Molecular analysis of polymorphic species of the genus *Marshallagia* (Nematoda: Ostertagiinae)

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**Abstract**

**Background:** The genus *Marshallagia* (Family Haemonchidae, subfamily Ostertagiinae) contains multiple species of nematodes parasitising the abomasum (or duodenum) of ruminants, in particular of Caprinae. Male specimens have been described to be polymorphic with the frequent/major morphotype initially described in the genus *Marshallagia* while the minor/rare morphotype was initially often placed in the genus *Grossospicularia*. Due to common morphological features, certain pairs of morphotypes were suggested to belong to the same species such as *Marshallagia marshalli*/*M. occidentalis*. However, molecular evidence to confirm these pairs of morphotypes belonging to the same species is missing.

**Methods:** In the present study, *Marshallagia* sp. were collected from domestic sheep in Uzbekistan. Male specimens were morphologically described with particular emphasis on the structure of the bursa copulatrix. After DNA isolation from morphologically identified specimens, PCRs targeting the ribosomal internal transcribed spacer 2 (ITS2) and mitochondrial cytochrome c oxidase subunit 1 (cox1) regions were conducted. After Sanger sequencing, maximum likelihood phylogenetic analyses and pairwise identities between sequences were calculated.

**Results:** The major morphotypes of *M. marshalli*, *M. schumakovitschi* and *M. uzbekistanica* and the minor morphotypes *M. occidentalis*, *M. trifida* and *M. sogdiana* were identified and their morphology was documented in detail. ITS2 sequences showed little variation and did not allow diagnosing species. In contrast, phylogenetic analysis of cox1 sequences identified highly supported clusters and verified that *M. marshalli*, *M. occidentalis* and *M. uzbekistanica* are different morphotypes of the species *M. marshalli* while *M. schumakovitschi* and *M. trifida* represent distinct morphotypes of *M. trifida*. For *M. sogdiana* no corresponding major morphotype could be identified in the present study. Due to a large barcoding gap, comparison of cox1 sequences in terms of percent identity was sufficient to reliably assign the sequences to a particular species without phylogenetic analysis.

**Conclusions:** The data presented here create a framework that will allow the classification of other members of the genus in the future and underline that parallel morphological and molecular analysis of specimens is crucial to improve the taxonomy of polymorphic species.

**Keywords:** *Marshallagia*, Polymorphic species, Parasitic nematodes, Ruminants, Barcoding gap
**Background**

Nematodes of the genus *Marshallagia* Orloff, 1933 are parasites of the abomasum and duodenum of free-ranging and domesticated ruminants and are most often associated with Caprinae from the Holarctic region [1, 2]. They have a considerable impact on the metabolism of host animals frequently leading to economic losses [3–5]. According to our data, the species of *Marshallagia marshalli* Ransom, 1907 is widespread in Uzbekistan and it is the most frequently detected member of this genus [6].

Drozdz [7, 8] described that morphological dimorphism is common among males of several species of the subfamily Ostertagiinae, i.e. presence of “major” and “minor” morphotypes in the same host. Major morphotypes are named due to their higher frequency relative to minor morphotypes, not due to differences in size. In contrast to males, females are monomorphic, and this situation has often led to the description of distinct species for these male morphotypes.

In the genus *Marshallagia*, five dimorphic species have been described by Drozdz [9]. Later, 12 dimorphic species of *Marshallagia* were differentiated by morphological descriptions incorporating the synlophe in males, the structure of the spicules as well as the genital cone in major and minor morphotype males [2]. In particular, structural characters of the spicules and genital cone were used to distinguish between major and minor morphotypes. The major morphotypes were usually placed in the genus *Marshallagia* while minor morphotypes were often allocated to *Grosspiculagia* Orloff, 1933. Thus, major and minor morphotypes of the same species have historically been described as different nominal species, often in separate genera [9, 10]. However, there are also morphological features that are shared between major and minor morphotypes but can be used to discriminate between species including in particular details of the morphology of esophagus and synlophe as well as the shape of the rays of the copulatory bursa [9]. For instance, *M. occidentalis* Ransom 1907 (minor morphotype) has clearly distinguishable morphological features discriminating it from *M. marshalli* (major morphotype), but there are clear hints that *M. marshalli* and *M. occidentalis* are synonyms and represent only different morphotypes of *M. marshalli*, whereas *M. mongolica* Shumakovitschi, 1950 appears to be a major morphotype while *M. grossipriculum* Li, Yin, Kong & Jang, 1987, which corresponds to *Marshallagia* sp. 1 of Drozdz [9], is most likely a minor morphotype of the same species. The same relationship presumably also applies to *M. shumakovitschi* Kadyrov, 1959 (major morphotype) and *M. trifida* (Guille, Marotel & Penisset, 1911) (= *Marshallagia* sp. 2 of Drozdz [9]) (minor morphotype) [2].

