Morphology, genetic characterization and molecular phylogeny of pinworm Skrjabinema longicaudatum n. sp. (Oxyurida: Oxyuridae) from the endangered Tibetan antelope Pantholops hodgsonii (Abel) (Artiodactyla: Bovidae)

Yi-Fan Cao¹²†, Hui-Xia Chen³†, Yang Li³, Dang-Wei Zhou¹⁴*, Shi-Long Chen¹⁴ and Liang Li³*

Abstract

Background: The Tibetan antelope Pantholops hodgsonii (Abel) (Artiodactyla: Bovidae) is an endangered species of mammal endemic to the Qinghai-Tibetan Plateau. Parasites and parasitic diseases are considered to be important threats in the conservation of the Tibetan antelope. However, our present knowledge of the composition of the parasites of the Tibetan antelope remains limited.

Methods: Large numbers of nematode parasites were collected from a dead Tibetan antelope. The morphology of these nematode specimens was observed using light and scanning electron microscopy. The nuclear and mitochondrial DNA sequences, i.e. small subunit ribosomal DNA (18S), large subunit ribosomal DNA (28S), internal transcribed spacer (ITS) and cytochrome c oxidase subunit 1 (cox1), were amplified and sequenced for molecular identification. Moreover, phylogenetic analyses were performed using maximum likelihood (ML) inference based on 28S and 18S + 28S + cox1 sequence data, respectively, in order to clarify the systematic status of these nematodes.

Results: Integrated morphological and genetic evidence reveals these nematode specimens to be a new species of pinworm Skrjabinema longicaudatum (Oxyurida: Oxyuridae). There was no intraspecific nucleotide variation between different individuals of S. longicaudatum n. sp. in the partial 18S, 28S, ITS and cox1 sequences. However, a high level of nucleotide divergence was revealed between the new species and its congeners in 28S (8.36%) and ITS (20.3–23.7%) regions, respectively. Molecular phylogenetic results suggest that the genus Skrjabinema should belong to the...
subfamily Oxyurinae (Oxyuroidea: Oxyuridae), instead of the subfamily Syphacidae or Skrabinemiiinae in the traditional classification, as it formed a sister relationship to the genus Oxyuris.

**Conclusions:** A new species of pinworm *Skrabinema longicaudatum* n. sp. (*Oxyurida: Oxyuridae*) is described. *Skrabinema longicaudatum* n. sp. represents the first species of *Oxyurida* (pinworm) and the fourth nematode species reported from the Tibetan antelope. Our results contribute to the knowledge of the species diversity of parasites from the Tibetan antelope, and clarify the systematic position of the genus *Skrabinema*.

**Keywords:** Tibetan antelope, Parasite, Nematoda, Morphology, Genetic data, Phylogeny

100,000–150,000 mature individuals (https://www.iucnredlist.org/species/15967/50192544). This species is listed as “Near Threatened” in the IUCN Red List of Threatened Species™ and also listed as Class I (Endangered in China) National Protected Wild Animal Species in China.

Parasites and parasitic diseases are considered to be important threats in wildlife conservation, as they can potentially impair the health of wildlife, decrease fitness, cause population declines and even contribute to local extinction [3–7]. Parasites are also significant pathogens of the Tibetan antelope [1, 2]. To date, 17 species of ectoparasites and endoparasites have been reported from the Tibetan antelope, including 5 species of oestrid and hippoboscid flies, 7 species of protozoans, 2 species of tapeworms and 3 species of nematodes [1, 2, 8–12].

In the present study, some nematode specimens were collected from the digestive tract of the Tibetan antelope, which were identified morphologically as a new species of the genus *Skrabinema* (*Oxyurida: Oxyuridae*) using light and scanning electron microscopy. The nuclear and mitochondrial DNA sequences, i.e. small subunit ribosomal DNA (*18S*), large subunit ribosomal DNA (*28S*), internal transcribed spacer (ITS) and cytochrome *c* oxidase subunit 1 (*cox1*), were also amplified and sequenced for molecular identification of this species. Moreover, in order to clarify the systematic status of the genus *Skrabinema*, phylogenetic analyses were performed using maximum likelihood (ML) inference based on *28S* and *18S* + *28S* + *cox1* sequence data, respectively.

