High diversity of *Rickettsia* spp., *Anaplasma* spp., and *Ehrlichia* spp. in ticks from Yunnan Province, Southwest China

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*Rickettsia*, *Anaplasma*, and *Ehrlichia* belonging to the order Rickettsiales are causative agents of tick-borne diseases in humans. During 2021, 434 ticks including *Rhipicephalus microplus* and *R. haemaphysaloides* were collected from three sampling sites in Yunnan Province, Southwest China, and analyzed for the presence of these bacteria. Nine bacterial species were identified, including two *Rickettsia* spp., three *Anaplasma* spp., and four *Ehrlichia* spp., some of which are potential human pathogens.

Genetic and phylogenetic analysis on 16S rRNA, *gltA*, *groEL*, *ompA*, *ompB*, and *sca4* genes indicated the presence of a novel spotted fever group *Rickettsia* (SFGR) named "*Candidatus Rickettsia shennongii*" in six of the 38 *R. haemaphysaloides* ticks from two locations, Dehong Autonomous Prefecture and Honghe City. Another SFGR species, *Candidatus Rickettsia jingxinensis* was detected in ticks from all three sites, with an overall positive rate of 62.67%. Three other human pathogenic species, *Anaplasma ovis* (1.38%, 6/434), *Ehrlichia canis* (16.36%, 71/434), and *E. chaffeensis* (0.23%, 1/434) were detected in these ticks and characterized. Moreover, *Ehrlichia* sp. (4.84%, 21/434), *E. minasensis* (7.37%, 32/434), *A. marginale* (6.91%, 30/434), and *Candidatus Anaplasma boleense* (1.15%, 5/434) were detected in *R. microplus* ticks, for which pathogenicity to humans remains to be determined. The results reveal the remarkable diversity of Rickettsiales bacteria in ticks from Yunnan Province, Southwest China. The high infection rate of some human pathogenic bacteria in ticks may indicate potential infection risk in humans, and it highlights the need for surveillance in local populations.

**KEYWORDS**

*Rickettsia*, anaplasma, ehrlichia, Yunnan Province, *Candidatus Rickettsia shennongii*
Introduction

Anaplasma spp. and Ehrlichia spp., belonging to order Rickettsiales are tick-borne intracellular bacteria, many of which are important pathogens of livestock. Anaplasma spp., Ehrlichia spp., as well as Rickettsia spp. bacteria also occasionally infect humans. Therefore, these bacteria are important for both veterinary and human public health (Gondard et al., 2017). SFGR represents a large group within the genus Rickettsia including Rickettsia rickettsii, R. conori, R. australis, R. honei, R. japonica, R. africæ, R. sibrica, etc. (Stenos et al., 2005). They are widely distributed and many of them are etiological agents of known human diseases, such as Rocky Mountain spotted fever (RMSF), A. phagocytophylum, Mediterranean spotted fever, Indian tick typhus, Queensland tick typhus, Flinders Island spotted fever, Japanese spotted fever, African tick bite fever, and North Asian tick-typus, etc. (Sentaursa et al., 2012; Graham et al., 2017; Rudakov et al., 2019; Zazueta et al., 2021). Of those, R. rickettsii mainly distributes in North and South America and was reported to infect more than 4,000 patients during 2009–2019 at Mexico-United States Border (Zazueta et al., 2021). From 1997 to 2004, 415 children cases of Mediterranean spotted fever caused by R. conori were recorded in Sicily, Italy (Colomba et al., 2006). Meanwhile, only sporadic infection cases were reported for R. australis, R. honei (in Australia), R. africæ (in Africa), R. japonica (Asia), and other SFG members (Parola et al., 2013). For the genus Anaplasma, Anaplasma marginale, A. centrale, and A. bovis are common pathogens causing severe or mild anaplasmosis in ruminants, especially cattle (Rjeibi et al., 2018). Meanwhile, A. phagocytophylum, A. bovis, and A. capra are confirmed human pathogens (Li H. et al., 2015; Ismail and McBride, 2017; Lu et al., 2019). In the past decades, a total of 110 infection cases of A. phagocytophylum were reported (Dumic et al., 2022). Among Ehrlichia species, E. ruminantium and E. minasensis are known to infect cattle, causing severe fever, anemia, and thrombocytopenia (Peter et al., 2020). Ehrlichia chaffeensis, E. ewingii, E. muris, E. muris. Subsp. caucalereisns, and E. canis are reported to infect humans, with syndromes ranging from febrile to multiple organ failure (Saito and Walker, 2016; Pritt et al., 2017). From 2012–2016, 6,786 cases of E. chaffeensis infection were reported in the United States (Mogg et al., 2020).