Even less is known about the occurrence of cryptic species in other frequently occurring representatives of the Ostertagiainae, especially in *M. marshalli* due to its presence in several host species [2, 11]. Regarding nomenclature, taxonomy and phylogeny of the genus *Marshallagia*, Hoberg et al. [2] proposed that additional studies are required combining morphological and molecular analysis from individual specimens to confirm descriptions of the major species *M. skrjabini* as well as the minor species *M. belockani* and *M. sogdana*. Moreover, it remains an open question as to whether or not *M. marshalli/M. occidentalis* is distributed in North America and in Eurasia. *Marshallagia marshalli/M. occidentalis* is the only species of the genus with a Holarctic distribution but detailed comparisons between North American and Central Eurasian populations have not been conducted. Therefore, it should be considered that there are two closely related species with more restricted geographic distribution patterns [2] and that reports of identification of *M. marshalli* from Eurasian ruminants are due to misidentification.

Based on studies by Drozdz [9], Hoberg et al. [2] and Wyrobisz et al. [12] listed five polymorphic species in the genus *Marshallagia*: *M. marshalli/M. occidentalis; M. lichtenfelsi/M. lichtenfelsi* f. minor Hoberg Abrams, Piligt & Jenkins, 2012; *M. mongolica/M. grossipriculum; M. schumakovitschi/M. trifida; and M. skrjabini/M. belockani*. However, to date no consensus on the species composition of the genus *Marshallagia* has been obtained.

Within the Strongylida, species distinction supported merely by morphological features is difficult, and requires confirmation by means of molecular methods. For instance, recent data have suggested that *Cooperia spatulata* is just a morphotype of *C. punctata* [13] while there is evidence that the small strongyle morphospecies *Cylicostephanus calicatus* and *C. minutus* are in fact cryptic species complexes of at least two and three genospecies, respectively [14]. The taxonomy of Ostertagiinae is mainly complicated by complex relationship between species and morphotypes in the genus *Teladorsagia*, but complexity may also be expected among other Ostertagiinae (e.g. in the genera *Ostertagia* and *Marshallagia*) [12]. Studies devoted to the problems of taxonomy of strongylid nematodes have shown that non-coding regions of rRNA genes, in ITS1 and particularly ITS2, are well suited to allow taxonomic discrimination and this also applies to members of the Ostertagiinae [15–24]. These studies have led to important insights in the evolution and solved general questions of phylogeny of Ostertagiinae including confirmation of conspecificity for *M. marshalli* and *M. occidentalis* [25]. Furthermore, questions regarding conspecificity of supposed major and
minor morphs and cryptic species within the genus of *Orloffia* Drozdz, 1965 were solved [26]. There is no unanimous opinion about the taxonomic independence and specific composition of the genus *Orloffia*. At the same time, there are several species and genera, for which important taxonomic and phylogenetic questions remain unresolved. Additional molecular analyses of ribosomal and mitochondrial DNA will allow understanding these problems in more depth.

Regarding species identification in terms of barcoding properties, mitochondrial DNA sequences were shown to be superior to ribosomal spacers [27]. Recent work on *Cooperia* spp. (Cooperiidae, Cooperinae) and *C. minutus* (Strongylidae, Cyathostominiae) revealed that combined analyses of mitochondrial and nuclear marker sequences improved species identification and phylogenetic analyses [13, 14].

This study aimed to provide sequence data on mitochondrial cytochrome *c* oxidase subunit 1 (*cox1*) and nuclear ribosomal intergenic spacer 2 (ITS2) DNA for some species or morphotypes of the genus of *Marshallagia*. The objective was to morphologically identify individual *Marshallagia* sp. specimens from Uzbekistan to the species and morphotype level followed by obtaining molecular ITS2 and *cox1* data from the same specimens in order to clarify the taxonomic status of the local *Marshallagia* species/morphotypes.