**Methods**

**Parasite collection**

A Tibetan antelope died naturally in the Hohxil National Nature Reserve, Qinghai Province, China. The digestive tract of this Tibetan antelope was sent to the Key Laboratory of Adaptation and Evolution of Plateau Biota (AEPB), Northwest Institute of Plateau Biology, Chinese Academy of Sciences for examination of parasites. Large numbers of nematode parasites were isolated from the caecum and colon. Specimens were fixed and stored in 5% glycerine plus 70% ethanol until study.

**Morphological observations**

For light microscopical studies, nematodes were cleared in lactophenol. Drawings were made using a Nikon microscope drawing attachment. For scanning electron microscopy (SEM), the anterior and posterior parts of specimens were re-fixed in 4% formaldehyde solution, post-fixed in 1% OsO4, dehydrated via an ethanol series and acetone, and then critical-point dried. Samples were coated with gold and examined using a Hitachi S-4800 scanning electron microscope at an accelerating voltage of 20 kV. Measurements (the range, followed by the mean values in parentheses) are given in micrometers (μm) unless otherwise stated.

**Molecular protocols**

Three female specimens were randomly chosen for molecular analysis. Genomic DNA from each sample was extracted using a Column Genomic DNA Isolation Kit (Shanghai Sangon, China) according to the manufacturer’s instructions. The partial *18S* region was amplified by polymerase chain reaction (PCR) using the forward primer *18S*-F (5′-CGC GAA TRG CTC ATT ACA ACA GC-3′) and the reverse primer *18S*-R (5′-GGG CGG TAT CTG ATC GCC GGC-3′) [13]. The partial *28S* region was amplified by PCR using the forward primer 28S-F (5′-AGC GGA AAA GAA ACT AA-3′) and the reverse primer 28S-R (5′-ATC CGT GTT TCA AGA CGG G-3′) [14]. The ITS-1 region of nuclear rDNA was amplified by PCR using the forward primer SS1 (5′-GGT GAA CCT GCG-3′) and the reverse primer SS2R (5′-AGT GCT CAA TGT GTC TGC AA-3′). The ITS-2 region of nuclear rDNA was amplified by PCR using the forward primer NC13 (5′-ATC GAT GAA GAA CGC AGC-3′) and the reverse primer NC2 (reverse: 5′-TCA GCC GGT GAG ACT ATC TAA-3′) [15]. The partial *cox1* region was amplified by PCR using the forward primer *cox1F2020* (5′-GAG TAC TAA TCA TAA GGA TAT TGG-3′) and the reverse primer *cox1R2020* (5′-ACA TAA ACY TCA GGA TGA CCA-3′), both newly designed in present study. The cycling conditions are as described previously [16]. PCR products were checked on GoldView-stained 1.5% agarose gels and purified with Column PCR Product Purification Kit (Shanghai Sangon, China).
Sequencing was carried out using a Dye Deoxy Terminator Cycle Sequencing Kit (v.2, Applied Biosystems, California, USA) and an automated sequencer (ABI-PRISM 377). Sequences were aligned using ClustalW2. The DNA sequences obtained herein were compared (using the algorithm BLASTn) with those available in the National Center for Biotechnology Information (NCBI) database (https://www.ncbi.nlm.nih.gov).

Phylogenetic analyses
Phylogenetic trees were constructed using maximum likelihood (ML) inference with MEGA X software based on the partial 28S and 18S + 28S + cox1 sequence data, respectively. *Pseudonymsus spirotheca* (Oxyurida: The-lastomatoidea: Pseudonymidae) was treated as the out-group. The ingroup includes the representatives of the Oxyuridae with the 28S and 18S + 28S + cox1 sequence data available in the GenBank database. We used a built-in function in the software MEGA X to select a best-fitting substitution model for the present sequences according to the Bayesian information criterion. The K2 (Kimura 2-parameter) + G model for the 28S sequence data, and the HKY (Hasegawa-Kishino-Yano) + G + I model for the 18S+28S+cox1 sequence data were identified as the optimal nucleotide substitution model, respectively. Nodal support for ML trees were tested using 1000 bootstrap replications, and bootstrap values exceeding 80% were showed in the phylogenetic trees.

