First molecular survey and identification of *Anaplasma* spp. in white yaks (*Bos grunniens*) in China

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SUMMARY

Anaplasmosis is caused by a group of obligate intracellular bacteria in the genus *Anaplasma*, which are transmitted by ticks and infect humans, domestic animals and wildlife. This study was conducted to determine the prevalence and molecular characterization of *Anaplasma* spp. in semi-wild white yaks sampled in Tianzhu Tibetan Autonomous County, northwest China. Out of 332 samples tested, 35 (10·9%) were positive for *Anaplasma* spp. The positive rates were 6·2% (20/322) and 5·3% (17/322) for *Anaplasma bovis* and *Anaplasma phagocytophilum* in white yaks, respectively. None of the sample was positive for *Anaplasma marginale*. Two (0·6%) samples were simultaneously positive to *A. bovis* and *A. phagocytophilum*. Sequence analysis of the 16S rRNA gene revealed two genotypes (ApG1 and ApG2) of *A. phagocytophilum* and two sequence types (ST1 and ST2) of *A. bovis* in white yaks. This study is the first to document the presence of *Anaplasma* in white yaks. Our findings extend the host range for *Anaplasma* species and provide more valuable information for the control and management of anaplasmosis in white yaks.

Key words: 16S rRNA gene, Anaplasma phagocytophilum, Anaplasma bovis, white yak, tick-borne disease.

INTRODUCTION

Anaplasmosis is a tick-borne infectious disease of a variety of wild and domestic animals and human beings throughout the world. It is caused by a group of obligate intracellular bacteria in the genus Anaplasma (Rickettsiales: Anaplasmataceae), including Anaplasma phagocytophilum, Anaplasma marginale, Anaplasma ovis, Anaplasma bovis and Anaplasma platys (Dumler et al. 2001). Among those, three Anaplasma species (A. phagocytophilum, A. marginale and A. bovis) have been detected in cattle and recognized as the agents of bovine anaplasmosis in China (Bai et al. 1987; Zhang et al. 2013; Yang et al. 2015). Recently, an A. platys-like pathogen was also identified in cattle from Xinjiang, Northwest China (Yang et al. 2015).

The white yak (*Bos grunniens*) is a semi-wild and endemic species in Tianzhu Tibetan Autonomous County (TTAC) that relies heavily on white yaks farming for milk, meat and local economy. These animals inhabit in the alpine steppe ecoregion at altitude over 3000 m, with a population of ~49 400. Previous

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studies have reported that the Tianzhu white yaks are infected with *Chlamydia abortus* (Qin et al. 2015a). Moreover, a high seroprevalence (17·76%) of tickborne *Babesia bigemina* was also observed in them (Qin et al. 2015b). Apart from the above reports, information on the *Anaplasma* infection is currently not available. The objective of this study was to determine whether and what species of *Anaplasma* agents infect white yaks in TTAC in Gansu Province, Northwest China. The molecular characterization of the identified *Anaplasma* strains was further analysed.

MATERIALS AND METHODS

Study sites

This study was carried out in TTAC in Gansu Province, Northwest China, with a total area of 7149 km². Sampling sites were located between longitude 102°07′–103°46′ east and latitude 36°31′–37°55′ north in Gansu Province, Northwest China. The annual average temperature here is –8 to 4°C, having an obvious vertical distribution of temperature.

Blood sampling and DNA preparation

The surveillance was performed from March to July in 2015 during the peak season of tick in rural areas

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Table 1. Primers and PCR amplification conditions

Pathogen and target gene	Primer name	Primer sequence (5′–3′)	Annealing temp (°C)	Amplicon size (bp)
A. phagocytophi- lum, 16S rRNA	EC9	TACCTTGTTACGACTT	55	1462
	EC12a	TGATCCTGGCTCAGAACGAACG		
	SSAP2f	GCTGAATGTGGGGATAATTTAT	55	641
	SSAP2r	ATGGCTGCTTCCTTTCGGTTA		
A. bovis, 16S rRNA	EC9	TACCTTGTTACGACTT	55	1462
	EC12a	TGATCCTGGCTCAGAACGAACG		
	AB1f	CTCGTAGCTTGCTATGAGAAC	55	551
	AB1r	TCTCCCGGACTCCAGTCTG		
A. marginale, msp4	MSP45	GGGAGCTCCTATGAATTACAGAGAATTGTTTAC	55	869
	MSP43	CCGGATCCTTAGCTGAACAGGAATCTTGC		
A. marginale, msp5	AM1 AM2	TGTCTAACCTTCTGCTGTTCGTTGC ACACGAAACTGTACCACTGCCATGC	60	643

in TTAC. EDTA whole-blood samples were taken from the jugular vein of 332 white yaks and collected in a sterile tube. Total DNA was extracted from 300 μ L of blood using the Gentra Puregene DNA purification kit (Qiagen, Beijing, China) following the instructions of the manufacturer.