**Methods**

**Parasite collection and examination**

All parasite material was collected at necropsy from domestic sheep (*Ovis aries*) from farms in the Kitob district (Kashkadarya region) and in the Shofirkon district (Bukhara region) in Uzbekistan. Mature *M. marshalli* and *M. occidentalis* worms were collected from the mucosa of the abomasum of domestic sheep (*Ovis aries*) in Kashkadarya region (June 2016) while *M. schumakovitschi*, *M. trifida*, *M. sogdiana* and *M. uzbekistanica* were collected from abomasum of sheep in Bukhara region (July 2016). Male specimens were manually cut in two parts: the posterior region was cleared in phenol alcohol (80% melted phenol and 20% ethanol) for examination of the morphological features and the anterior and middle parts were fixed in 70% ethanol for the molecular studies (Table 1).

**Morphological identification**

*Marshallagia* species were identified according to morphological and morphometrical characters using literature data [1, 2, 9]. All adult male worms isolated from each sheep were morphologically analyzed to identify parasite species. The species identification was established based on caudal bursa according to the features proposed by above literature, especially morphological characters and measurements of spicules, dorsal ray and gubernaculum (Table 1).

An equal mixture of lactic acid and glycerin was used to enlighten the posterior part of the studied nematodes without additional staining. Also included in the present paper are morphological studies made by the authors analyzing specimens (paratypes or sifotypes) of species (*M. marshalli* Ransom, 1907 and *M. schumakovitschi* Kadyrov, 1959) in the Central Helminthological Museum FGGNU, Russian Institute of Parasitology, Animal and Plant named after K. I. Skrjabin, Moscow. For this purpose, a microscope ML 2000 equipped with a digital camera (Meiji, Saitama, Japan) was used.

**DNA extraction**

For DNA isolation, at least a single specimen of each species/morphotype was used. Before isolation of genomic DNA, the ethanol was removed and the adult nematodes were washed with sterile water and DNA was extracted using the NucleoSpin® Tissue Kit (Macherey-Nagel, Düren, Germany) in accordance with the manufacturer’s

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**Table 1** Specimens of *Marshallagia* species collected from hosts in Uzbekistan and additional paratype males from the Central Helminthological Museum (Moscow, Russia) for morphological analyses

| Parasite species          | Locality                  | No. of specimens | Collection    |
|---------------------------|---------------------------|------------------|---------------|
| *Marshallagia marshalli*  | Kashkadarya district, Uzbekistan | 26 ♂ | CPIZ 10270a    |
| *M. occidentalis*         | Kashkadarya district       | 22 ♂ | CPIZ 10278     |
| *M. schumakovitschi*      | Bukhara district, Uzbekistan | 17 ♂ | CPIZ 10271     |
| *M. trifida*              | Bukhara district           | 18 ♂ | CPIZ 10291     |
| *M. sogdiana*             | Bukhara district           | 12 ♂ | CPIZ 10293     |
| *M. uzbekistanica*        | Bukhara district           | 5 ♂  | CPIZ 10292     |
| *Marshallagia marshalli*  | Volgograd district, Russia  | 2 ♂  | CHM 14768b     |
| *M. schumakovitschi*      | Osh district, Kirgizstan    | 2 ♂  | CHM 22289      |

*a* CPIZ - Collection of the Parasitology Institute of Zoology Uzbekistan Academy of Sciences, Uzbekistan  
*b* CHM - Central Helminthological Museum Russian Institute of Parasitology, Animal and Plant named after K.I. Skrjabin, Russia
The DNA was eluted with 50 µl elution buffer provided in the kit and stored at −20 °C until further use. The extracted DNA was quantified on a Take3 plate in an Epoch plate reader (Biotek, Berlin, Germany).

**PCR and cloning**

PCRs were conducted using (i) a combination of the forward and reverse primers flanking the complete ITS2 region [28] and (ii) a partial cox1 gene fragment [29] (Table 2). PCR reactions contained 0.2 mM dNTPs, 250 nM of each primer, 0.4 U Phusion Hot Start II High-Fidelity DNA polymerase (Thermo Fisher Scientific, Darmstadt, Germany) and 2 µl template DNA in 20 µl 1× HF buffer. PCRs were performed on a C1000 or S1000 PCR cycler (Bio-Rad, Feldkirchen, Germany). PCR products were purified using DNA Clean & Concentrator®-5 (Zymo Research, Freiburg, Germany) and amplification products were analyzed by agarose gel electrophoresis in 1.0–1.5% agarose gels. Purified fragments were ligated into the StrataClone Blunt PCR Cloning Vector pSC-B-amp/kan (Agilent, Waldbronn, Germany) and transformed into StrataClone SoloPack competent *Escherichia coli* cells according to the manufacturer’s protocol. Plasmid DNA was purified using the EasyPrep1 Pro kit (Biozym, Hessisch Oldendorf, Germany) and sent for sequencing to LGC Genomics (Berlin, Germany).