Results

Family Oxyuridae Cobbold, 1864

Genus *Skrjabinema* Werestschagin, 1926

*Skrjabinema longicaudatum* n. sp.

Type-host
Tibetan antelope *Pantholops hodgsonii* (Abel) (Artiodactyla: Bovidae: Caprinae).

Type-locality
Hoh Xil Nature Reserve near Wudaoliang (35° 26′ N, 93° 17′ E), Qinghai Province, China.

Type-specimens
Holotype: male (HBNU-N-2020M001L); allotype: female (HBNU-N-2020M002L); paratypes: 9 females (HBNU-N-2020M003L) deposited in the College of Life Sciences, Hebei Normal University, Hebei Province, China; paratypes: 2 males and 100 females (KLAEPB No.019001) deposited in the Key Laboratory of Adaptation and Evolution of Plateau Biota, Northwest Institute of Plateau Biology, Chinese Academy of Sciences, Qinghai Province, China.

Site in host
Caecum and colon.

Prevalence and intensity
A single Tibetan antelope examined with 124 worms.

ZooBank registration
To comply with the regulations set out in Article 8.5 of the amended 2012 version of the *International Code of Zoological Nomenclature* (ICZN) [17], details of the new species have been submitted to ZooBank. The Life Science Identifier (LSID) of the article is urn:lsid:zoobank.org:pub:9194626F-7C3B-445C-BD36-0AF06E39C46F. The LSID for the new name *Skrjabinema longicaudatum* is urn:lsid:zoobank.org:act:3A5AB2D4-5B82-4CBE-8CF8-76B97783694E.

Etymology
The specific epithet is derived from a combination of the Latin words *longus*-(long) and *caudatum*-(cauda), and refers to the unusually long tail in the female of the new species.

Description

General
Small-sized, whitish nematodes. Body cylindrical, maximum width at slightly posterior to mid-body. Cephalic vesicle indistinct in both sexes (Fig. 1a, c). Lateral alae present in both sexes (Figs. 1c, 2a, c, d). Sexual dimorphism prominent in cephalic structure (Figs. 1b, 3a, b, d). Cuticle with remarkable transverse annulations in anterior part of body (Fig. 2d, e). Buccal cavity very small, without cuticular tooth or other ornamentation. Oesophagus divided into short pharynx, cylindrical corpus, indistinct isthmus and ovoid posterior bulb with valves (Fig. 1a). Nerve-ring situated at about 1/4 of total oesophageal length (Fig. 1a). Excretory pore located in body wall depression, posterior to oesophageo-intestinal junction (Figs. 1a, 2a, b). Deirids not observed.

Male
[Based on 3 mature specimens; Figs. 1c, e, g, h, 3d]: Body 1.92–2.85 (2.38) mm long; maximum width 141–151 (146). Oral aperture simple, triradiate, surrounded by three small, more or less triangular lips with small apical median notch (Fig. 3d). Interlabia or
interlabial projections absent (Fig. 3d). Oesophagus 400–454 (427) in total length, representing 15.7–20.9 (17.6)% of body length; pharynx + corpus + isthmus 255–308 (280) long, size of bulb 143–146 (145) \( \times \) 131–132 (132). Nerve-ring at 170–200 (185) and excretory pore at 760–890 (828) from cephalic extremity, respectively. Lateral alae narrow, extending from about level of nerve-ring to anterior region of cloaca (Fig. 1c). Posterior extremity of body distinctly curved ventrally (Fig. 1h). Spicule single, pointed at distal end, 74–81 (76.7) long, representing 3.15–3.85 (3.43) % of body length (Fig. 1e, g, h). Gubernaculum small, well sclerotized, about 48 long (Fig. 1g, h). Phasmids present slightly posterior to cephalic extremity with pointed tip (Fig. 1g, h). Tail 33 long, ending in short finger-like tip (Fig. 1g, h). Phasmids present slightly posterior to cloaca.

