Genetic characteristics of *Giardia duodenalis* from sheep in Inner Mongolia, China

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**Abstract** – *Giardia duodenalis* is an important zoonotic pathogen for both human and animal health. Although there have been reports on *G. duodenalis* infections in animals all over the world, information regarding the prevalence and genetic characteristics of *G. duodenalis* in sheep in Inner Mongolia, China, is limited. In this study, 209 sheep fecal specimens were collected in this autonomous region. We established that the prevalence of *G. duodenalis* was 64.11% (134/209), as determined using nested PCR detection and sequences analysis of the small subunit ribosomal RNA (SSU rRNA) gene. Based on the beta-giardin (*bg*) locus, the glutamate dehydrogenase (*gdh*) locus, and the triose phosphate isomerase (*tpi*) locus to study genetic characteristics, both assemblages A (2.99%, 4/134) and E (97.01%, 130/134) were found. Five novel nucleotide sequence of assemblage E were detected, two at the *bg* locus, two at the *gdh* locus, and one at the *tpi* locus. Multilocus genotyping yielded four assemblage E and two assemblage A multilocus genotypes (MLGs), including four novel assemblage E MLGs and one novel assemblage A MLG. Results of this study indicated that *G. duodenalis* was highly prevalent in sheep in Inner Mongolia. This study is the first to use the multilocus genotyping approach to identify *G. duodenalis* in sheep from this region.

**Key words:** Inner Mongolia, Sheep, *Giardia duodenalis*.

**Introduction**

*Giardia duodenalis* (also known as *G. lamblia* or *G. intestinalis*) is a common intestinal parasite that is wide-spread among vertebrate hosts, including humans, livestock, and wildlife, worldwide [8, 34]. *Giardia duodenalis* infections often remain asymptomatic, but can cause severe diarrhea and chronic disease in humans [6, 16, 24]. Investigations and case reports on *G. duodenalis* infections in humans are common in China. The large number of epidemiological investigations...
conducted at the start of this century suggested that the average infection rate was 0.85% (197/23,098) [18], with the highest infection rate (9.46%, 7/74) reported by one study carried out in a pediatric hospital in China [32]. As sheep have been found to have unexpectedly high levels of infection, they have long been considered a potential reservoir for human infections [10, 25, 31].

Extensive analysis of protein and DNA polymorphisms have long been considered findings indicating that G. duodenalis is a species complex, whose members show little variation in their morphology, and the major genetic groups are now described as assemblages (may correspond to distinct species) [30]. Studies have shown that G. duodenalis can be sub-classified into at least 8 genetically different assemblages (A–H) [29], of which assemblage A and assemblage B are considered to be zoonotic, while the remaining assemblages (C–H) seem to be host-specific. However, in recent studies, assemblage C, D, E and F has been found in a few human cases [1, 7, 26, 37]. Studies on sheep have identified a predominance of G. duodenalis assemblage E, while assemblage A occurred infrequently [10, 25, 27, 31] and assemblage B was rarely found [5, 23].

For the past few years, the reported infection rate with G. duodenalis in sheep and goats in China was 6.07% (418/6890) [18]. Among these, almost all the cases of G. duodenalis infections in sheep were caused by assemblages E and A, with assemblage E being particularly prevalent. However, there are few reports on G. duodenalis infection rates in goats and sheep in Inner Mongolia [36, 40].

In recent years, multilocus genotyping (MLG) of the beta-giardin (bg), glutamate dehydrogenase (gdh) and triose phosphate isomerase (tpi) loci has increasingly been used to characterize G. duodenalis infections in humans and animals [4, 11, 33]. This method has been favored because PCR assays targeting these loci have been shown to have different sensitivities, and occasionally different genotyping results [19, 22]. However, most earlier studies characterized G. duodenalis in sheep using individual loci, and thus far, there are only a few reports on genotyping G. duodenalis from sheep in China using MLG analysis [39].

The Ordos fine-wool sheep is a unique breeding animal in Wushen Banner, Ordos City, Inner Mongolia Autonomous Region. Its fur and meat have extremely high economic benefits and are the main local economic animals, playing an important role in animal husbandry in this area. Giardia duodenalis has influence on the growth and development of sheep, which may in turn affect the economic benefits for local farmers [2]. However, data on G. duodenalis infection in Ordos fine-wool sheep are rare.

