Genomic Characteristics of Emerging Intraerythrocytic Anaplasma capra and High Prevalence in Goats, China

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Anaplasma capra is an emerging tickborne human pathogen initially recognized in China in 2015; it has been reported in ticks and in a wide range of domestic and wild animals worldwide. We describe whole-genome sequences of 2 A. capra strains from metagenomic sequencing of purified erythrocytes from infected goats in China. The genome of A. capra was the smallest among members of the genus Anaplasma. The genomes of the 2 A. capra strains contained comparable G+C content and numbers of pseudogenes with intraerythrocytic Anaplasma species. The 2 A. capra strains had 54 unique genes. The prevalence of A. capra was high among goats in the 2 endemic areas. Phylogenetic analyses revealed that the A. capra strains detected in this study were basically classified into 2 subclusters with those previously detected in Asia. Our findings clarify details of the genomic characteristics of A. capra and shed light on its genetic diversity.

A naplasma capra is an emerging tickborne zoonotic pathogen in the genus Anaplasma, family Anaplasmataceae, and was initially identified in blood samples from asymptomatic goats (Capra

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Members of the family Anaplasmataceae have complex life cycles involving vertebrate hosts and hematophagous ticks, many of which have emerged as human pathogens. The genus Anaplasma was proposed according to the phylogenetic analyses based on 16S rRNA and groEL sequences (17) and initially encompassed 6 species: A. phagocytophilum, A. marginale, A. centrale, A. ovis, A. platys, and A. bovis. Subsequently, 2 candidate novel species (A. capra and A. odocoilei) and other unclassified genovariants (1,18-20) were included in the List of Prokaryotic Names with Standing in Nomenclature (https://www.bacterio. net) pending validation. To date, 5 Anaplasma species have been known to infect humans: A. phagocytophi*lum*, *A. capra*, *A. ovis*, *A. platys*, and *A. bovis* (21). Since the A. marginale genome sequence was reported in 2005 (22), a total of 24 A. marginale genomes (23), 32 A. phagocytophilum genomes (24,25), 1 A. centrale genome

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(26), 2 A. ovis genomes (27), and 1 A. platys genome (28) have been sequenced and deposited in GenBank. Although A. capra has been extensively detected in ticks and animal hosts worldwide, no genome of the emerging pathogen has been determined so far, which has hindered us from better understanding its genetic features and pathogenesis. Considering A. capra is an intraerythrocytic pathogen and abundant in blood samples of host goats (1,29), we separated erythrocytes from the blood of infected goats to enrich the bacteria and generated the entire genome of A. capra using metagenome assembly to promote better understanding of this emerging pathogen, to compare the characteristics of A. capra genomes with previously published genomes of other Anaplasma and related species, and to evaluate intraspecies genetic diversity of A. capra in different geographic locations and tick species across China.

Materials and Methods

Sample Collection and Preparation

We collected EDTA blood samples from 3 flocks of goats in Shandong Province and a flock of goats in Guizhou Province, China (Appendix Figure 1, https://wwwnc.cdc.gov/EID/article/29/9/23-0131-App1.pdf), during September 2021-July 2022. Meanwhile, we prepared blood smears for some goats. We collected host-seeking ticks in the same areas where the infected goats lived by dragging white flags over vegetation. An entomologist (Y.S.) identified all ticks to the species level and developmental stage. We extracted DNA from each goat blood sample or tick by using a High Pure PCR Template Preparation Kit (Roche, https://www.roche.com) according to the manufacturer's instructions.

PCRs and Sequencing

We conducted a nested PCR specific for the citrate synthase (*gltA*) gene of *A. capra* (Appendix Table 1) to screen all goat blood and tick samples, as previously described (1). We amplified all the positive samples for *gltA* by specific PCRs targeting the 16S rRNA, *msp4*, and *groEL* genes of *A. capra* (Appendix Table 1). We sequenced all amplicons to confirm the correctness of PCR results and conducted a SYBR Greenbased quantitative PCR (qPCR) targeting different regions of the *gltA* gene by using a specific primer (Appendix Table 1).

Fluorescence In Situ Hybridization

We used fluorescence in situ hybridization (FISH) to observe the *A. capra* on blood smears. We designed

the probe on the basis of the 16S rRNA full-length sequence of *A. capra* (Appendix Table 2) and labeled it with Quasar 570. We resuspended the pooled FISH probes in a final concentration of 25 μ mol/L in RNase-free storage buffer, which we protected from light and stored at -20°C. We performed FISH on the prepared blood smear with a commercial kit (Biosearch Technologies, https://www.biosearchtech. com), according to the manufacturer's instructions.

Enrichment of A. capra for Genomic Sequencing

We separated erythrocytes from infected goats by conducting gradient centrifugation using cell separation solution (Eppendorf, https://www.eppendorf. com) for 20 min at 200 \times g at 4°C. Then, we added 4 times volume of precooled (4°C) erythrocyte lysis buffer (Solarbio, http://www.solarbio.net) to the isolated erythrocytes by gentle pipetting to ensure adequate mixing. After placing the lysis solution at 4°C for 10 min, we centrifuged the solution at $350 \times g$ for 10 min to remove residual blood cells. After that treatment, we maximally removed the host DNA in samples. Finally, we centrifuged the supernatant at 20,000 \times g at 4°C for 30 min. We resuspended the pooled A. capra for DNA extraction by using the High Pure PCR Template Preparation Kit (Roche). We then constructed a sequencing library by using the AxyPrep MAG PCR Clean Up Kit (Fisher Scientific, https://www. fishersci.com) for an MGI sequencing set (https:// en.mgi-tech.com). We prepared the sequencing library according to the Whole Genome Sequencing Library Preparation Protocol (MGI). We sequenced the paired-end libraries with a read length of 2×150 bp on a DNBseq-T7 platform at Grandomics Gene Technology Beijing Co. Ltd (Beijing, China).