**Sequence comparisons and phylogenetic analyses**

Sequences from the present study were analyzed together with sequences previously deposited in GenBank. As an outgroup, two sequences per gene from the species *Teladorsagia circumcincta*, were included. Accession numbers of all sequences from specimens investigated in the present study are provided in Table 3. The ITS2 and cox1 sequences were aligned using MAFFT (multiple sequence alignment using fast Fourier transformation) in the Q-INS-I modus that takes predicted RNA secondary structures into account [30] and the M-COFFEE modus of T-Coffee (Tree-based Consistency Objective Function for alignment Evaluation) [31], respectively. The cox1 alignment was manually edited to ensure that codons were not interrupted by gaps. For calculation of relative identity (%) between sequences, alignments were analyzed using the dist.dna function in the ape 4.0 (Analyses of Phylogenetics and Evolution) package [32] in R 4.0.0 statistics software [33]. Identities were calculated as “raw” identities and pairwise deletion of positions with gaps was turned on. Comparisons of sequences within the genus *Marshallagia* were sorted into the individual intraspecies comparisons and a single category containing all interspecies comparisons. The identity in percent for all these comparisons within the genus *Marshallagia* were compared using the Kruskall-Wallis test followed by a Conover-Iman post-hoc test with the function posthoc.kruskal.conover.test as implemented in the R package PMCMR 4.3 [34]. All P-values below 0.05 were considered to be statistically significant. Scatter plots were visualized using GraphPad Prism 5.03 (GraphPad, La Jolla, USA).

Phylogenetic analyses were conducted on a single gene level. First, substitution saturation tests were conducted according to Xia et al. [35] using DAMBE 5 (Data Analysis in Molecular Biology and Evolution) software [36]. DAMBE 5 was also used to split the cox1 alignment into

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**Table 2** Primers and PCR conditions used for molecular analyses of nematodes

| Primer      | Sequence (5’–3’) | Initial denaturation | Denaturation/annealing/extension | Final elongation |
|-------------|------------------|----------------------|----------------------------------|-----------------|
| NC1         | ACGTCTGTGTCAGGGTGTTC | 98 °C for 30 s       | 40×: 98 °C for 10 s; 55 °C for 30 s; 72 °C for 30 s | 72 °C for 10 min |
| NC2         | TTAGTTTCTTTCCTCCGCT |                      |                                  |                 |
| COI_Nema_Fw | GAAAGGTCTTATCATAGATATGG | 95 °C for 2 min      | 35×: 95 °C for 1 min; 48 °C for 1 min; 72 °C for 1 min | 72 °C for 5 min |
| COI_Nema_Rv | ACCTCAGGATGACCAAAAAAYCAA |                      |                                  |                 |

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**Table 3** Cytochrome c oxidase subunit 1 (cox1) and second internal transcribed spacer (ITS2) sequences of *Marshallagia* species from Uzbekistan and the GenBank database from different geographical origins used in this study

| Species                  | Voucher | GenBank ID | ITS2  |
|--------------------------|---------|------------|-------|
| *Marshallagia marshalli* | M2      | MT116991   | MT110920 |
| *M. marshalli*           | M12     | MT116992   | MT110919 |
| *M. occidentalis*        | M14     | MT116997   | MT110967 |
| *M. schumakovitschi*     | M3      | MT116993   | MT110926 |
| *M. schumakovitschi*     | M6      | MT116994   | MT110928 |
| *M. schumakovitschi*     | M8      | MT116995   | MT110929 |
| *M. schumakovitschi*     | M10     | MT116996   | MT110927 |
| *M. trifida*             | M5      | MT116998   | MT110827 |
| *M. trifida*             | M9      | MT116999   | MT110828 |
| *M. sogdiana*            | M4      | MT117000   | MT110824 |
| *M. sogdiana*            | M7      | MT117001   | MT110825 |
| *M. sogdiana*            | M13     | MT117002   | MT110826 |
| *M. uzbekistanica*       | M1      | MT116990   | MT110829 |
one partition for the first and second and another partition for the third codon position. Maximum likelihood phylogenetic trees were calculated using IQ-TREE [14] on the IQ-TREE server (http://iqtreet.cibiv.univie.ac.at). Using the ModelFinder option of IQ-TREE [37], automatic determination of the best model applying the Bayesian information criterion was performed including models with FreeRate heterogeneity. Ultrafast bootstrapping (1000 bootstrapped replicates) [38] and the Shimodaira-Hasegawa approximate likelihood ratio test (SH-aLRT) (1000 replicates) [39] were used to obtain node support statistics. The command line in IQ-TREE for ITS2 sequences was: iqt -s infile.fas -st DNA -m TESTNEW -bb 1000 -alrt 1000. For cox1 sequences, separate models were fitted for codon positions 1 and 2 vs codon position 3 using the command line: iqt -s COI_FcC_infile.fas -sp partition_file.txt -pre infile.fas -m TESTNEW -bb 1000 -alrt 1000. Phylogenetic trees were visualized in FigTree 1.1.4 and further edited in CorelDraw 20.