Located in southwest China, Yunnan Province covers a vast area with diverse climates and unique biodiversity resources. The extraordinary biological and ecological diversity of this area makes the extensive diversity of Rickettsiales bacteria possible. In recent decades, much attention has been paid to Rickettsiales bacteria circulating in this area. Numerous studies have been performed and many bacterial species belonging to the order Rickettsiales have been characterized in Yunnan Province (Liang et al., 2012; Liu et al., 2020; Jiao et al., 2021). One study revealed high positive rates of SFGR in domestic animals (goats, dogs, and cattle), and proved the existence of R. helongjiangensis and a distinct Rickettsia (Li et al., 2012). Another study reported the presence of R. raoultii and Ca. R. jingxinensis in ticks from Yunnan Province (Li et al., 2020). In Jiao et al. (2021) detected multiple tick-borne pathogens circulating in Rhipicephalus microplus ticks from Yunnan, including Ca. R. jingxinensis, A. marginale, and Coxiella burnetii. Furthermore, A. capra, A. phagocytophylum, and Candidatus Neoehrlichia mikurensis were also detected in R. microplus ticks from Yunnan Province (jiao et al., 2021). Although many studies on tick-borne Rickettsiales bacteria have been performed in Yunnan Province, further exploration is still needed. To improve our knowledge on the biodiversity and epidemiology of Rickettsiales bacteria, we collected ticks from goats and cattle in three locations of Yunnan Province, and characterized Rickettsia, Anaplasma, and Ehrlichia in them.

Materials and methods

Sample collection and processing

During 2021, ticks were collected from three locations in Yunnan Province: Ruili county-level city of Dehong Dai-Jingpo Autonomous Prefecture (97.85°E, 24.01°N), Zhaoyang District of Zaotong city (103.71°E, 24.32°N), and Shiping county of Honghe city (102.49°O, 23.71°N; Figure 1). Ticks were collected from cattle and goats, then brought to China CDC alive. Tick species were morphologically characterized by a sophisticated arthropod taxonomist based on the characteristics of the capitula, body, legs, anal groove, and caudal appendage (Namgyal et al., 2021). For further confirmation, randomly selected ticks were undergone molecular analysis by sequencing the mitochondrial cytochrome oxidase I (COI) gene sequences (Lu et al., 2013).

DNA extraction, PCR detection, and amplification of key genes

After washing three times with sterile phosphate-buffered saline (PBS), ticks were individually homogenized in 500μl of PBS.
using a Mixer Mill MM 400 (Retsch, Hann, Germany). The total DNA of each tick was extracted in 80 μl of eluate using a Mollusk DNA Extraction Kit (Omega Bio-Tek, United States) according to the manufacturer’s instructions.

Each tick DNA sample was screened by PCR for the presence of bacterial DNA including Rickettsia spp., Anaplasma spp., and Ehrlichia spp. Rickettsial DNA was detected by nested PCR as described previously using primers targeting the outer membrane protein A (ompA) gene, resulting in amplification of a 755-bp fragment (Lu et al., 2017). The DNA samples were also screened for Anaplasma and Ehrlichia DNA as described previously, amplifying a 550–850bp fragment of the 16S rRNA gene (Guo et al., 2016). Negative control with distilled water in the PCR master mixture and positive control (DNA of R. felis, A. marginale and E. canis, respectively) were included in each test. After 1% agarose gel electrophoresis, amplification products were subjected to sequencing.

