INTRODUCTION

Sheep keds, a kind of ectoparasites from Hippoboscidae (Diptera: Hippoboscoidea) that subsist solely on blood [1,2], parasitize on sheep. However, they have also been reported to parasitize on other domestic and wild animals, including goats [3], rabbits [1], dogs [4], Tibetan antelope [2,5], European bison [6], red foxes [7] as well as humans [4].

Sheep keds have been widely distributed geographically. It was originally found in most parts of Europe, Northwest Africa, Mongolia and northern India, and later found in Kenya, South Africa, Japan, Australia, New Zealand and most parts of North America [2]. However, sheep keds parasitizing on sheep and Tibetan antelopes were only reported in few areas in China such as Xinjiang [2,8], Qinghai [2,5], Liaoning [9], and Gansu [10,11]. In addition, adult and pupae of sheep keds were found on the imported sheep, sheep skin and wool in quarantine ports of Shandong [12], Beijing [13], and Zhejiang [14].

Sheep keds could cause damage to skin, inflammation, anemia, slow weight gain, poor wool quality, and low yield for sheep [1,2,10,15]. Moreover, it is a media for insect-borne pathogens such as Trypanosoma melophagium [16], Anaplasma ovis [17], blue-tongue virus [18], Bartonella schoenbuchensis, Bartonella chomelii [19], B. melophagi [15,20], and Bartonella spp. In China, Borrelia garinii and Borrelia valaisiana-like groups [8] were detected in sheep keds on Tibetan; Rickettsia raoultii and Rickettsia slovaca were detected in sheep keds on Xinjiang [2]; Bartonella, Arsenophonus, Wolbachia [10,11], Enterobacter, Shewanella, Acinetobacter, Bacillus, Halomonas, and Staphylococcus [10] were detected in sheep keds on Gansu. Sheep keds have caused significant economic losses to the world’s animal husbandry.

Bartonella initially belongs to Rickettsiales, Bartonellaceae. According to later genetic analysis, it then belongs to Rhizobiales, Bartonellaceae, Bartonella. It is a widely distributed facultative intracellular parasite and transmitted by hematophagous arthropods among vertebrates [16]. At least 30 Bartonella that
cause disease in human body have been reported so far [15]. Nowadays, B. bovis, B. melophagi, B. capreoli, B. chomelii, B. dromedarii, and B. schoenbuchensis have been identified in cloven-hoofed animals [20]. Wolbachia spp. belongs to Rickettsiales, Anaplasmataceae, Wolbachia. It is the most widely distributed intracellular symbiotic bacteria in the world. About 40% of arthropods and over 65% of insect species naturally carry Wolbachia [21,22]. At present, phylogenetic analysis based on multiple genes was used to divide Wolbachia into 16 supergroups [23]. It has been given wide attention because it can participate in the reproductive regulation of arthropod hosts in various ways.

Bartonella and Wolbachia are distributed globally. They are rarely studied in China and widely distributed in Xinjiang. It is necessary to detect the pathogens of sheep keds such as Bartonella and Wolbachia. It will make an important contribution to the health of humans and animals.

**MATERIALS AND METHODS**

Sheep keds samples
In July 2016, more than 200 sheep keds specimens were collected from sheep of local fair in Yaha Town of Kuqa in Aksu, Xinjiang (1,029 m above sea level; 41°44’N, 83°14’E). Seven experimental specimens were selected from them and preserved in 70% ethanol.

In June 2017, more than 200 sheep keds specimens were collected from each of the 5 sheep from the local fair in Yaha Town of Kuqa in Aksu, Xinjiang. These sheep keds specimens were placed in sampling vials with sufficient air and transported immediately to the laboratory for cryopreservation. Two sheep keds specimens were randomly selected from each sheep as experimental specimens.

In this study, 17 samples were processed individually.

DNA extraction, PCR and sequence analysis
Sheep keds preserved in 70% ethanol were washed by using 50% ethanol, 30% ethanol, 10% ethanol. And they were followed by washing with distilled water. The cryopreserved sheep keds were washed twice with distilled water. After the elution, they were dried by sterile filter paper and cut into pieces, and then placed into 2 ml sterile tubes. The kit was operated according to instructions of TaKaRa MiniBEST Universal Genomic DNA Extraction Kit Ver. 5.0 (TaKaRa, Dalian, China). And then the sheep keds genomic DNA was extracted and eluted with 50 μl Elution Buffer twice. Finally, the genomic DNA was collected and then stored at -20°C.

The Bartonella gltA gene and the Anaplasma 16S rRNA gene were amplified according to the instructions of the kit of Premix Taq™ (TaKaRa Taq™ Version 2.0) (TaKaRa). Bartonella gltA gene amplification primers were 5'-GGT ACC YAC AGA AGA AGT CC-3' and 5'-ATG GCA AAA AGA AGA GGA TTA AAC A-3' [24,25]; the expected amplification product size was 379 bp. The genus-specific set of primers for 16S rRNA gene were used for Wolbachia DNA amplification, and the primers sequence were 5'-GCT ACC YAC AGA AGA AGT CC-3' and 5'-TAG CAC TCA TCG TTT ACA GC-3' [26]; the expected amplification product size was 345 bp. The PCR system was 50 μl and the annealing temperature was 55°C and 54°C, respectively.