Female
[Based on 10 mature specimens; Figs. 1a, b, d, f, i, 2a–i, 3a–c, e, f]: Body 9.92–12.1 (11.1) mm long; maximum width 396–574 (475). Cephalic extremity with three anchor-shaped lips, each lip with 2 triangular lateral lobes not attached to cephalic rim (Figs. 1b, 3a–c). Interlabia digitiform, between lateral lobes of lips (Figs. 1b, 3a, b). Four large cephalic papillae and 2 small amphidial pores present (Figs. 1b, 3a, b). Oesophagus 832–881 (853) in total length, representing 6.98–8.78 (7.75) % of body length; pharynx + corpus + isthmus 634–703 (671) long, size of bulb 168–188 (181) \( \times \) 139–188 (161). Nerve-ring at 198–248 (225) and excretory pore at 1.24–1.92 (1.77) mm from cephalic extremity, respectively. Lateral alae extending from long distance posterior to base of cephalic extremity and ending at about middle of tail (Fig. 2a, c, d, f, h). Vulva a transverse slit, very small, with rudimentary lips observed under SEM, located at 2.97–3.37 (3.16) mm from cephalic extremity, representing 25.1–31.1% (28.7%) of body length (Figs. 1d, 3e). Egg asymmetrical, flattened at one side, embryonated or nonembryonated, thick-shelled, with smooth surface, 40–59 (51) \( \times \) 20–40 (30) \( (n = 20) \) (Figs. 1i, 2i, 3f). Anus with small pre anal lip (Fig. 2g). Tail slender, very long, 2.63–3.13 (2.90) mm, with pointed tip, representing 23.0–27.7 (26.3) % of body length (Figs. 1f, 3f). Phasmids not observed.

Genetic characterization
Partial 18S region
Three 18S sequences of S. longicaudatum n. sp. obtained herein were all 678 bp in length and represent one genotype. There are two species of Skrjabinema with 18S sequence registered in GenBank, namely S. kamosika (AB699690) and Skrjabinema sp. (EF180060). Pairwise comparison of 18S sequences between S. longicaudatum n. sp. and the two species of Skrjabinema displayed 0.29–1.18% nucleotide divergence. The 18S sequences of S. longicaudatum n. sp. are deposited in the GenBank database under the accession numbers MW020179-MW020181.

Partial ITS region
Three ITS sequences of S. longicaudatum n. sp. obtained herein were all 1079 bp in length and represent one genotype. There are two species of Skrjabinema with ITS sequence registered in GenBank, namely S. kamosika (AB699691) and Skrjabinema sp. (AB367796). Pairwise comparison of ITS sequences between S. longicaudatum n. sp. and the other two species of Skrjabinema displayed 20.3–23.7% nucleotide divergence. The ITS sequences of S. longicaudatum n. sp. are deposited in the GenBank database under the accession numbers MW020057-MW020059.

Partial 28S region
Three 28S sequences of S. longicaudatum n. sp. obtained herein were all 819 bp in length and represent one genotype. There is only one species of Skrjabinema, namely S. ovis (KY990019) with 28S sequence registered in GenBank. Pairwise comparison of ITS sequences between S. longicaudatum n. sp. and S. ovis displayed 8.36% nucleotide divergence. The 28S sequences of S. longicaudatum n. sp. are deposited in the GenBank database under the accession numbers MW020098-MW020100.

Partial cox1 region
Three cox1 sequences of S. longicaudatum n. sp. obtained herein was 360 bp in length. There is no species of Skrjabinema with cox1 sequence registered on GenBank. The cox1 sequences of S. longicaudatum n. sp. are deposited in the GenBank database under the accession numbers MW021552-MW021554.