Results and discussion
Morphological identification
Six morphotypes of Marshallagia species were found within the present material. Major and minor Marshallagia species were isolated from domesticated sheep in Uzbekistan, separated according to their morphological identification (Fig. 1) and morphometric comparison (Table 4) and assigned to M. marshalli, M. schumakovitschi, M. uzbekistanica, M. occidentalis, M. trifida and M. sogdiana. As detailed above, Drozdz [7, 8] described the phenomenon of regular co-occurrence of rather rare (minor) species in pairs with the more numerous (major) species, which, together with subtle morphological features, lead to the hypothesis that they represent different morphotypes of the same species. Based on this hypothesis, the analyzed specimens of Marshallagia were divided into six separate morphotypes that were grouped into three species with co-existing morphotypes (Fig. 1). The original micrographs of the specimens that were used for molecular analysis are presented for one exemplary individual per morphotype: M. marshalli (Fig. 1a); M. schumakovitschi (Fig. 1b); M. uzbekistanica (Fig. 1c); M. occidentalis (Fig. 1d); M. trifida (Fig. 1e); and M. sogdiana (Fig. 1f). A detailed morphological description of the different morphotypes is given in Additional file 1: Text S1.

The most reliable characters for differentiation among species of Marshallagia, and specifically of the major morphotypes of the respective species, include the placement of the trifurcation of the spicule tips, the form of the dorsal and ventral processes (relative length, curved or straight), and the chitinized structure of the tip of respective processes.

The three major morphotypes designated as M. marshalli, M. schumakovitschi and M. uzbekistanica (Fig. 1a–c) differ by the distal ends of spicule processes, strongly curved for the first one and with tubercles for the second and third species. All three major morphotypes are characterized by the absence of a gubernaculum (Fig. 1a–c).

Marshallagia marshalli (Ransom, 1907) is the type-species for the genus. Ransom [40, 41] described this species and the minor morphotype, M. occidentalis, based on specimens in domesticated sheep (O. aries L.) from North America. In specimens of the major morphotype of M. marshalli the spicules are strongly curved in lateral view, 210–310 µm in length, 42–54 µm in width; eyelet at trifurcation prominent; with dorsal and ventral process nearly equal in length; gubernaculum absent or strongly chitinized (Fig. 1a). The M. marshalli morphotypes studied by us corresponded to the morphology of the parasite type of M. marshalli (CHM 14768) (Table 1).

Marshallagia schumakovitschi differs from M. marshalli in the structure of the spicule. The most distal sixth part of the spicules is divided into three processes.
Dorsal and ventral processes are nearly equal in length, the ventral process terminates in a simple tip, which may be bent; the dorsal process is weakly sclerotized, blunt and not strongly recurved and extends to near the termination of the main shaft (Fig. 1b).

Marshallagia uzbekistanica is a morphologically atypical form placed in the genus Marshallagia in the original description. Spicules are asymmetric (Fig. 1c). The characteristic of M. uzbekistanica turned out to be a peculiar structure of the spicules, which are weakly chitinized at the proximal end and granular in structure [42].

Minor morphs represented in the present study by the morphotypes M. occidentalis, M. trifida and M. sogdiana correspond to the diagnosis of the genus Grosspiculagia, which is now considered to be a synonym of Marshallagia. In contrast to the major morphs, all minor morphs have thick spicules that are split into three processes. The two more massive processes have cap-shaped distal ends and sometimes a hook-like outgrowth. In contrast to the major morphs, all minor morphs have a transparent, sometimes subtle gubernaculum (Fig. 1d–f).

Marshallagia occidentalis represents the minor morphotype of M. marshalli [9]. Near the middle of the spicule length, spicules are divided into three processes: two ventral and one dorsal. A gubernaculum is present but to the rear its diameter is strongly reduced (Fig. 1d).