For further characterization and phylogenetic analysis of the detected bacterial strains, the partial sequences of the gltA gene (citrate synthase), the groEL gene (heat shock protein), and a long fragment of the 16S rRNA gene were obtained for representative Rickettsia, Anaplasma, and Ehrlichia positive samples. Additionally, nearly complete sequences of ompB and sca4 genes were obtained for the putative novel Rickettsia species. The primers used were described previously (Roux and Raoult, 2000; Sekeyova et al., 2001; Kang et al., 2014; Guo et al., 2016; Lu et al., 2017) and they are listed in Supplementary Table S1. All recovered sequences have been deposited in the GenBank database (GenBank numbers listed in Supplementary Table S2).

Genetic and phylogenetic analysis of obtained sequences

Sequences obtained in this study were analyzed by BLASTn and Clustal W within Molecular Evolutionary Genetics Analysis (MEGA) software, version 7.0 (Kumar et al., 2016). Phylogenetic analysis was also performed using PhyML 3.0. Confidence values for each branch of the phylogenetic trees were determined by bootstrap analysis with 1,000 replicates. Sequences recovered in this study were aligned with the reference sequences retrieved from the GenBank database.

Results

Sample collection

During March to July 2021, 434 ticks (397 R. microplus and 37 R. haemaphysaloides) were collected from three locations in Yunnan Province. The three sampling locations (Ruili County-level City of Dehong Dai-Jingpo Autonomous Prefecture, Zhaoyang District of Zhaotong City, and Shiping County of Honghe City) were shown in Figure 1. Tick species were initially determined by morphological examination and then further determined by amplification and analysis of the COI gene (Lu et al., 2013). Except for four nymphs from Zhaotong, all ticks were adult, and most were fully or partially engorged. The COI sequences of ticks have been submitted to GenBank. The Accession numbers are OM959242-OM959313, OM959315-OM959321, and OM977035-OM977043. The tick species, quantity, and vertebrate hosts in different sampling sites were shown in Supplementary Table S3.

Rickettsia spp.

Nested PCR targeting the conserved domain of the ompA gene was performed to screen for Rickettsia in the collected tick samples. Based on agarose gel electrophoresis, DNA sequencing, and comparison with BLASTn, two Rickettsia species were initially identified: Ca. R. jingxinensis and a putative novel species. For further characterization, the 16S rRNA gene (1,172 bp), as well as the gltA (1,004 bp) and groEL (1,038–1,042 bp) genes were successfully amplified from the total tick DNA samples.

DNA of Ca. R. jingxinensis was detected in ticks from all three sites, with strikingly high positive rates varying from 37.50 to 84.69% (48 of 128 ticks from Dehong, 166 of 196 ticks from Zhaotong, and 70 of 110 ticks from Honghe; Table 1). All Ca. R. jingxinensis strains were detected in R. microplus ticks except for one from R. haemaphysaloides. All the ompA sequences shared 100% identity with Ca. R. jingxinensis isolate Meixian-Hl-107 (MH932061.1) and Xian-Hl-79 (MH932069.1) reported in Haemaphysalis longicornis from ShaanXi Province, Northern China (Guo et al., 2019). The 16S rRNA, gltA and groEL genes shared 99.90–100% identity with each other in reference to homologous genes. The 16S rRNA and gltA gene sequences were 100 and 99.90% identical to other Ca. R. jingxinensis strains from China. Meanwhile, the groEL sequences shared highest 99.90% identity with Uncultured Rickettsia sp. clone tick28 (ON409665) and Uncultured Rickettsia sp. clone tick26 (ON409664) we previously identified in Ngawa, Sichuan Province, which actually also represent Ca. R. jingxinensis strains.