PCR and sequence analysis

The total DNA was detected by nested PCR for the presence of A. phagocytophilum and A. bovis 16S rRNA gene and A. marginale major surface protein 4 (msp4) and major surface protein 5 (msp5) genes as previously described (de la Fuente et al. 2001; Kawahara et al. 2006; Zhou et al. 2007; Zhang et al. 2013). The PCR primers for 16S rRNA, msp4 and msp5 amplification were included in Table 1. The reaction was performed in an automatic thermocycler (Bio-Rad, Hercules, CA, USA) in a total volume of 25 μ L containing 2·5 μ L of 10 × PCR buffer (Mg²⁺ Plus), 2·0 μL of each dNTP at 2·5 mM, 1.25 U of Taq DNA polymerase (TaKaRa, Dalian, China), $2.0 \,\mu\text{L}$ of template DNA, $1.0 \,\mu\text{L}$ of each primer (20 pm) and 16.25 µL of distilled water. DNA extracted from whole blood of sheep infected with A. phagocytophilum and cattle infected with A. marginale and A. bovis was used as the positive control, and sterile water was used as the negative control for each run. Cycling conditions for PCR amplification were: 4 min of denaturation at 94 °C, 35 cycles at 94 °C for 30 s, annealing for 30 s (annealing temperatures of primers was listed in Table 1), and 72 °C for 1-1.5 min (dependent on the target gene), with a final extension step at 72 °C for 10 min. PCR products were determined by UV transillumination in a 1.0% agarose gel following electrophoresis and staining with ethidium bromide.

Positive PCR products were purified (TaKaRa Agarose Gel DNA purification Kit Ver.2·0, TaKaRa, China), cloned into pGEM-T Easy vector (Promega, USA) and transformed into

Escherichia coli JM109 competent cells (TaKaRa, China). Two recombinant clones were selected for sequencing using BigDye Terminator Mix (Sangon, China). The obtained sequences were analysed by a BLASTn search in GenBank or by using the Clustal W method in the MegAlign software (DNAStar, Madison, WI). Phylogenetic trees were constructed based on the sequence distance method using the neighbour-joining (NJ) algorithm with the Kimura two-parameter model of the Mega 4·0 Software (Tamura et al. 2007).

Nucleotide sequence accession numbers

The GenBank accession numbers obtained in this study were as follows: KT824824–KT824833 (Ap20-a, Ap20-b, Ap31-a, Ap31-b, Ap54-a, Ap54-b, Ap5-a, Ap5-b, Ap12-a and Ap12-b) for *A. phagocytophilum* and KT824834–KT824851 (Ab20, Ab21-b, Ab21-c, Ab23-a, Ab23-b, Ab26-a, Ab26-b, Ab30-a, Ab30-b, Ab6-a, Ab6-b, Ab9-a, Ab9-b, Ab11-a, Ab11-b, Ab19-a, Ab19-b, Ab20-a) for *A. bovis*.

RESULTS

Out of 332 samples tested, 35 (10.9%) were positive for Anaplasma spp. The positive rates were 6.2% (20/322) and 5.3% (17/322) for $A.\ bovis$ and $A.\ phagocytophilum$ in white yaks, respectively. None of the sample was positive for $A.\ marginale$. Two (0.6%) samples were simultaneously positive to $A.\ bovis$ and $A.\ phagocytophilum$.