Marshallagia trifida is the minor morphotype of M. schumakovitschi and can be identified based on its spicule structure [2, 9]. The ventral process of the spicules is strongly curved; the dorsal process extends to the tip of the main shaft of the spicule. The gubernaculum is fusiform (Fig. 1e).

Marshallagia sogdiana is the minor morphotype of M. skrjabini Asadov, 1954 and was transferred to that species as a new combination of morphs [2]. The proximal ends of the spicules are characterized by the presence of a peculiar, disc-like structure (Fig. 1f). In the middle of the spicule it is divided into three processes: two ventral and one dorsal. The distal spicules have a membrane in the form of a sheath. A gubernaculum is present.

Thus, based on morphological characters and morphometric comparison of males of Marshallagia sp., two pairs of major and minor morphotypes were identified: M. marshalli/M. occidentalis, M. schumakovitschi/M. trifida while for the pair M. skrjabini/M. sogdiana only the minor morphotype was found (Table 4). In addition, M. uzbekistanica was identified as an unusual major morphotype. The data on the morphology of these species from samples collected in Uzbekistan presented here closely corresponds to the previously published data by Asadov [43], Ivashkin [1] and Hoberg et al. [2]. Herein, to scrutinize the identity of the morphs of the genus Marshallagia, molecular studies were conducted in particular to confirm which major and minor morphotypes belong to the same species.

### Table 4

Morphometric data for the male specimens (major and minor morphotypes) of the morphospecies of Marshallagia in domestic sheep from Uzbekistan, based on new observations during the current study.

| Characters                              | M. marshalli (n = 26) | M. occidentalis (n = 22) | M. schumakovitschi (n = 17) | M. trifida (n = 18) | M. sogdiana (n = 12) | M. uzbekistanica (n = 5) |
|-----------------------------------------|-----------------------|--------------------------|-----------------------------|---------------------|----------------------|--------------------------|
| Body length                             | 680–1400 (1050 ± 39)  | 1000–1600 (1330 ± 51)    | 1140–1380 (1190 ± 73)       | 830–17000 (1396 ± 88) | 940–1850 (1355 ± 177) | 734–1586 (1240 ± 28)     |
| Body width                              | 125–257 (207 ± 9)     | 180–260 (210 ± 8)        | 112–201 (175 ± 4)          | 131–212 (170 ± 9)   | 129–235 (169 ± 21)    | 294–428 (340 ± 8)        |
| Diameter of anterior region             | 11–26 (21 ± 1)        | 36–46 (41 ± 1)           | 21–33 (26 ± 3)             | 41–62 (50 ± 3)      | 39–65 (47 ± 4)        | 17–23 (21 ± 4)           |
| Distance from cervical capsule to anterior extremity | 292–491 (381 ± 13) | 371–479 (414 ± 8)       | 391–455 (405 ± 15)         | 337–581 (424 ± 28)  | 165–215 (188 ± 9)     | 401–448 (429 ± 10)       |
| Distance from nerve-ring to anterior extremity | 239–391 (318 ± 7) | 291–416 (350 ± 10)      | 220–388 (290 ± 12)         | 251–411 (320 ± 10)  | 247–395 (295 ± 12)    | 151–221 (173 ± 9)        |
| Esophagus length                        | 211–319 (242 ± 18)   | 751–991 (890 ± 20)      | 781–957 (860 ± 20)         | 672–932 (774 ± 31)  | 719–793 (748 ± 23)    | 65–96 (78 ± 9)            |
| Esophagus width                         | 26–39 (33 ± 3)        | 62–93 (71 ± 5)           | 58–87 (65 ± 4)             | 59–72 (62 ± 3)      | 53–68 (59 ± 4)        | 63–74 (68 ± 4)            |
| Spicule length                          | 211–311 (250 ± 10)   | 218–373 (280 ± 9)       | 218–301 (240 ± 10)         | 241–391 (295 ± 9)   | 254–383 (284 ± 15)    | 102–147 (124 ± 7)         |
| Spicule width                           | 42–54 (48 ± 3)        | 52–61 (57 ± 2)           | 53–59 (53 ± 3)             | 43–57 (49 ± 4)      | 47–51 (48 ± 5)        | 38–46 (41 ± 3)            |
| Dorsal ray length                       | 321–483 (370 ± 11)   | 139–317 (200 ± 20)      | 251–375 (300 ± 10)         | 231–338 (259 ± 11)  | 223–315 (253 ± 11)    | 223–378 (241 ± 11)        |
| Bursa length                            | 285–452 (360 ± 8)    | 653–746 (690 ± 8)       | 311–415 (357 ± 10)         | 611–884 (709 ± 12)  | 593–875 (745 ± 12)    | 278–548 (366 ± 13)        |
nuclear ITS2 rRNA (321–325 bp excluding the primers) and mitochondrial partial cox1 (696 bp) genes were amplified, cloned and sequenced. All sequences were deposited in GenBank under the accession numbers provided in Table 3.