Notably, a putative novel SFGR species was identified in R. haemaphysaloides ticks from Dehong (3/31, 9.68%) and Honghe (3/7, 42.86%; Table 1). The ompA sequences of these were 100% identical to each other, and they shared 98.31% identity with R. riofficephali str. 3-7-female6-CWPP (CP003342.1) and 98.17% with R. massiliiae MTU5 (CP000683.1). The 16S rRNA (1,172 bp) gene sequences of all strains from Dehong and Honghe were identical, despite the geographic separation, and they shared 100.0% identity with Rickettsia sp. strain HB-9543 N2 (MT433470), 99.91% identity with R. raoultii isolate Tomask (MK304546) and R. conorii str. Malish 7 (AE006914.1). The gltA (1,004 bp) gene sequences were found to be 99.70–99.80% identical to Rickettsia sp. strain HB-9543 N2 (MT433498), 99.30–99.40% to R. massiliiae MTU5 (CP000683.1) and R. riofficephali str. HJ#5 (CP013133). Regarding the groEL (1,042 bp) gene, all

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1 http://www.ncbi.nlm.nih.gov/BLAST
sequences shared high similarity with R. rhipicephali str. HJ#5 (CP013133, 99.42%) and R. rhipicephali str. 3-7-female6-CWPP (CP003342; 99.33%). For further confirmation, the ompB (4704–4707 bp) and sca4 (2560–2566 bp) genes were successfully recovered. Similar to other genes, the ompB gene shared 99.68–99.77% similarity to Rickettsia sp. strain HB-9543 N2 (MT434988) and 98.14–98.22% similarity with R. rhipicephali str. 3-7-female6-CWPP (CP003342.1), while the sca4 gene shared 99.73–99.96% identity with Rickettsia sp. strain HB-9543 N2 (MT434990) and 98.10–98.37% identity to R. rhipicephali str. HJ#5. Given all ticks in this study were collected from both Dehong and Honghe city, an additional six-nucleotide insertion (AAGAAA) in the groEL (854–1,206 bp) gene was detected, with a positive rate of 31.07% (32/103). The 16S rRNA, gltA (926 bp), groEL (857 bp), and ompB (2560–2566 bp) genes share 99.65–100.00% identity with previously reported R. microplus strains in the phylogenetic trees (von Fricken et al., 2021). All three genes (16S rRNA, 854–1,206 bp, gltA, 539 bp, groEL, 846 bp) are 100% identical to previously reported A. microplus strains, and they are closely clustered with other A. microplus strains in the phylogenetic trees (Figure 3). A widely distributed animal pathogen, A. marginale was detected in R. microplus ticks from all three locations, with prevalence rates from 4.08 to 15.53% (Table 1). All three genes (16S rRNA: 854–1,206 bp, gltA, 672 bp, groEL 857 bp) genes share 99.65–100.00% identity with Ca. A. boleense strains detected in other locations of China. Anaplasma ovis was detected in ticks from Dehong and Honghe, mostly from R. haemaphysaloides. All the three genes (16S rRNA, 854–1,206 bp, gltA, 539 bp, groEL, 846 bp) are 100% identical to previously reported A. ovis sequences, and they are closely clustered with other A. ovis strains in the phylogenetic trees (Figure 3). As a widely distributed animal pathogen, A. marginale was detected in R. microplus ticks from all three locations, with prevalence rates from 4.08 to 15.53% (Table 1). All three genes (16S rRNA: 854–1,206 bp, gltA, 672 bp, groEL 857 bp) share 100% identity with previously reported A. marginale sequences. To be noticed, it has been observed that the detection rates of Anaplasma spp. in ticks from the environment and those removed from livestock are different (von Fricken et al., 2021). Given all ticks in this study were removed from cattle and goats, the detection rates might be influenced.