To characterize the *Anaplasma* spp. detected in yaks, positive samples were sequenced. Twenty-eight sequences were obtained in this study: 10 for *A. phagocytophilum* and 18 for *A. bovis*. The partial 16S rRNA gene sequences of *A. phagocytophilum* (599 bp) and *A. bovis* (511 bp) were analysed. Alignment of these sequences revealed two genotypes of *A. phagocytophilum* in white yaks (ApG1

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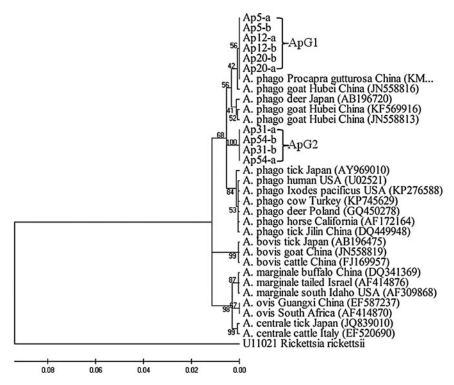


Fig. 1. Phylogenetic analysis of *A. phagocytophilum* (*A. phago*) based on 16S rRNA gene partial sequences. An alignment of 16S rRNA sequences from position 694 to 1334 of the sequence (based on strain ES34, GenBank accession no. AB196720) was used to construct this tree. *Rickettsia rickettsii* is used as an outgroup.

and ApG2). The similarity between ApG1 (Ap54-a, Ap54-b, Ap31-a and Ap 31-b) and ApG2 (Ap5-1, Ap5-b, Ap12-a, Ap12-b, Ap20-a and Ap20-b) was 98·5%. ApG1 were 98·8% identical to strains MR-23 (GenBank accession no. KP276588) isolated from *Ixodes pacificus* in the USA and Trbrt45 (GenBank accession no. KP745629) isolated from cow in Turkey. ApG2 were 100% identical to strain JC3-3 (GenBank accession no. KM186948) that was detected in the Mongolian gazelle from China. The 16S rRNA of *A. bovis* sequences identified in yaks were 99·6–100% identical to each other and to strain Ab4a (GenBank accession no. KJ639885) identified in red deer from Qilian-Mountain in Northwest China.

Phylogenetic analysis of 16S rRNA gene was conducted with *A. phagocytophilum* and *A. bovis* sequences in this study and selected sequences of *Anaplasma* spp. deposited in GenBank (Figs 1 and 2). The results revealed that ApG1 clustered independently from all known *A. phagocytophilum* sequences, ApG2 displayed a close relationship with the sequence amplified from Mongolian gazelle found in China (GenBank accession no. KM186948) (Fig. 1). All *A. bovis* strains from white yaks were classified into *A. bovis* cluster and contained two sequence types (ST1 and ST2) (Fig. 2).

DISCUSSION

In the present study, molecular survey and characterization of *Anaplasma* pathogens was performed

in white yaks. Our findings clearly demonstrated that the presence of *A. phagocytophilum* and *A. bovis* in white yaks from TTAC.

Anaplasma phagocytophilum has been recognized as an emerging pathogen of veterinary and human health significance (Chen et al. 1994). In addition to humans, A. phagocytophilum infection has been reported in some domestic animals, such as sheep, goats and cattle (Aktas et al. 2011, 2012; Altay et al. 2014). Anaplasma phagocytophilum is a wide spread tick-borne infection causing human granulocytic anaplasmosis and tick-borne fever in domestic ruminants, responsible for serious economic loss to sheep and cattle industry (Stuen, 2007). Several molecular surveys of A. phagocytophilum have been performed in ticks and domestic ruminants in China. In this study, A. phagocytophilum infection was reported in white yaks. The positive rate of A. phagocytophilum in white yaks (17/322, 5·3%) was much lower than in black yaks (51/158, 32.3%) and cattleyaks (7/20, 35.0%) conducted in Gannan Tibetan Autonomous Prefecture in Gansu Province, and was slightly lower than that in cattle (8/125, 6·4%) in Xinjiang Province (Yang et al. 2013, 2015). In general, A. phagocytophilum is associated with Ixodes ticks, including I. pacificus, Ixodes dentatus and Ixodes scapularis in the USA (Goethert and Telford, 2003; Teglas and Foley, 2006); Ixodes ricinus and Ixodes trianguliceps in Europe (Bown et al. 2008; Aktas et al. 2010); Ixodes persulcatus in Asia (Cao et al. 2003). However, the information

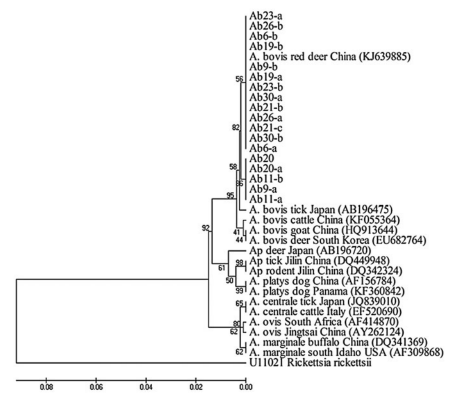


Fig. 2. Phylogenetic analysis of *A. bovis* based on 16S rRNA gene partial sequences. An alignment of 16S rRNA sequences from position 60 to 610 of the sequence (based on strain ES1019, GenBank accession no. HQ913644) was used to construct this tree. *Rickettsia rickettsii* is used as an outgroup.