Phylogenetic analyses
Phylogenetic trees based on the partial 28S sequence data showed that representatives of the family Oxyuridae were divided into three monophyletic clades. Clade I included members of the genera Syphacia, Passalarius, Syphatineria, Syphabulae and Rauschtieneria, representing the subfamily Syphacinae. Clade II contained species of the genera Oxyurus and Skrjabinema, representing the subfamily Oxyurinae. Clade III included species of the genus Trypanoxyurus, representing the subfamily Enterobini (Fig. 4). Skrjabinema longicaudatum n. sp. displayed a sister relationship to S. ovis.
Phylogenetic tree constructed based on the 18S+28S+cox1 sequence data had similar topology to the phylogenetic results using the partial 28S sequence data, in which representatives of the Oxyuridae also divided into three monophyletic clades (Fig. 5). Species of *Trypanoxyuris* and *Enterobius* formed clade I,
representing the subfamily Enterobiinae. The members of *Syphabulea* and *Syphacia* grouped together (clade II), belonging the subfamily Enterobiinae. Clade III included representatives of *Oxyuris* and *Skjabinema*, representing the subfamily Oxyurinae. *Skjabinema longicaudatum* n. sp. clustered together with *S. kamosika*.

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**Fig. 2** Scanning electron micrographs of *Skjabinema longicaudatum* n. sp. (Oxyurida: Oxyuridae) from the endangered Tibetan antelope *Pantholops hodgsonii* (Abel) (Artiodactyla: Bovidae) in China. 

- **a** Anterior part of female (lateral ala and depressed region around excretory pore arrowed), lateral view. 
- **b** Magnified image of depressed region and excretory pore (excretory pore arrowed). 
- **c** Magnified image of the original position of lateral ala. 
- **d** Anterior part of female (lateral alae arrowed), dorsal view. 
- **e** Magnified image of transverse annulations in the anterior part of body. 
- **f** Posterior extremity of female (anus arrowed), ventral view. 
- **g** Magnified image of anus. 
- **h** Magnified image of the ending position of caudal ala. 
- **i** Magnified image of eggs in uterus in different views.
The genus *Skrjabinema* Werestschagin, 1926 (Oxyuridea: Oxyuridea) currently includes 10 nominal species reported from various ruminants worldwide, namely *S. alata* Mönnig, 1932, *S. africana* Mönnig, 1932, *S. caprae* Schad, 1959, *S. chubuki* Gagarin & Sapozhnikov, 1968, *S. kamosika* Hasegawa, Sato, Suzuki & Kaneshiro, 2012, *S. ovis* (Skrjabin, 1915), *S. parva* Dikmans, 1942, *S. rupicaprae* Böhm & Gebauer, 1930, *S. skrabinii* Gagarin & Sapozhnikov, 1968 and *S. tarandi* Skrjabin & Mizekowitsch, 1930 [18–24]. However, some of these species have not been sufficiently well described, especially the details of cephalic structure.

*Skrjabinema ovis* is the type-species of this genus, which has been widely reported from goats and sheep in Asia, Europe, America and Australia [21].

**Fig. 3** Scanning electron micrographs of *Skrjabinema longicaudatum* n. sp. (Oxyurida: Oxyuridae) from the endangered Tibetan antelope *Pantholops hodgsonii* (Abel) (Artiodactyla: Bovidae) in China. **a** Cephalic region of female, sub-apical view. **b** Cephalic region of female (amphidial pores arrowed), apical view. **c** Cephalic region of female, lateral view. **d** Cephalic region of male, apical view. **e** Magnified image of vulva. **f** Magnified image of egg, lateral view.
species has been recorded from *Capra aegagrus hircus* (Linnaeus), *Ovis aries* Linnaeus and *Procapra przewalskii* Büchner in China [25]. The new species differs from *S. ovis* in the absence of cephalic vesicle in females (vs the presence of remarkable cephalic vesicle in the female in *S. ovis*), slightly shorter spicules (0.074–0.081 mm long in the new species vs 0.09–0.12 mm long in *S. ovis*) and longer gubernaculum (0.048 mm long in *S. longicaudatum* n. sp. vs 0.019–0.025 mm long in the latter). *Skrabinema longicaudatum* n. sp. has the spicule without a dilated proximal end, longer gubernaculum (0.019–0.025 mm long) and the caudal alae ending about half-way along the tail in female, which is different from that of *S. parva* (the proximal end of spicule extends into a goblet-shaped, the gubernaculum 0.01–0.016 mm long and the caudal alae ending close to the tail tip in female). The absence of sub-interlabial projections in the cephalic region in the male distinguishes the new species from *S. kamosika*, *S. tarandi* and *S. caprae* (the presence of sub-interlabial projections in the cephalic end in the male). Moreover, the caudal alae of the female in *S. tarandi* and *S. caprae* are very long (ending near the tail tip vs caudal alae ending at about half the tail length in the female of *S. longicaudatum*). *Skrabinema longicaudatum* n. sp. differs from *S. rupicaprae* by having a larger body size of male (1.92–2.85 vs 1.54–1.79 mm long in *S. rupicaprae*), a relatively shorter spicule (spicule representing 3.15–3.85 % of body length in the former vs representing 4.47–5.20 % of body length in the latter) and a slightly longer gubernaculum (0.048 mm long in the new species vs 0.025 mm long in *S. rupicaprae*). The new species can be distinguished from *S. chubuki* and *S. skrjabini* by having morphologically different lips in the female (triangular lateral lobes of lip small, not attached to the cephalic rim in *S. longicaudatum* n. sp. vs triangular lateral lobes of lip very large, attached to the cephalic rim in the latter two species).