**Ehrlichia spp.**

Of the 434 ticks screened from three locations, 125 (28.80%) were positive for Ehrlichia including four species: *E. minasensis*, *E. canis*, *E. chaffeensis*, and *Ehrlichia sp.* (Table 1). In *R. microplus* ticks from both Dehong and Honghe city, an *Ehrlichia* sp. species closely related to *Ehrlichia* sp. strain WHBMXZ-43 was identified, which was first identified in *R. microplus* ticks from Wuhan city, South-Central China. The 16S rRNA genes of the detected strains share 99.84–99.91% identity with this *Ehrlichia*, while both gltA and groEL genes share 100% identity with this strain.

In *R. microplus* ticks from Honghe city, *E. minasensis* was detected, with a positive rate of 31.07% (32/103). The 16S rRNA, *A. boleense* was detected in *R. microplus* ticks from Dehong, Zhaotong, and Honghe, with positive rates of 2.06% (2/97), 0.51% (1/196), and 1.94% (2/103), respectively (Table 1). The 16S rRNA (855 bp), gltA (672 bp), and groEL (857 bp) genes share 99.65–100.00% identity with *Ca. A. boleense* strains detected in other locations of China. Anaplasma ovis was detected in ticks from Dehong and Honghe, mostly from *R. haemaphysaloides*. All the three genes (16S rRNA, 854–1,206 bp, gltA, 539 bp, groEL, 846 bp) are 100% identical to previously reported *A. ovis* sequences, and they are closely clustered with other *A. ovis* strains in the phylogenetic trees (Figure 3). As a widely distributed animal pathogen, *A. marginale* was detected in *R. microplus* ticks from all three locations, with prevalence rates from 4.08 to 15.53% (Table 1). All three genes (16S rRNA: 854–1,206 bp, gltA, 539 bp, groEL 846 bp) share 100% identity with previously reported *A. marginale* sequences. To be noticed, it has been observed that the detection rates of *Anaplasma spp.* in ticks from the environment and those removed from livestock are different (von Fricken et al., 2021). Given all ticks in this study were removed from cattle and goats, the detection rates might be influenced.

### Table 1: Prevalence of Rickettsiales bacteria in ticks in Dehong, Zhaotong, and Honghe cities of Yunnan Province, Southwest China.

| City       | R. microplus (%) | R. haemaphysaloides (%) | Dehong       | Zhaotong       | Honghe       | Total (%) |
|------------|------------------|-------------------------|--------------|----------------|--------------|-----------|
| Dehong     | 48/97 (49.48)    | 0/31 (0.00)             | 166/196 (84.69) | 69/103 (66.99) | 1/7 (14.29) | 284/434 (65.44) |
| Zhaotong   | 0/97 (0.00)      | 3/31 (9.68)             | 0/196 (0.00)  | 0/103 (0.00)   | 3/7 (42.86) | 6/434 (1.38)    |
| Honghe     | 2/97 (2.06)      | 0/31 (0.00)             | 1/196 (0.51)  | 2/103 (1.94)   | 0/7 (0.00)  | 5/434 (1.15)    |

*a* Positive samples/total samples.

*b* Fifty-four samples were co-infected with *A. marginale* and four were co-infected with *Ehrlichia* sp.

*c* Nine were co-infected with *E. minasensis*, two were co-infected with *Ehrlichia* sp., and nine were co-infected with both *E. minasensis* and *Ehrlichia* sp.