After the amplification products were identified by 1% gel electrophoresis, the positive amplification products were sequenced using the ABI PRISM™ 3730 XL DNA Analyzer. Sequencing results were performed using the BLAST online platform (https://blast.ncbi.nlm.nih.gov/Blast.cgi?PROGRAM=blastn&PAGE_TYPE=BlastSearch&LINK_LOC=blasthome), Primer Premier 5.0, DNAMAN, DNAStar, and Molecular Biology Software (MEGA 5.0) to conduct the sequence analysis and phylogenetic tree construction. The evolutionary history was inferred by using the Maximum Likelihood method based on the Tamura-Nei model.

**RESULTS**

Detection of B. melophagi
PCR was conducted on 17 sheep keds samples from Kuqa County, Aksu Prefecture, Xinjiang using the Bartonella gltA gene amplification primers, and all samples were positive (100%, 17/17). The size of the amplified products was identical as expected. The amplified products of gltA gene from randomly selected 5 sheep keds were sequenced. The sequences results were analyzed and compared by the BLAST. The results were over 99% in similarity when compared with the Bartonella gltA gene sequence in the database. Moreover, the top 5 gene sequence hosts in similarity were all ectoparasites of Hippoboscidae (Diptera: Hippoboscoidea) (Table 1). A total of 42 nucleotide sequences based on Bartonella gltA gene (312 bp) and Rickettsia gltA gene (330 bp) of outgroup sequence showed that 5 segments of the Bartonella gltA gene sequences collected in this study was clustered together with other B. melophagi gltA gene sequences from other sheep keds hosts (Fig. 1).
### Table 1. BLAST comparison analysis results of Bartonella gltA gene and Wolbachia 16S rRNA gene

| Pathogen | No. This study | First  | Second | Third | Fourth | Fifth |
|----------|----------------|--------|--------|-------|--------|-------|
| Bartonella | XJNJ-1 (MG701233) | AY724769, 99% (349/349), M. ovinus | KY224150, 99% (345/349), M. ovinus | AY724768, 99% (345/349), M. ovinus | KY224151, 99% (345/349), M. ovinus | AJ564633, 98% (342/349), deer ked |
|          | XJNJ-2 (MG701234) | KY224150, 100% (349/349), M. ovinus | KY224150, 99% (345/349), M. ovinus | KY224151, 99% (345/349), M. ovinus | KY224151, 99% (345/349), M. ovinus | AJ564633, 99% (346/351), deer ked |
|          | XJNJ-3 (MG701235) | KY224150, 99% (345/349), M. ovinus | KY224150, 99% (345/349), M. ovinus | KY224151, 99% (345/349), M. ovinus | KY224151, 99% (345/349), M. ovinus | AJ564633, 97% (340/349), deer ked |
|          | XJNJ-4 (MG701236) | KY224150, 99% (347/349), M. ovinus | KY224150, 99% (347/349), M. ovinus | KY224151, 99% (347/349), M. ovinus | KY224151, 99% (345/349), M. ovinus | AJ564633, 98% (343/350), deer ked |
|          | XJNJ-5 (MG701237) | KY224150, 99% (340/341), M. ovinus | KY224150, 99% (340/341), M. ovinus | KY224151, 99% (338/341), M. ovinus | KY224151, 99% (335/341), M. ovinus | AJ564633, 98% (333/341), M. ovinus |
| Wolbachia | XJNJ-1 (MG701229) | EU780456, 96% (302/316), Cimex lectularius | KY224164, 95% (300/316), M. ovinus | KY224164, 95% (300/316), M. ovinus | KY224163, 95% (300/315), M. ovinus | KT796584, 95% (300/315), Neelus incertus |
|          | XJNJ-2 (MG701230) | KY224164, 99% (299/303), M. ovinus | KY224163, 99% (299/303), M. ovinus | KY224163, 99% (299/303), M. ovinus | KY224163, 99% (299/303), M. ovinus | KT796584, 99% (299/303), Cercopithifilaria japonica |
|          | XJNJ-3 (MG701221) | EU780456, 98% (312/317), C. lectularius | KY224164, 99% (303/305), M. ovinus | KY224163, 99% (303/305), M. ovinus | KY224163, 99% (302/306), M. ovinus | KT796584, 99% (302/306), Bartonella australis |
|          | XJNJ-4 (MG701232) | EU780456, 99% (303/306), C. lectularius | KY224164, 99% (302/306), M. ovinus | KY224163, 99% (302/306), M. ovinus | KY224163, 99% (302/306), M. ovinus | KT796584, 99% (302/306), Bartonella australis |

**Fig. 1.** Phylogenetic tree of Bartonella based on gltA gene. The evolutionary history was inferred by using the Maximum Likelihood method based on the Tamura-Nei model. The analysis involved 42 nucleotide sequences. There were a total of 309 positions in the final dataset. Evolutionary analyses were conducted in MEGA5. Sequences of this work were marked with black triangle (▲).
Sequence analysis of *Wolbachia* of sheep keds