The objectives of this study were to investigate the distribution of G. duodenalis assemblages/genotypes in sheep in Inner Mongolia based on MLG analysis, and analyze their genetic characteristics, assess the zoonotic transmission risk, and elucidate the public health significance of this protozoan parasite.

**Materials and methods**

**Ethics statement**

This study was performed in accordance with the recommendations of the Guide for the Care and Use of Laboratory Animals (Publication Year: 2010, ISBN: 9780309154000). The research protocol was reviewed and approved by the Research Ethics Committee of Henan Agricultural University (approval no. LVR1AE2018-007). Permission was obtained from the farm owners before fecal sample collection. In this study, all fecal samples were carefully collected from the rectum of each sheep without causing discomfort.

**Sample collection**

The Inner Mongolia Autonomous Region straddles three major regions of northeast China, north China and northwest China. The area has a plateau-type geology, with a complex and diverse temperate continental monsoon climate. Inner Mongolia makes good use of its local geographical and meteorological features, and is the largest grassland pastoral area in China, with animal husbandry making an important economic contribution.

To study the infection rate and aggregation distribution of G. duodenalis in this animal population, we selected the representative commercial farm at our study site that has the highest intensity of Ordos fine-wool sheep (>3 months old). There were no symptoms of diarrhea in the flock during sample collection. Fresh feces were collected from animals by rectal sampling and stored in a 2.5% (w/v) potassium dichromate solution in clean containers. Stool samples were of normal shape. All fecal specimens were transported to the laboratory with an ice pack at 4 °C immediately after collection. DNA extraction was performed within 48 h.

**DNA extraction and PCR amplification**

DNA extraction was performed using commercial E.Z.N.A Stool DNA kits (Omega Bio-Tek Inc., Norcross, GA, USA), following the manufacturer’s recommendations. Extracted DNA samples were stored at −20 °C until PCR analysis.

The DNA samples were analyzed using nested PCR amplification of the small subunit ribosomal RNA (SSU rRNA) gene to determine the G. duodenalis infection rate [3]. Additionally, to determine the multilocus genotypes (MLGs) of the G. duodenalis isolates detected in this study, all G. duodenalis positive isolates were tested using nested PCR based on the bg [15], gdh [4] and tpi [28] loci (Table 1). Using an Applied Biosystems 2720 Thermal Cycler (Applied Biosystems, Foster City, CA, USA), PCR reactions for G. duodenalis loci were conducted in 25 µL systems: 2.5 µL 10× PCR buffer, 2 µL dNTPs (1.25 mM each), 0.3 µL each primer (25 µM each), 0.2 µL rTaq DNA polymerase (1 unit/µL each) (Takara Shuzo Co., Ltd), 2 µL of DNA sample, 17.7 µL double distilled water.

The secondary PCR products were separated by 1% agarose gel electrophoresis, following staining with DNA Green (TIANDZ, Beijing, China), observed, photographed, and recorded on a Tanon 3500 Gel Image Analysis System (TANON, Shanghai, China).

**Sequence and phylogenetic analyses**

All the secondary PCR amplicons of the SSU rRNA, bg, gdh and tpi genes from G. duodenalis-positive samples were
Table 1. Primer sequences and reaction conditions used in nested PCR amplifications.