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**Anaplasma spp.**

Three *Anaplasma* species were identified; *Candidatus Anaplasma boleense, A. ovis*, and *A. marginale*. DNA of *Ca. A. boleense* was detected in *R. microplus* ticks from Dehong, Zhaotong, and Honghe, with positive rates of 2.06% (2/97), 0.51% (1/196), and 1.94% (2/103), respectively (Table 1). The 16S rRNA (855 bp), gltA (672 bp), and groEL (857 bp) genes share 99.65–100.00% identity with *Ca. A. boleense* strains detected in other locations of China. Anaplasma ovis was detected in ticks from Dehong and Honghe, mostly from *R. haemaphysaloides*. All the three genes (16S rRNA, 854–1,206 bp, gltA, 539 bp, groEL, 846 bp) are 100% identical to previously reported *A. ovis* sequences, and they are closely clustered with other *A. ovis* strains in the phylogenetic trees (Figure 3). As a widely distributed animal pathogen, *A. marginale* was detected in *R. microplus* ticks from all three locations, with prevalence rates from 4.08 to 15.53% (Table 1). All three genes (16S rRNA: 854–1,206 bp, gltA, 539 bp, groEL 846 bp) share 100% identity with previously reported *A. marginale* sequences. To be noticed, it has been observed that the detection rates of *Anaplasma spp.* in ticks from the environment and those removed from livestock are different (von Fricken et al., 2021). Given all ticks in this study were removed from cattle and goats, the detection rates might be influenced.

**Ehrlichia spp.**

Of the 434 ticks screened from three locations, 125 (28.80%) were positive for *Ehrlichia* including four species: *E. minasensis*, *E. canis*, *E. chaffeensis*, and *Ehrlichia* sp. (Table 1). In *R. microplus* ticks from both Dehong and Honghe city, an *Ehrlichia* sp. species closely related to *Ehrlichia* sp. strain WHBMXZ-43 was identified, which was first identified in *R. microplus* ticks from Wuhan city, South-Central China. The 16S rRNA genes of the detected strains share 99.84–99.91% identity with this *Ehrlichia*, while both gltA and groEL genes share 100% identity with this strain.

In *R. microplus* ticks from Honghe city, *E. minasensis* was detected, with a positive rate of 31.07% (32/103). The 16S rRNA,
gltA, and groEL gene sequences all share 100% identity with *E. minasensis* strain B1 isolated from cattle in Brazil. In the phylogenetic trees based on these three genes, the two *E. minasensis* strains (Honghe-25 and Honghe-42) were both located in the same clade with *E. minasensis* strain B11 (Figure 4). This result indicated the prevalence and the high genetic conservation of *E. minasensis* in Yunnan Province.

Of the 196 *R. microplus* ticks tested from Zhaotong city, two *Ehrlichia* species were detected: *E. canis* (71/196, 36.22%) and *E. chaffeensis* (1/196, 0.51%; Table 1). The prevalence in male, female, and nymph ticks was shown in Supplementary Table S5. The 16S rRNA gene sequences of detected *E. canis* strains (1,406 bp) share 99.57% similarity with *E. canis* clone CuD125 (MK507008.1) and *E. canis* strain YZ-1 (CP025749.1), etc. The gltA and groEL sequences are 100% identical between the detected strains. The gltA (973 bp) sequences are 97.84% identical to *E. canis* strain YZ-1 (CP025749.1) and *E. canis* isolate T85D7 (MW382940.1), while the groEL sequences share 98.66% identity with *E. canis* strain Mossesane (MG953295.1). Phylogenetic analysis clearly revealed that all strains are assembled with other *E. canis* strains (Figure 4). Additionally, an *E. chaffeensis* strain was detected in a single *R. microplus* tick. All 16S rRNA, gltA, and groEL genes share 100% similarity with *E. chaffeensis* str. West Paces, *E. chaffeensis* Arkansas, and other previously reported *E. chaffeensis* strains. These appear to be the only gltA and groEL sequences of *E. chaffeensis* from China submitted to GenBank.

**Discussion**

Yunnan Province is a widely recognized biodiversity hotspot in China and worldwide (Li R. et al., 2015). In the present study, we revealed the extensive diversity and high positive rate of Rickettsiales bacteria in ticks from Yunnan Province. Nine
bacterial species belonging to the genus Rickettsia, Anaplasma, and Ehrlichia were detected, including a novel SFGR species. Our result may contribute to the current knowledge of the biodiversity of Rickettsiales bacteria circulating in this area.

