Host specificity of *Enterocytozoon bieneusi* genotypes in Bactrian camels (*Camelus bactrianus*) in China

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Abstract

**Background:** *Enterocytozoon bieneusi* is an obligate, intracellular fungus and is commonly reported in humans and animals. To date, there have been no reports of *E. bieneusi* infections in Bactrian camels (*Camelus bactrianus*). The present study was conducted to understand the occurrence and molecular characteristics of *E. bieneusi* in Bactrian camels in China.

**Results:** Of 407 individual Bactrian camel fecal specimens, 30.0% (122) were *E. bieneusi*-positive by nested polymerase chain reaction (PCR) based on internal transcriber spacer (ITS) sequence analysis. A total of 14 distinct *E. bieneusi* ITS genotypes were obtained: eight known genotypes (genotype EbpC, EbpA, Henan-IV, BEB6, CM8, CHG16, O and WL17), and six novel genotypes (named CAM1 to CAM6). Genotype CAM1 (59.0%, 72/122) was the most predominant genotype in Bactrian camels in Xinjiang, and genotype EbpC (18.9%, 23/122) was the second-most predominant genotype. Phylogenetic analysis revealed that six known genotypes (EbpC, EbpA, WL17, Henan-IV, CM8 and O) and three novel genotypes (CAM3, CAM5 and CAM6) fell into the human-pathogenic group 1. Two known genotypes (CHG16 and BEB6) fell into the cattle host-specific group 2. The novel genotypes CAM1, CAM2 and CAM4 cluster into group 8.

**Conclusions:** To our knowledge, this is the first report of *E. bieneusi* in Bactrian camels. The host-specific genotype CAM1 was the predominant genotype, which plays a negligible role in the zoonotic transmission of *E. bieneusi*. However, the second-most predominant genotype, EbpC, has greater zoonotic potential.

**Keywords:** *Enterocytozoon bieneusi*, Bactrian camel, Genotype, Host specificity

Background

Microsporidia are a diverse group of emerging obligate intracellular eukaryotic fungi and there are approximately 1300 microsporidian species in 160 genera [1]. To date, there are at least 14 microsporidian species reported to be infectious to humans [2]. *Enterocytozoon bieneusi* is the most frequently detected species in humans [3], as well as in domestic animals and wildlife [4], and even in environmental water samples [5].

More than 200 *E. bieneusi* genotypes have been identified in humans and animals by polymerase chain reaction (PCR) based on ribosomal internal transcribed spacer (ITS) gene sequence analysis [2, 6]. Molecular phylogenetic analysis has shown that all *E. bieneusi* ITS genotypes are clustered into nine large groups, including the potentially zoonotic group 1, and some host-specific groups (Group 2 to Group 9) [7].

The Bactrian camel (*Camelus bactrianus*) was the major means of transportation on the ancient Silk Road. Today, the population of Bactrian camels in China has been estimated at 242,000, most of which are domesticated in desert and semi-desert areas of northwestern China and play an important role in the livelihood of pastoralists through providing milk and meat [8]. There are some reports of intestinal pathogen infections in camels and Bactrian camels in the Middle East countries and China, such as *Eimeria* spp. and Cryptosporidium.
spp. [9, 10]. However, *E. bieneusi* infection has not been previously reported in Bactrian camels.

This study was undertaken to better understand the prevalence of *E. bieneusi* in Bactrian camels and assess the host specificity of *E. bieneusi* infections in Bactrian camels in China.

**Methods**

**Specimen collection**

A total of 407 individual fresh fecal specimens from Bactrian camels were collected from 18 different grazing Bactrian camel groups in 11 collection sites of Xinjiang Uygur Autonomous Region (hereinafter referred to as Xinjiang) of northwestern China. Only one specimen was collected per animal. These specimens were collected during August and September of 2013 and from July 2016 to July 2017 (Table 1). The grazing Bactrian camel groups were kept outdoors and shared pastures with cattle, sheep, goats and wild animals, and each group had approximately 30–300 animals.

After animal defecation, about 50–100 g of each fresh specimen was collected immediately from the ground using sterile gloves. Each specimen was collected in a plastic container and marked with the specimen number and site. The specimens were transported to the laboratory and stored in 2.5% (w/v) potassium dichromate solution at 4 °C before DNA extraction.

**DNA extraction and PCR amplification**

Approximately 200 mg of each fecal specimen was washed at least three times with distilled water by centrifugation at 5000× *g* for 5 min to remove the potassium dichromate. DNA was extracted using the E.Z.N.A.R® Stool DNA Kit (Omega Biotek Inc., Norcross, GA, USA) according to the manufacturer’s instructions. For *E. bieneusi* screening, nested PCR assays were used to amplify an rRNA gene fragment containing the entire internal transcriber spacer (ITS) [6]. Each specimen was analyzed in duplicate using positive and negative controls. The secondary PCR products were examined by electrophoresis in a 1.5% agarose gel and visualized after staining with GelRed™ (Biotium Inc., Hayward, CA, USA).