| Gene | Primer sequences (5’ – 3’) | Nucleotide fragment (bp) | Annealing temperature (°C) | Reference |
|------|----------------------------|--------------------------|-----------------------------|-----------|
| SSU rRNA | Gia2029 (AAGTGGTTGCGACCGGACTC) | 292 | 55 | [3] |
|  | Gia2150c (CTGTGCCTACCTTGGATGT) | 50 | 59 | |
|  | RH1 (CATTGGTCTGATGTCGCC) | 50 | 59 | |
|  | RH4 (AGTCGACCTGTAGTCTCCGACCAGG) | 50 | 59 | |
| bg | BG1 (AAACCCCGACCCACCTCACCCGCAGTGC) | 511 | 65 | [15] |
|  | BG2 (GAGGCCGCCCTGATGTTTCGAGACGAC) | 50 | 59 | |
|  | BG3 (GAAACGAGCTCAGTAGGTCG) | 55 | 59 | |
|  | BG4 (CTGCAGCTCGTTCGTTGT) | 55 | 59 | |
| gdh | Gdh1 (TTGGGCTTTYAGTACAAACTC) | 520 | 50 | [4] |
|  | Gdh2 (ACCTCCTGTCGGGTGGGCGCA) | 50 | 50 | |
|  | Gdh3 (ATGGCYGAAGCTYCAAGGCAAGCT) | 50 | 50 | |
|  | Gdh4 (GGGGCGCARGGGCATGATGCGA) | 50 | 50 | |
| tpi | AL3543 (AAATATGCGCTGCTGTGCG) | 530 | 50 | [28] |
|  | AL3544 (CCCTTCATCGGCGAACCAC) | 50 | 50 | |
|  | AL3545 (GTGCCGACCACACCCCGTGCG) | 50 | 50 | |

Results

Giardia duodenalis prevalence, and distribution of assemblages

A total of 134 (64.11%, 95% CI: 57.6–70.7%) G. duodenalis-positive fecal samples were identified using the nested PCR analysis of the SSU rRNA genes in this study. The genetic diversity of the G. duodenalis-positive samples was determined by sequencing the bg, gdh and tpi genes, and a total of 39, 72 and 32 sequences, respectively, were obtained for these three genetic loci. Assemblage E (n = 130) and assemblage A (n = 4), were detected, based on the SSU rRNA gene.

Assemblage A and E

Of the bg sequences, 7 were identified as assemblage A, and 32 were identified as assemblage E. Sequence A1 (n = 4) was identical to AY655702, and A2 (n = 3) had one single-nucleotide polymorphism (SNP) relative to AY072723 (Table 2). Assemblage E sequences were designated as E1 (n = 11), E2 (n = 12), E3 (n = 7), E4 (n = 1), and E5 (n = 1). The E3 and E4 had one SNP each (A170G and A455G) relative to KT922250 and KT922248, respectively, and one sequence each was identical to MK610388, KT922250, and KP635098. At gdh sequences, 8 were identified to assemblage A, and 64 were identified to assemblage E. All 8 assemblage A sequences were identical to the genotype A1 sequence (AY178735) (Table 2). Among the assemblage E isolates, E3 and E6 had one SNP each (G369A and A455G) relative to MK645797 and MK645792, respectively. The remaining sequences were identical to counterparts in the database (E1, E2, E4 and E5 were identical to KT369778, KT369785, KY432862, and MK645788, respectively). Using KT369778 as the reference sequence, the intra-assemblage substitutions in assemblage E at the gdh gene can be seen in Table 3.

Sequence analysis of the tpi locus revealed that 12 successfully amplified isolates were identified as assemblage A, and 20 were assemblage E. A1 was identical to L02120, and A3 had

Statistical analysis

The infection rates and 95% confidence intervals (CI) were calculated by the Wald method in SPSS, version 22.0 (SPSS Inc., Chicago, IL, United States). Differences in corresponding infection rates among locations were examined by the Chi-square test, and differences were considered significant at p < 0.05.

Nucleotide sequence accession numbers

The representative nucleotide sequences generated in this study were submitted to the GenBank database under the accession numbers MK442896–MK442915.
two SNPs (A100G and C363T) relative to EU041754. Among the assemblage E sequences, E4 was identified to be a novel sequence, and the remaining sequences were consistent with KT369763, KT922262, and MF671903, respectively. The intra-assemblage substitutions in assemblage E at the tpi gene can be seen in Table 3.

### Multilocus genotyping

Using multilocus sequence typing, 4 assemblage A and 5 assemblage E isolates were successfully sequenced at all three loci (Table 4). To study the relationships between the different isolates in more detail, we performed a phylogenetic analysis based on a dataset of concatenated bg + gdh + tpi gene sequences. Data from the specimens were not included in the MLG analysis when a mixed infection was detected at one of the three loci.