Rickettsia spp. have been recognized as pathogens of potential public health importance. Herein, the DNA of Ca. R. jingxinensis was detected in all three locations with extremely high positive rates varying from 37.50 to 84.69%. Cadidatus Rickettsia jingxinensis is a SFGR first identified in Ha. longicornis from northeast China in 2016 (Liu et al., 2016). It has since been detected in ticks from multiple provinces in China, including Shaanxi, Guangxi, Sichuan, and Yunnan (Guo et al., 2019; Liu et al., 2020; Jiao et al., 2021; Lu et al., 2022). Furthermore, it was also reported in Korea, Thailand, and India, spanning from East Asia to South Asia (Takhampunya et al., 2019; Bang et al., 2021) (GenBank No. MN463681–MN463688). Previous studies revealed strikingly high positive rates of this Rickettsia in certain areas (for instance, 44.4–69.7% in H. longicornis from Shaanxi and 24.61% in R. microplus from Yunnan; Guo et al., 2019; Jiao et al., 2021). Our findings confirmed the widespread circulation of Ca. R. jingxinensis in China, as well as its high positive rate in ticks removed from livestock. Because most ticks were fully or partially engorged in the present work, it is not clear whether the Rickettsia DNA was from the blood meal or the tick itself. In another word, the possibility remains that the livestock in these locations may be infected by Ca. R. jingxinensis. These results remind us that the potential risk of Ca. R. jingxinensis infecting animals should be considered, and more attention should be paid to its role in human/animal diseases.

In R. haemaphysaloides ticks from Dehong and Honghe, a novel SFGR species named Ca. R. shennongii was identified. According to the gene sequence-based criteria for the identification of a new Rickettsia species (Fournier et al., 2003), genetic analysis of key genes clearly indicated that these strains represent a novel Rickettsia species. Sequences from all strains were almost identical, and they formed distinct clades in the phylogenetic trees. Genetically, this Rickettsia is closely related to R. raoultii, R. massiliae, and R. conorii (16S rRNA gene shares 99.91% identity with R. raoultii and R. conorii, while groEL gene shares 99.33% identity with R. massiliae), all of which are recognized human pathogens causing spotted fever. Therefore, the pathogenicity of this Rickettsia is a concern. Interestingly, this Rickettsia was only detected in R. haemaphysaloides, a widely distributed three-host tick in China and South Asian countries (Zhou et al., 2006). This species is known to harbor pathogens including A. phagocytophilum, A. ovis, R. rhipicephali, R. slovaca, R. massiliae, and Babesia microti (Kuo et al., 2018; Li et al., 2018; Ghafar et al., 2020; Ali et al., 2021). Our results may suggest the possibility of R. haemaphysaloides ticks harboring Ca. R. shennongii.
Herein, the DNA of three *Anaplasma* species was detected in all three locations including *Candidatus* *Anaplasma* boleense, *A. ovis*, and *A. marginale*. *Anaplasma ovis* is a widely-distributed pathogen affecting sheep, goats, and wild ruminants. As the etiological agent of ovine anaplasmosis first reported in 1956, the pathogenicity of *A. ovis* has been well studied since. *Anaplasma ovis* infection in sheep is usually subclinical and mostly manifested as hemolytic anemia (Dahmani et al., 2019). Other main clinical manifestations include extreme weakness, anorexia, and weight loss, but these manifestations mostly occur under poor health conditions (Bauer et al., 2021). In 2010, a human anaplasmosis case caused by an *A. ovis* variant was reported in Cyprus, with clinical symptoms of fever, hepatosplenomegaly, and enlarged lymph nodes (Chochlakis et al., 2010), providing evidence that *A. ovis* might be a potential zoonotic pathogen that may occasionally infect humans. In the present study, ticks from Dehong and Honghe were both tested positive for *A. ovis*, suggesting that *A. ovis* circulation is common in Yunnan Province, and surveillance in local populations is needed.