on the epidemiology of the potential tick vectors is unclear in the study sites, therefore the investigation of tick species transmitting A. phagocytophilum in TTAC should be further carried out. To date, no transovarial transmission of A. phagocytophilum was described, except for Dermacentor albipictus ticks (abnormal feeding systems) (Baldridge et al. 2009). Identification of reservoir hosts is important to prevent and control of A. phagocytophilum, that play a critical role in the maintenance of the agent in nature. Our results indicated that white yaks are part of the natural maintenance cycle of A. phagocytophilum. Anaplasma phagocytophilum can cause persistent infection in ruminants and other animals for several years (Brown and Barbet, 2016). The white yak serve as reservoir host may facilitate further spread of infection. In this study, sequence analysis showed two 16S rRNA genotypes of A. phagocytophilum (ApG1 and ApG2) existed in white yaks (Fig. 1). Considerable strain variation of A. phagocytophilum has been reported in different hosts or geographic locations, and four geographically dispersed ecotypes were identified based on groESL gene and showed significantly different host ranges in Europe (Jahfari et al. 2014). It has been showed that the A. phagocytophilum ecotypes I in cattle were in the same group as those that infected human beings (Jahfari et al. 2014). Although no human case was currently reported in TTAC, A. phagocytophilum infection in white yaks warrants further investigation.

Historically, A. bovis was usually reported in buffalo and cattle from South America and Africa (Ooshiro et al. 2008). Recently, it has also been reported in cattle in continental Europe (Ceci et al. 2014; Aktas and Ozubek, 2015). Aside from the aforementioned study, the agent was identified in other ruminant and non-ruminant animals such as deer, cats, dogs and rabbits sheep, goats, (Kawahara et al. 2006; Ooshiro et al. 2008; Sakamoto et al. 2010; Liu et al. 2012; Tateno et al. 2013; Ben Said et al. 2015). In China, molecular evidence for the presence of A. bovis was reported in several wild and domestic animals including Reeves' muntjac, Mongolian gazelle, red deer, sika deer, cattle and goats (Liu et al. 2012; Yang et al. 2014, 2015; Li et al. 2015). In the present study, A. bovis infection was found in white yaks. The infection rate (6.2%, 20/322) was significantly lower than the 49.6% prevalence in goats in central and Southern China (Liu et al. 2012), and 42.7% in sheep and 23.8% in goats in Tunisia (Ben Said et al. 2015). However, it is almost comparable with 4.8% prevalence in cattle in Xinjiang, Northwest China (Yang et al. 2015). Sequences analysis revealed that A. bovis identified in this study were divided into two sequence types, indicating two different strains infected white yaks in TTAC. The ST1 has been identified in red deer in the Qilian-Mountain area that is near the study site, indicating A. bovis ST1 circulates in this region and has deer

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and white yaks as hosts. Previous report suggested that infection with *A. phagocytophilum* strain precludes infection with other strains; this could help to maintain the host tropism among strains (Rejmanek *et al.* 2012). *Anaplasma bovis* has a wide range of hosts, and the host tropism of *A. bovis* strains has not been demonstrated. However, both ST1 and ST2 of *A. bovis* were found in a white yak in this study (Ab9-a and Ab9-b) (Fig. 2).

Furthermore, A. marginale was known to cause severe disease in Northern China (Bai et al. 1987). However, none of the white yaks in TTAC were positive for A. marginale using two PCR assays based on msp4 and msp5 genes. More samples should be investigated to determine the ability of white yak as the host for A. marginale in the future study. In addition, coinfection of A. phagocytophilum and A. bovis occurred in two (0.6%) of the sampled animals. Although Anaplasma species infect different cell types (Dumler et al. 2001; Rar and Golovljova, 2011), coinfection increases the difficulties in diagnosis and treatment of bovine anaplasmosis.

In summary, this study is the first report to document the presence of *Anaplasma* in white yaks. Our findings extend the host range for *A. phagocytophilum* and *A. bovis* and demonstrate that white yaks play an important part in the natural life cycles of *Anaplasma* spp. Further studies are needed to investigate the potential tick vectors of these pathogens in this alpine steppe ecoregion.

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