**Sequencing and phylogenetic analysis**

The positive secondary PCR amplicons were sent to a commercial company (GENEWIZ, Suzhou, China) for sequencing. The sequence accuracy was confirmed with bidirectional sequencing, and the sequences obtained were aligned with reference sequences downloaded from...
GenBank to determine the genotypes, using the program ClustalX 2.0 (http://www.clustal.org/).

The genotypes of *E. bieneusi* isolated in this study were compared with known *E. bieneusi* ITS genotypes with a neighbor-joining analysis in the Mega 5 program [6]. A bootstrap analysis was used to assess the robustness of the clusters using 1000 replicates. The established nomenclature system was used in naming the *E. bieneusi* ITS genotypes [11].

**Nucleotide sequence accession numbers**
The nucleotide sequences reported in this paper have been submitted to the GenBank database at the National Center for Biotechnology Information under the accession numbers: MG602791-MG602796.

**Statistical analysis**
Chi-square test was used to compare the prevalence of *E. bieneusi* infections and predominant genotypes distributions. Differences were considered significant at *P* < 0.05.

**Results and discussion**
Of all 407 individual Bactrian camel fecal specimens, 30.0% (122) were *E. bieneusi*-positive based on the ITS sequence analysis. The majority of the grazing Bactrian camel groups, 14 out of 18 (77.8%), were positive for *E. bieneusi*. Among them, Urumqi-1 had the highest infection rate (84.6%, 11/13) (*χ²* = 67.728, *df* = 17, *P* < 0.001); the other infection rates ranged from 16.7–52.9% (Table 1).

To the best of our knowledge, this is the first report of *E. bieneusi* in Bactrian camels, and the pathogen is widespread in Xinjiang, northwestern China. In China, the average prevalence of *E. bieneusi* in animals ranges from 0.9% (4/426) in rabbits [12] to 45.6% (426/934) in pigs [13]. However, *E. bieneusi* infection has only been reported in some animals in northwestern China (Table 2), the average prevalence ranging from 1.1% (4/353) in white yaks [14] to 47.8% (22/46) in sheep [2]. In Xinjiang, only dairy calves [15] and grazing horses [16] have been previously reported to have *E. bieneusi* infections, with a prevalence of 16.5% (85/514) and 30.9% (81/262), respectively. The high prevalence in Bactrian camels found in this study may be the result of free feeding and drinking water, and mixed feeding with cattle, sheep, goats and other animals in the same pastures, and with the poor veterinary service.

A total of 14 distinct *E. bieneusi* ITS genotypes were obtained from 122 positive specimens from Bactrian camels. Among them, eight were known genotypes (EbpC, EbpA, Henan-IV, BEB6, CM8, CHG16, O and WL17), and six were novel genotypes (named CAM1 to CAM6). The sequences of the novel genotypes, CAM1 to CAM6, consisted of insertions, deletions, and substitutions compared with known genotypes. In the present study, the novel genotype CAM1 was the most prevalent (59.0%, 72/122) and was significantly predominant (*χ²* = 589.836, *df* = 13, *P* < 0.001) in Bactrian camels in Xinjiang. The other novel genotypes CAM2 (*n* = 8), CAM4 (*n* = 5), CAM3 (*n* = 1), CAM5 (*n* = 1) and CAM6 (*n* = 1) were also identified. The genotype EbpC was the predominant identified of the known genotypes (18.9%, 23/122) in Bactrian camels in Xinjiang, followed by genotype EbpA (*n* = 5). However, the other known genotypes Henan-IV, BEB6, CM8, CHG16, O and WL17, were identified in only one specimen each, though they are not listed in Table 2.

**Table 2** *Enterocytozoon bieneusi* infections and genotype distributions in animals in northwestern China: summary of previous literature