Mönnig [18] described *S. alata* and *S. africana* based on only female specimens in South Africa. Both species are differentiated from the new species by distinctly smaller body size of female (4.61–5.80 mm long in *S. alata* and *S. africana* vs 9.92–12.1 mm long in *S. longicaudatum* n. sp.). In addition, *S. africana* differs

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**Fig. 4** Maximum likelihood (ML) tree constructed from the partial 28S gene data showing the phylogenetic relationships of representatives of the family *Oxyuridae*. *Pseudonymus spirotheca* (*Oxyuridae: Pseudonymidae*) was chosen as the outgroup.
from *S. longicaudatum* n. sp. by having the caudal alae of the female ending very near the tail tip (vs caudal alae ending at about half of tail length in *S. longicaudatum*). *Skrjabinema alata* differs from the new species by having a relatively longer oesophagus (oesophagus representing 12.8–14.3 % of body length in *S. alata* vs representing 6.98–8.78 % of body length in *S. longicaudatum*). Moreover, *S. longicaudatum* n. sp. can be easily distinguished from all its congeners by the unusually long tail in the female (tail 2.63–3.13 mm, representing 23.0–27.7 % of body length vs not over 1.60 mm, representing 6.68–20.6 % of body length in the other species of *Skrjabinema*).

It is difficult to identify and discriminate the pinworms using traditional methods due to their extraordinary morphological similarity and sometimes the male worms being unavailable [26]. Molecular approaches have been employed for identification and discrimination of pinworms in some previous studies [24, 26–32]. However, to date, genetic data of pinworms available in the GenBank database remain limited, which has hindered the further studies of DNA-based taxonomy, population genetics and phylogenetics of this group of nematode parasites.

In the present study, we amplified and sequenced the partial 18S, 28S, ITS and cox1 alignments of our specimens for future use in the molecular identification of this new species. There was no intraspecific nucleotide variation between different individuals of *S. longicaudatum* n. sp. in the partial 18S, 28S, ITS and cox1 sequences. However, a high level of nucleotide divergence was revealed between the new species and its congeners in 28S (8.36%) and ITS (20.3–23.7%) regions, respectively. The more slowly evolving 18S gene may be not suitable for species identification of *Skrjabinema*, because of very low level of interspecific nucleotide variation detected between different species of *Skrjabinema* (0.29–1.18%). However, the 18S gene could be chosen to provide resolution at higher taxonomic levels. It is the first time to report the cox1 sequence of *Skrjabinema* species.

The systematic position of *Skrjabinema* is still under debate. Skrabin [22] placed this genus into the subfamily Syphaciinae Railliet, 1916 in Syphaciidae Skrabin & Schikhobalova, 1951. Erkulov & Moldopiyazova [33] proposed a new subfamily Skrjabineminae for the genera *Skrjabinema* and *Citellina*. Hugot [34] reduced the family Syphaciidae to a subfamily in Oxyuridae and did not recognise the validity of *Skrjabinema*. The present phylogenetic analyses based on the partial 28S and 18S+28S+cox1 sequence data supported the genus *Skrjabinema* to be a member of the subfamily Oxyurinae, with a sister relationship with the genus *Oxyuris*, which agrees well with recent molecular phylogenetic results [26].