### Internal transcribed spacer 2

Identity between all ITS2 sequences of the genus *Marshallagia* from the present study (*n* = 13) or from GenBank revealed between 89.7% and 100% identity with 85% of the pairwise comparisons showing > 95% identity. A maximum likelihood phylogenetic tree calculated from ITS2 sequences from the present study plus those available in GenBank using two representative *T. circumcincta* sequences as an outgroup is shown in Additional file 2: Figure S1. The phylogram reveals that the ITS2 sequence contains virtually no phylogenetic signal since (i) there are barely any clusters of sequences showing high statistical support, (ii) sequences assumed to come from different species were virtually identical and (iii) the sequences assigned to the same species are found scattered all over the tree. This indicates that ITS2 sequences are not suitable to address taxonomic or even phylogenetic questions within the genus *Marshallagia*. This is in agreement with other recent studies analyzing closely related Strongyloidea nematodes showing that ITS2 is an excellent marker to identify the genus, but that closely related species differ only minimally in their ITS2 sequence and that the phylogenetic signal obtained from ITS2 sequences of *Marshallagia* specimen was not reliable [13, 14].

### Cytochrome c oxidase subunit 1 gene

The phylogenetic tree calculated from cox1 sequences from the present study or downloaded from GenBank identified seven highly supported clusters (named I-VII from basal to distal operational taxonomic units (OTUs) in Fig. 2) with very low variability within the clusters. A maximum likelihood phylogenetic tree calculated from cox1 sequences from the present study plus those available in GenBank was included since data are annotated as *Marshallagia lanceata* in GenBank. Branch support is presented with results of the rapid bootstrap analysis before and of the Shimodaira-Hasegawa approximate likelihood ratio test behind the slash. Seven highly supported clusters (indicated by Roman numbers I-VII from basal to distal) are considered valid species: *O. lanceata* (I); *M. sogdiana* (II); *M. trifida* (III) including the morphotype *M. schumakovitschi*; the presumably misidentified *M. occidentalis* from China (IV); *M. hsui* (V); *M. marshalli* (VI) including the morphotypes *M. occidentalis* and *M. uzbekistanica* from Uzbekistan; and *M. mongolica* (VII) including the morphotype *M. grossospiculum* and a presumably misidentified *M. marshalli* from China. If more than one morphotype is present in a cluster, the valid species name is printed in bold. Numbers after species names show voucher designations from Lv et al. [44] while M1–M14 are voucher designations from the present study. Abbreviations: CN, China; UZ, Uzbekistan; ?, apparently misidentified specimens.
identities between 93.7% and 100% while all interspecies comparisons are in the range of 84.2–89.6%. Comparisons between different morphotypes of the same species are in the same range as comparisons between identical morphotypes. Since ranges of intra- and inter- species comparisons are not overlapping (barcoding gap), a simple calculation of identities between sequences will be sufficient to assign a sequence to a particular species allowing diagnosis without phylogenetic reconstruction.

**Polymorphic Marshallagia species**

The data presented here clearly show that *M. marshalli* is a polymorphic species and that *M. occidentalis* and *M. uzbekistanica* should be considered synonyms and that their descriptions detail the morphology of rare morphotypes of *M. marshalli*. In addition, the morphotypes *M. schumakovitschi* and *M. trifida* belong to the same species. Although *M. schumakovitschi* represents the major morphotype, this name should be considered a synonym of *M. trifida* since *M. trifida* was described earlier and this name has therefore priority according to the rules of zoological nomenclature. Data from Lv et al. [44] furthermore suggest that *M. grossospiculum* is a minor morphotype of the species *M. mongolica* and only the latter represents a valid species name. However, absence of morphological data in the publication by Lv et al. [44] suggests that further confirmation is warranted.