In this study, *Ca. A. boleense* was detected in ticks from all three sites of Yunnan Province. *Candidatus* *Anaplasma* boleense was first identified in ticks from Bole City in the Xinjiang Uygur Autonomous Region, Northwest China (Kang et al., 2014). In recent years, it has been detected in mosquitoes and rodents (FJ182047) in multiple provinces of China (Guo et al., 2016). In 2018, *Ca. A. boleense* was found to infect deer, boars, buffalos, cows, and bats in Peninsular Malaysia (Kob et al., 2018). In 2020, *Ca. A. boleense* was reported to infect marsh deer in Argentina, South America (Orozco et al., 2020). Furthermore, some sequences submitted to the GenBank database suggest that this *Anaplasma* exists in Australia (MH500004) and South Africa (MK814450), suggesting that it is distributed worldwide. However, few studies have been performed on the pathogenicity of *Ca. A. boleense* toward humans or other animals. The wide host range, as well as its wide geographical distribution, suggest that this *Anaplasma* species merits further investigation.

*Ehrlichia* is a genus closely related to human diseases. In this study, the DNA of 4 *Ehrlichia* species were identified, including *E. canis*, *E. chaffeensis*, *E. minasensis*, and *Ehrlichia sp.*, of which *E. canis* and *E. chaffeensis* are recognized human pathogens (Perez et al., 1996; Bouza-Mora et al., 2017). The most prevalent species was *E. canis* (36.22% positive rate in ticks from Zhaotong), the causative agent of canine monocytic ehrlichiosis. In China, *E. canis* has been detected in various hosts including ticks, goats, dogs, and deer in multiple locations (Xu et al., 2015; Li et al., 2016; Qiu et al., 2016; Yu et al., 2016; Lu et al., 2017; Zhang et al., 2017). Notably, *E. canis* is considered a potential agent of human disease (Perez et al., 1996; Bouza-Mora et al., 2017), although human infection cases have never been reported in China. As shown in Figure 2, the strains we detected are most closely related to *E. canis* strain WHBMXZ-124, a strain identified in *R. microplus* from Wuhan, China. However, considerable genetic distance exists between them, suggesting that they might represent a variant circulating in China. As a recently recognized species closely related to *E. canis*, *E. minasensis* is believed to have evolved from highly variable strains of *E. canis*, and it has been discovered in Canada, Brazil, France, Pakistan, Ethiopia, etc. (Li et al., 2019). Other than ticks, it also infects mammals including cattle and cervid. In China, there is only one report of *E. minasensis* in Hainan Province, South China (Li et al., 2019). In our current study, *E. minasensis* was detected in *R. microplus* ticks from Honghe city, with a high positive rate of 31.07%. To the best of our knowledge, this is the only report of *E. minasensis* circulating in China. Due to its pathogenicity to mammals, the close relationship to human pathogenic species *E. canis*, and the high infection rate in ticks, its pathogenicity to humans and animals should be further explored.

In summary, this study revealed substantial diversity of *Rickettsia*, *Ehrlichia*, and *Anaplasma* in ticks from Yunnan Province, including a novel *Rickettsia* species. Notably, some of these bacteria are human pathogens (*E. canis*, *E. chaffeensis*, *A. ovis*, etc) and the positive rates are high. Considering the frequent contact between humans and tick hosts (goats and cattle), these results indicate the potential risk of zoonosis transmitted from ticks to humans. Additionally, the human pathogenicity of *Ca. R. shennongii* should be further studied and surveillance in these areas is clearly needed.

**Data availability statement**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary material.

**Ethics statement**

This study was approved by the Ethics Committee of the National Institute for Communicable Disease Control and Prevention, Chinese Center for Disease Control and Prevention.

**Author contributions**

KL conceived the study. KL and ML designed the experiments. KL, JT, and JH collected the samples. ML and WW performed the experiments. HZ, KL, and WG performed data analysis. HZ, HJ, and KL wrote the manuscript. All authors contributed to the article and approved the submitted version.

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Conflict of interest
The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material
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