048| 0.0000   |
| nad5      | 1599         | 397|1202| 979|223 | 0.4461| 0.7815| 0.0000   |
| nad6      | 471          | 51 | 420| 336| 84 | 0.5625| 0.6901| 0.0000   |
| mlL       | 1003         | 233| 770| 592|178 | 0.3559| 0.7514| 0.0000   |
| mS        | 686          | 202| 484| 393| 91 | 0.3135| 0.6901| 0.0000   |

*Iss indicates index of substitution saturation. #Iss.c denotes critical index substitution saturation for an asymmetrical tree.
In China, since the first case with human thelaziasis was described in 1917, more than 600 cases scattered across almost the entire country have been reported (Wang et al., 2014). Human cases typically occur in rural villages associated with poor health and socioeconomic settings (Shen et al., 2006). Until now, the etiological agent of all human thelaziasis reported in China was identified as *T. callipaeda*, but few cases reported detailed morphological characteristics. In this study, we firstly reported three cases with clinical eyeworms in central China, and we examined these worms under light microscopy and scanning electron microscopy. The key characteristics of the *T. callipaeda* isolates in our study were similar to those of *T. callipaeda* isolates from dogs in southern Italy (Otranto et al., 2003). In addition, the mouth profile of the *T. callipaeda* male was first described in this study. In comparison with the isolates from Italy, the sizes of the parasites collected here were relatively smaller; the males measured 6.4–10.3 mm in length and 307–389 μm in width (7.7–12.8 mm in length and 338–428 μm in width for the Italy isolates), and the females measured 11.2–15.3 mm in length and 311–459 μm in width (12–18.5 mm in length and 370–510 μm for the Italy isolates).

In recent years, the mitochondrial genomes of spirurid nematodes have been extensively studied (Ghedin et al., 2007; Yatawara et al., 2010; Park et al., 2011; McNulty et al., 2012; Liu et al., 2015; Yilmaz et al., 2016). These whole mt genomes will serve as a useful dataset for studying the genetics, systematics and phylogenetic relationships of spirurid nematodes. In accordance with the two known mt genomes of *T. callipaeda* (NC_018363 and AP017700), all *T. callipaeda* isolates from central China contained 12 PCGs, 22 transfer RNA genes, two ribosomal RNA genes and a non-coding region but lacked an *atp8* gene (Liu et al., 2013a). The gene content and arrangement are

(Fuentes et al., 2012).
the same as those of Acanthocheilonema viteae, Brugia species, Chanderella quiscali, Dirolfilaria species, Loa loa, Onchocerca species, Wuchereria bancrofti, Gongylonema pulchrum, Setaria digitata (Ghedin et al., 2007; Yatawara et al., 2010; McNulty et al., 2012; Yilmaz et al., 2016), but distinct from those of Dracunculus medinensis, Philometroides sanguineus, Gnathostoma doloresi, and Helicometra longissimum (Su et al., 2016; Sun et al., 2016). The mt genome sequence of T. callipaeda has high a A+T content (above 70%), which is consistent with the mt genomes of nematodes from the order Spirurida (Yatawara et al., 2010; McNulty et al., 2012). All PCGs had ATG, TTT, ATA, or TTG as their initiation codons, and TAA or TAG as their termination codons, which was consistent with a previous study (Liu et al., 2013a). The sizes of the 22 tRNA genes identified here were similar to those of other spirurid nematodes, indicating their similar functions (Liu et al., 2013b; Yilmaz et al., 2016). The locations of rrnL and rrnS were also identical to previously published mt genomes of Thelazia. The nucleotide sequences of the mtDNA of five T. callipaeda isolated from different geographical locations were conspicuously similar, which indicated that the influence of genetic variations on the phylogenetic distances in T. callipaeda is relatively small (Zhang et al., 2015). The mt genome of the T. callipaeda isolate from Guangdong (NC_018363) was the most divergent of the entire set. The results of sequence divergence of PCGs suggested that nad4L is the most conserved protein-coding gene. Nad6 showed high sequence variability among the five isolates, indicating that this gene is suitable as a genetic marker for population genetic studies of T. callipaeda from different geographic origins.

Although nuclear ribosomal genes (rDNA) have been verified for phylogenetic studies of a wide range of organisms (Nadler et al., 2007; Zhang et al., 2017), in comparison with a single gene or a single region of rDNA, the complete mt genome sequences have significantly improved the power of phylogenetic analysis and made possible more accurate resolution of taxonomic relationships even at deep levels (Brabec et al., 2016). For spirurid nematodes, the mt genomes have been applied particularly to studies regarding phylogeny and evolution (Park et al., 2011; McNulty et al., 2012). To date, there are 24 mitogenomes of the order Spirurida in GenBank representing eight families. Our study brings a new perspective by presenting mitogenomes of the order Spirurida in GenBank representing et al., 2013a, 2015; Yilmaz et al., 2016). To date, there are 24 studies regarding phylogeny and evolution (Park et al., 2011; Liu et al., 2013a, 2015; McNulty et al., 2012, 2016). The mt genome of the T. callipaeda whole mitochondrial genomes of three T. callipaeda isolates from different geographical locations were sequenced using next-generation sequencing methods. The genome featured 12 PCGs, 22 transfer RNA genes, two ribosomal RNA genes and a non-coding region, which was similar to that of most of other spirurid nematodes. The mtDNA nucleotide sequences of these isolates from different locations were conspicuously similar. The nad6 gene showed high sequence variability among all isolates, which is worth considering for future population genetic studies of T. callipaeda. Phylogenetic analyses support the monophyly of T. callipaeda isolates from different hosts and distinct geographical locations. However, the taxonomic position of Spirocerca lupi is still ambiguous.

CONCLUSION

In this study, three cases of thelaziasis were reported in central China, and all clinical worms were identified as T. callipaeda according to several key characteristics examined under light microscopy and scanning electron microscopy. In addition, the whole mitochondrial genomes of three T. callipaeda isolates from different geographical locations were sequenced using next-generation sequencing methods. The genome featured 12 PCGs, 22 transfer RNA genes, two ribosomal RNA genes and a non-coding region, which was similar to that of most of other spirurid nematodes. The mtDNA nucleotide sequences of these isolates from different locations were conspicuously similar. The nad6 gene showed high sequence variability among all isolates, which is worth considering for future population genetic studies of T. callipaeda. Phylogenetic analyses support the monophyly of T. callipaeda isolates from different hosts and distinct geographical locations. However, the taxonomic position of Spirocerca lupi is still ambiguous.

AUTHOR CONTRIBUTIONS

Conceived and designed the experiments: XZ, ZW, and JC. Performed the experiments: XZ, YS, JD, PJ, and RL. Analyzed the data: XZ. Contributed reagents/materials/analysis tools: JC, XZ, and ZW. Wrote the paper: XZ, ZW, and JC. All authors read and approved the final version of the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2017.01335/full#supplementary-material

2https://www.ncbi.nlm.nih.gov/taxonomy
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