Multilocus genotyping yielded two assemblages A MLGs and four assemblage E MLGs. One assemblage A MLG was identical to the AI-1, and the assemblage A MLG was considered a novel MLG (named AI-novel (IM)) which had genetic distance with AI-1 and AI-2; AI-novel (IM) and AI were in the same cluster in the phylogenetic analysis (Fig. 1). The MLG-E2 and MLG-E3 from Inner Mongolia (IM) found in this study were genetically distinct from those found in sheep from other areas in China (Fig. 2).
Discussion

*Giardia duodenalis* is an important intestinal parasite that has a global distribution in humans and a diverse range of other animals [8]. There have been reports of *G. duodenalis* infection in sheep in various regions of China, including Heilongjiang [20, 38], Henan [17], Jilin, Liaoning, Shandong [17], and Qinghai [21]. However, data on *G. duodenalis* infections in sheep in Inner Mongolia are limited, with only one published report [36]. This study reports on the occurrence and genetic characteristics of *G. duodenalis* infections in sheep in Inner Mongolia, China.

The results of this study showed that the occurrence rate of *G. duodenalis* in sheep was 64.11%, which is considerably higher than previously reported for sheep in Inner Mongolia (4.27%, 16/375) [36]. It is also higher than the infection rates of *G. duodenalis* reported for sheep from other regions of China, such as Heilongjiang (4.64%, 25/539) [38], Henan (5.24%, 100/1906) [17, 33], Jilin (0%, 0/48), Liaoning (0%,...
In previous studies, apart from one study in which two assemblage B isolates were identified in sheep from Heilongjiang province [38], *G. duodenalis* infections in Chinese sheep were all reported to be caused by either assemblage E or assemblage A [18], which is consistent with the results of this study. Assemblage E is apparently the most common *G. duodenalis* genotype in sheep [8]. In this study, assemblage E accounted for 97.01% in sheep infected with *G. duodenalis*, which is also consistent with previous reports [33, 38]. Assemblage E is commonly found in hoofed animals, including sheep, and is not considered anthropotic. However, several human cases have been reported in Egypt, Brazil and Australia [1, 7, 9, 12, 37], and additional research is therefore needed to study the public health risks of assemblage E.

Four assemblage E MLG genotypes were identified in total, all of which were new assemblage E MLG genotypes, indicating that assemblage E had high genetic diversity. The phylogenetic analysis of the concatenated sequences of assemblage E MLGs revealed that assemblage MLG-E2 (IM) and assemblage MLG-E3 (IM) found in this study were genetically distinct from the assemblages found in sheep in Qinghai and Henan Provinces, China [14, 33]. These differences were mainly due to the genetic variation of the bg locus. MLG-E1 (IM) were placed in the major cluster of MLGs from Tibetan sheep in Qinghai, whereas MLG-E4 (IM) clustered with MLGs from sheep in Henan Province (Fig. 1).

The phylogenetic analysis of the concatenated sequences of the assemblage A MLGs revealed that AI-novel (IM) was a new MLG that belonged to sub-assemblage AI. Sequences obtained from the assemblage A MLG isolates belonged to the sub-assemblage AI, which has been more commonly identified in animals than humans [8, 35]. Although *G. duodenalis* found in this study have limited zoonotic potential, a threat to public health cannot be ignored.

These MLGs results suggest that there was no significant geographic isolation of *G. duodenalis* genotypes in three regions in China. This may be because the Inner Mongolia Autonomous Region is the largest grassland pastoral area in China, and sheep fed here will be distributed to various regions of the country, thus promoting gene exchange of *G. duodenalis* in various regions. However, there may be specific genotypes in different regions for different breeding environments, and this still needs to be investigated in extensive further research. In conclusion, the results of this study showed that there was a high prevalence of *G. duodenalis* in sheep from Inner Mongolia, in northwest China. Both assemblages A and E were found, with assemblage E being the most prevalent type. Two new bg gene sequences, two new gdh gene sequences, and one new tpi gene sequence was identified. Multilocus genotyping yielded four new assemblage E MLGs and one new sub-assemblage A MLG. In addition, further studies on the zoonotic potential and geographic isolation of *G. duodenalis* from other regions are required to provide additional data.

**Conflict of interest**

The authors declare that they have no competing interests relevant to this article.

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