| Region    | Host                  | No. of specimens | No. infected (%) | Genotype (no.)                                                                 | Reference |
|-----------|-----------------------|------------------|-----------------|--------------------------------------------------------------------------------|-----------|
| Xinjiang  | Dairy cattle          | 514              | 85 (16.5)       | J (57), I (19), BEB4 (4), D (2), EbpC (2), CC4 (1)                          | [15]      |
| Xinjiang  | Grazing horses        | 262              | 81 (30.9)       | EbpC (21), EbpA (20), BEB6 (9), CHG19 (2), CM6 (4), CM7 (2), CM8 (1),       | [16]      |
|           |                       |                  |                 | CS-1 (1), CS-4 (1), D (1), G (3), horse1 (4), horse2 (2), O (4), Peru8 (1) |           |
|           |                       |                  |                 | XJH1 (2), XJH2 (1), XJH3 (1), XJH4 (1)                                    |           |
| Gansu     | White yaks            | 353              | 4 (1.1)         | BEB4 (2), I (1), WCY1 (1)                                                  | [14]      |
| Ningxia   | Dairy cattle          | 109              | 51 (46.8)       | J (25), CM8 (14), I (8), O (1), CHC1 (1), CHC2 (1), CHC3 (1)               | [6]       |
| Qinghai   | Yaks                  | 327              | 23 (7.0)        | BEB4 (16), I (1), J (1), CHN11 (4), CHN12 (1)                              | [18]      |
| Shaanxi   | Sheep                 | 46               | 22 (47.8)       | BEB6 (4), CHG1 (3), CHG2 (3), CD6 (3), CHG5 (2), E (1), F (1),            | [2]       |
|           |                       |                  |                 | CHG14 (1), CHG16 (1), CHG24 (1)                                            |           |
| Shaanxi   | Monkeys               | 197              | 25 (12.7)       | D (10), BEB6 (4), MH (7), XH (2), BS (2)                                   | [19]      |
| Shaanxi   | Dairy cattle          | 198              | 39 (19.7)       | I (21), J (16), CHN1 (1), CSX1 (1)                                        | [20]      |
| Shaanxi   | Beef cattle           | 173              | 34 (19.7)       | I (19), J (14), CSX2 (1)                                                   | [20]      |
| Shaanxi   | Golden takins         | 191              | 28 (14.7)       | BEB6 (10), D (8), I (6), TEB1–4 (each 1)                                   | [21]      |
| Shaanxi   | Cashmere goats        | 315              | 50 (15.9)       | SX1 (43), CHG1 (7)                                                         | [22]      |
| Shaanxi   | Dairy goats           | 170              | 56 (32.9)       | CHG1 (35), SX1 (13), BEB6 (6), CHG2 (2)                                    | [22]      |
have been commonly reported in many other types of animals.

The phylogenetic analysis of the ITS genotypes revealed the following clusters: Group 1, Group 2 and Group 8. The six known genotypes (EbpC, EbpA, WL17, Henan-IV, CM8 and O) and three novel genotypes (CAM3, CAM5 and CAM6) identified in this study fell into the human-pathogenic Group 1 (Fig. 1), which is the genotype of major zoonotic potential suggesting that the Bactrian camels play a potential role in \textit{E. bieneusi} transmission to humans [11]. In contrast, the two known genotypes CHG16 and BEB6 fell into the cattle host-specific Group 2. The three novel genotypes CAM1, CAM2 and CAM4 clustered into Group 8 (69.7%, 85/
122) (Fig. 1), suggesting that the host-specific genotype CAM1 in Bactrian camels exhibits less zoonotic potential compared to the genotypes clustered into the human-pathogenic group.

In previous studies, *E. bieneusi* genotype EbpC and EbpA were reported in humans and various animals and were also the predominant genotypes in the reports of humans and pigs in China [13, 17]. Genotypes EbpC and EbpA were the most common *E. bieneusi* genotypes in grazing horses in Xinjiang [16], and genotype EbpC was also identified in dairy calves in Xinjiang [15]. Similarly, the *E. bieneusi* zoonotic genotypes EbpC and EbpA were identified in Bactrian camels in the present study. However, there were no published reports of genotypes EbpC and EbpA in animals in Gansu, Ningxia, Qinghai, and Shaanxi, northwestern China (Table 2). Further investigations of the epidemiology and host specificity of *E. bieneusi* in humans and other animals in Xinjiang might be informative.

**Conclusions**

The present study demonstrated a widespread occurrence of *E. bieneusi* in Bactrian camels in Xinjiang, China. The host-specific genotype, CAM1, was the most predominant genotype, which plays a negligible role in the zoonotic transmission of *E. bieneusi*. The second-most predominant genotype, EbpC, in addition to other genotypes of zoonotic potential, was also commonly identified in Bactrian camels in this study. Bactrian camels could serve as a vector for *E. bieneusi* transmission to humans and other animals, and *vice versa*.

**Abbreviations**

ITS: Ribosomal internal transcribed spacer; PCR: Polymerase chain reaction

**Acknowledgments**

We thank Liwen Bianji, Edanz Editing China (http://www.liwenbianji.cn/ac), for editing the English text of a draft of this manuscript.

**Funding**

This study was partly supported by the National Key Research and Development Program of China (2017YFD0501305), and the Key Laboratory of Tarim Animal Husbandry, Science, and Technology, Xinjiang Production & Construction Group Open Project Fund (HS2017142), and the President Foundation of Tarim University (TDZKCX201701).

**Availability of data and materials**

The data sets supporting the conclusions of this article are included within the article. The newly generated sequences were deposited in the GenBank database under the accession numbers MG602791-MG602796.

**Authors’ contributions**

LXZ and MQ designed the study. BJ, AYZ and ZLW participated in specimen collection and analysis. JQL, BJ, MQ and ZHC contributed to data analysis. LXZ, JQL and MQ wrote the manuscript. All authors read and approved the final manuscript.

**Ethics approval and consent to participate**

This study was conducted in accordance with the Chinese Laboratory Animal Administration Act (1988). The study protocol was reviewed and approved by the Research Ethics Committee of Tarim University. Appropriate permission was obtained from the owner before the collection of fecal specimens.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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**Received: 2 January 2018 Accepted: 12 March 2018**

**Published online: 02 April 2018**

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