PCR was conducted on 17 sheep keds samples from Kuqa County, Aksu Prefecture, Xinjiang using the *Anaplasma* 16S rRNA gene amplification primers. The results showed that 15 samples were positive with the positive rate at 88.23% (15/17) and the size of the amplification products was consistent as the expected size. The 16S rRNA gene amplification products from 5 randomly selected sheep keds were sequenced and then 4 sequence results obtained were compared and analysed by the BLAST. Three segments of them have the highest similarity at 99% after comparing with *Wolbachia* 16S rRNA gene sequences in the database. Another segment has the similarity at 96%. The top 5 gene sequence in similarity all contain 2 segments of the gene sequences from the host sequence of sheep keds (Table 1). A total of 40 nucleotide sequences of supergroups A-N based on *Wolbachia* 16S rRNA gene (247 bp) and *Rickettsia* 16S rRNA (250 bp) of outgroup sequence showed that 4 segments of *Wolbachia* 16S rRNA gene sequence collected in this study was clustered together with the *Wolbachia* sequence of 2 sheep keds hosts and *Wolbachia* sequence of other supergroup F (Fig. 2). The 3 segments designated as supergroup F with 1 segment need further analysis with other genes. In this study, 4 *Wolbachia* 16S rRNA gene fragments from sheep keds collected from Aksu Prefecture of Xinjiang were named as *Wolbachia* XJNJ-1, XJNJ-2, XJNJ-3, and XJNJ-4 strains; accession numbers were MG701229, MG701230, MG701231, and MG701232, respectively, when submitting to GenBank.
DISCUSSION

From routine PCR, sequencing, and sequence analysis on sheep keds in Aksu Prefecture of southern Xinjiang, China, the results showed that the Bartonella was carried by all the 17 sheep keds and the Wolbachia was carried by 15 out of them. According to the literature on the differential analysis of Bartonella gltA gene [25,27-29] and Wolbachia 16S rRNA gene in different host [30-33], it was confirmed that Bartonella detected in this study was B. melophagi, 3 of the Wolbachia were supergroup F. Previous reports showed that Bartonella and Wolbachia were detected in sheep keds in Gansu, China [10,11], but Bartonella and Wolbachia were not classified. The sequences of B. melophagi gltA gene (KY224150 and KY224151) and Wolbachia 16S rRNA gene (KY224163 and KY224164) from Xinjiang sheep keds were found in the GenBank database. However, it is short of details in the literature. This is the first study reported that Wolbachia supergroup F found in sheep keds and provided the molecular evidence that sheep keds carried B. melophagi and Wolbachia supergroup F in the Aksu Prefecture of southern Xinjiang, China. The 2 pathogens were found in sheep keds around Taldimakan Desert for the first time.

It is reported previously that Bartonella was detected in sheep and sheep keds [10,11,15,19,20,34]. This study also confirmed the widespread prevalence of B. melophagi in sheep keds in Kuqa County, Aksu Prefecture, China. Currently, no report has revealed the relation between B. melophagi and sheep, but it has been reported that B. bovis was associated with bovine endocarditis [35] and previous reports mentioned that B. melophagi was isolated from 2 female patients with pericarditis and skin lesions with animal exposure history [36]. This study is in agreement with the literature which showed it may be a symbiotic relationship between B. melophagi and sheep keds [19] or B. melophagi is well adapted to sheep and sheep keds, and frequently transferred between 2 hosts [20]. Sheep keds may play an important role in the spread of B. melophagi among sheep. The exact role and mode of action need further study. Close attention should be given that B. melophagi existed in the local population of Xinjiang.

This is the second report in the world that Wolbachia is detected in sheep keds. It also provides another evidence that Wolbachia is the most widely distributed intracellular symbiotic bacteria. Three Wolbachia strains in this study are attributed to supergroup F. It is the first report of the supergroup F Wolbachia on sheep keds. In this study, we need to further analyze and identify the subgroups of 1 strain in combination with other Wolbachia gene fragments. In addition, genetic analysis is based on the Wolbachia 16S rRNA gene. The classification of supergroups B, N, K, and I is relatively confusing, thus, more accurate and detailed classification requires multi-gene tandem or whole genome integrated data analysis. Wolbachia plays a regulatory role in host reproductive behavior by means such as inducing host cytoplasm incompatibility and parthenogenesis [11]. Further studies are needed to determine its role in regulating the reproductive behavior and regulation mode.

Bartonella and Wolbachia are zoonosis pathogens. They use vampire arthropods as their parasitic host, storing hosts or the media. They are highly valued by veterinary medicine and by the medicine in the control of infectious diseases or vector biological control.

ACKNOWLEDGMENTS

This study was funded by the National Natural Science Foundation of China (No. 31460655) and the open project of Key Laboratory of Tarim Animal Husbandry Science and Technology, Xinjiang Production & Construction Corps (HS201501, HS201801).

CONFLICT OF INTEREST

The authors declare no conflicts of interest.

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