It remains unclear whether all *Marshallagia* species are polymorphic but the new data presented here confirm certain pairs of morphotypes that must be considered to be the same species. For other species, the availability of cox1 sequences will allow to group more morphotypes once new sequences from morphologically identified and documented material become available.
**Marshallagia** DNA sequences from specimens of unclear origin

The phylogenetic tree in Fig. 2 contains several GenBank entries from samples of doubtable origin. In particular, there is no available description for *M. hsiui* and the name is considered to be a *nomen nudum* [2]. The sequences obviously belong to a *Marshallagia* species but without a detailed, publicly available morphological description, it is impossible to decide whether they represent an undescribed species or if they should be assigned to one of the many described species for which no *cox*1 reference sequences are currently available.

The *M. occidentalis* specimens from China for which no morphological description was provided by Lv et al. [44] have presumably been identified incorrectly. The sequences apparently represent *Marshallagia* sequences but the assignment to *M. occidentalis* is in contrast to all data summarized by Hobberg et al. [2] and to the results of the present study, which all consider *M. occidentalis* as a minor morphotype of *M. marshalli*. Without voucher material deposited in a museum, the taxonomic value of these sequences will remain very limited and only after identical/highly similar sequences will be reported with a detailed morphological description and deposited voucher material, the sequences will be of epidemiological and taxonomic value.

**Conclusions**

Species limits remain poorly defined within the genus *Marshallagia* (and presumably other, particularly polymorphic Ostertaginiae as well) where subtle morphological differences, high morphological or genetic variability, incomplete descriptions, and circumscribed differential diagnoses hinder identification [2, 43, 46, 47]. It was shown in the present study that at a number of described and named taxa from Eurasia are in fact only synonyms of previously established major or minor morphotypes. For holarctic species such as *M. marshalli* with its different morphotypes, it would be highly interesting to compare Eurasian and North American specimens regarding mitochondrial genotypes to determine if there is further population genetic structuring or even different genospecies on both continents. In any case, free access to species descriptions, including approaches to overcome language barriers such as very limited accessibility of original parasitic nematode descriptions in Russian for western parasitologists and *vice versa*, would be required to improve the situation. Missing of accurate and detailed figures and difficult access to representative type-specimens is especially problematic and complicates the possibility of complete and direct comparisons among otherwise similar species and respective morphotypes [2]. Accordingly, comprehensive revision of the genus *Marshallagia* appears warranted but is currently unfortunately unrealistic.

**Supplementary information**

**Supplementary information** accompanies this paper at https://doi.org/10.1186/s13071-020-04265-1.

**Additional file 1:** Text S1. Detailed morphological description of the *Marshallagia* species and morphotypes.

**Additional file 2:** Figure S1. Maximum likelihood phylogenetic tree calculated based on ITS2 sequences. *Teladorsagia circumcincta* sequences were used as the outgroup. Rapid bootstrapping and Shimodaira-Hasegawa approximate likelihood ratio test results are shown before and after the slash, respectively. M1-M14 indicate voucher designations from the present study.

**Abbreviations**

rRNA: ribosomal RNA; ITS1: internal transcribed spacer 1; ITS2: internal transcribed spacer 2; *cox*1: cytochrome c oxidase subunit 1 gene.

**Acknowledgments**

The authors thank Sabrina Ramünke for constant assistance provided regarding molecular studies during the whole period of work.

**Authors' contributions**

AK conceived and designed the study and prepared the first draft of the manuscript. OA and RK collected specimens for the study. KS carried out morphometric analysis of species. AK and OA performed the morphological description. AK carried out the molecular work and JK completed the bioinformatical and statistical analysis. JK and GVS oversaw the study and manuscript preparation. All authors read and approved the final manuscript.

**Funding**

Open access funding provided by Projekt DEAL. This study has been carried out within the framework of the DEUTSCHER AKADEMISCHER AUSTAUSCHDIENST (DAAD)—funding program 57313677, Bilateral Exchange of Academics, 2017, that supported the visit of AK to the Institute of Parasitology and Tropical Veterinary Medicine, Freie Universität Berlin, Germany as a host, and part of the study was supported by project number FA-A8-TO04, Institute of Zoology, Uzbekistan Academy of Sciences.

**Availability of data and materials**

All data generated or analyzed during this study are included in this article and its additional files. Sequence data were deposited in the GenBank database under the accession numbers listed in Table 3.

**Ethics approval and consent to participate**

The parasite samples used in the study were collected from sheep, when the animals were slaughtered for meat and we obtained the specimens from the abomasum and intestines with the permission of the farm owners.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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Received: 28 March 2020 Accepted: 29 July 2020

Published online: 12 August 2020
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