![Fig. 5 Maximum likelihood (ML) tree constructed from the partial 18S+28S+cox1 gene data showing the phylogenetic relationships of representatives of the family Oxyuridae. Pseudonymus spirotheca (Oxyurida: Pseudonymidae) was chosen as the outgroup.](image-url)
Our present knowledge of the composition of the nematode parasites of the Tibetan antelope remains limited. In the light of available literature, only three species of nematodes have been recorded from the Tibetan antelope, including *Nematodirus* sp., *Marshallagia mongolica* Schumakoviech, 1938 and *M. marshalli* (Ransom 1907) (Rhabditida: Strongyloidea) [9–11]. *Skrjabinema longicaudatum* n. sp. represents the first species of Oxyurida (pinworm) and the fourth nematode species reported from the Tibetan antelope.

Conclusions
A new species of pinworm *Skrjabinema longicaudatum* n. sp. (Oxyurida: Oxyuridae) is described using light and scanning electron microscopy, based on specimens collected from the endangered Tibetan antelope. *Skrjabinema longicaudatum* n. sp. represents the first species of Oxyurida (pinworm) and the fourth nematode species reported from the Tibetan antelope. The nuclear and mitochondrial DNA sequences (i.e. 18S, 28S, ITS and *cox*1) were amplified and sequenced for molecular identification of this new species. Phylogenetic analyses using maximum likelihood (ML) inference based on 28S and 18S + 28S + *cox*1 sequence data suggested that the genus *Skrjabinema* should belong to the subfamily Oxyurinae (Oxyuridae), instead of the subfamily Syphaciidae or Skrjabinemiinae in the traditional classification, as it formed a sister relationship to the genus *Oxyuris*. Our results contribute to the knowledge of the species diversity of parasites from the Tibetan antelope, providing useful genetic data for molecular identification and phylogeny of the Oxyuridae, and also clarified the systematic position of the genus *Skrjabinema*.

Abbreviations
AEPB: Key laboratory of adaptation and evolution of plateau biota; SEM: Scanning electron microscopy; K2: Kimura 2-parameter; HKY: Hasegawa-Kishino-Yano; PCR: Polymerase chain reaction; ML: Maximum likelihood; 18S: Small subunit ribosomal DNA; 28S: Large subunit ribosomal DNA; ITS: Internal transcribed spacer; *cox*1: Cytochrome c oxidase subunit 1; L: Lip; i: Interlabia; tl: Triangular lateral lobes of lip; cp: Cephalic papillae.

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Authors’ contributions
YFC, DWZ and SLC carried out sample collection. LL, HXC, DWZ, YFC and YL identified the nematode specimens and analyzed data. LL, HXC, YFC and DWZ designed the study, conducted the phylogenetic analyses and wrote the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials
The nuclear and mitochondrial DNA sequences of *Skrjabinema longicaudatum* were obtained in this study and deposited in the GenBank database under the accession numbers MW021552-MW021554 (*cox*1 sequences), MW020057- MW020059 (ITS sequences), MW020098-MW020100 (28S sequences), MW020179-MW020181 (18S sequences). Type specimens of the new species were deposited in College of Life Sciences, Hebei Normal University, Hebei Province under the accession numbers HBNU-N-2020M001-3L, and the Key Laboratory of Adaptation and Evolution of Plateau Biota, Northwest Institute of Plateau Biology, Chinese Academy of Sciences, Qinghai Province, under the accession numbers KLAEPB No.019001, China.

Ethics approval and consent to participate
This study was conducted under the protocol of the Ethical Commission of the Northwest Institute of Plateau Biology, Chinese Academy of Sciences and Hebei Normal University. All applicable institutional, national and international guidelines for the protection and use of animals were followed.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no conflict of interest.

Author details
1 Key Laboratory of Adaptation and Evolution of Plateau Biota, Chinese Academy of Sciences, Beijing, 100093, People’s Republic of China.
2 College of Life Sciences, Hebei Normal University, Shijiazhuang 050024, Hebei Province, People’s Republic of China.
3 University of Chinese Academy of Sciences, Beijing 100093, People’s Republic of China.

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