Genome Assembly and Comparative Analyses

We mapped the clean reads to the goat (Capra hircus) reference genome (GenBank accession no. GCF_001704415) by using SAMtools 1.14 (30) to discard host-derived reads. We de novo assembled contigs from the unmapped reads by using metaSPAdes 3.15.3 (31). We performed contig binning by using MetaBAT 2.15 (32) and evaluated assembly quality by using CheckM version 1.1.3 in linage_wf mode, which searches for universal single-copy marker genes and deduces completeness and contamination on the basis of presence and absence of these genes (33). We generated G+C content, genome completeness, and annotation information and depicted them by using an approach described previously (34,35). We estimated average nucleotide identity (ANI) and DNA-DNA

hybridization (DDH) by using fastANI 1.32 (36) and GGDC (https://ggdc.dsmz.de/ggdc.php).

Phylogenetic Analyses

We deposited in GenBank the results of the phylogenetic analysis of the whole genomes of the 2 *A. capra* strains and all the genomes of *Anaplasma* species by using Orthofinder 2.5.4 (*37*), after eliminating the poorly aligned positions and divergent regions by using Gblocks 0.91b. We aligned trimmed sequence by using Muscle 5.1 (R.C. Edgar, unpub. data, https:// doi.org/10.1101/2021.06.20.449169) and constructed the phylogenetic tree by using iqtree 2.2.0.3 (*38*). Furthermore, we conducted phylogenetic analyses on *A. capra gltA, groEL*, 16S rRNA, and *msp4* genes obtained from infected goats and ticks by using the maximumlikelihood method in MEGA11 (*39*).

Functional Analysis of Predicted Genes

To find difference in the Kyoto Encyclopedia of Genes and Genomes (KEGG) between the 2 strains of *A. capra* and other species in the genus *Anaplasma*, were annotated orthogroup sequences by using KOfam 1.4.0 (40) and illustrated them using a Venn diagram. We used the software eggNOG-Mapper 2.1.7 to determine the Clusters of Orthologous Group (COG) categories for protein encoding regions (41).

Results

Forty-three (59.7%) of 72 goat blood samples were positive for *gltA* gene of *A. capra*. We chose 2 blood samples (1 from a 2-year-old female goat in Shandong Province and another from a 10-month-old female goat in Guizhou Province) (Appendix Figure 1) for next-generation sequencing because they had high bacterial loads (8.4×10^6 *gltA* gene copies/mL blood for the goat in Shandong Province and 2.0×10^6 *gltA* gene copies/mL blood for the goat in Guizhou Province) as estimated by qPCR (Appendix Table 1). In addition, we visualized *A. capra* by specific FISH in erythrocytes on the blood smear prepared from the goat in Shandong Province for next-generation sequencing (Figure 1).

The metagenome sequencing resulted in >38 million 150-bp clean reads from each sample. Despite primary removing of host DNA, 95.9% and 93.3% of reads in the 2 samples were mapped to the goat genome and discarded. The remaining reads were subsequently de novo assembled into contigs by using the SPAdes 3.15.3 with meta parameters (*31*). The 2 assembled *A. capra* genomes were named *A. capra* str. BIME1 (GenBank accession no. GCA_025628785.1) and *A. capra* str. BIME2 (GenBank accession no. GCA_025628805.1), and had a higher level of completeness (99.79% for BIME1 and 99.36% for BIME2). The genome of *A. capra* was the smallest (\approx 1.07 Mb) among those in the genus *Anaplasma* and the second smallest genome of the family Anaplasmataceae, just after *Neorickettsia sennetsu* (0.859 Mb) (24). The genome sequences of the 2 strains shared 99.89% nucleotide similarity with each other.

We compared the 2 A. capra genomes with other representative species strains in the genus Anaplasma (Appendix Table 3). The G+C content (48.3% for both) of the 2 A. capra genomes was similar to those of A. ovis, A. marginale, and A. centrale, which are all intraerythrocytic pathogens. The A. capra genomes yielded a total of 929 and 932 genes, of which 862 and 863, respectively, represented coding sequences. They possessed 37 tRNAs and a complete ribosomal RNA operon, in which the 16S rRNA gene was separated from the 23S-5S rRNA gene pair (Figure 2) as displayed by other members of the order Rickettsiales (42). The 2 strains of A. capra and other intraerythrocytic Anaplasma species, including A. ovis, A. centrale, and A. marginale, contained comparable numbers of pseudogenes that have lost functions owing to mutation accumulation and are observed more frequently in obligate intracellular bacteria where the lost gene functions are compensated by the host cells (43). Of note, A. phagocytophilum has ≈4-fold more pseudogenes than the other Anaplasma species (Appendix Table 3).

The estimated values of ANI and DDH between A. *capra* and other *Anaplasma* species suggested that A. *capra* were distinct from the other species. On the basis of ANI values, A. capra str. BIME1 was most similar to A. marginale, whereas A. capra str. BIME2 was most similar to A. ovis. The DDH results revealed that both A. *capra* strains were most close to *A*. *marginale* (Appendix Table 4). The phylogenetic analysis based on the single copy genes revealed that the 2 A. capra strains together occupied a distinct branch and were more closely related to A. ovis, A. marginale, and A. centrale than to A. phagocytophilum and A. platys in the genus Anaplasma (Figure 3, panel A). To explore the gene differences in species in the genus Anaplasma, we used Orthofinder (37) to identify the homologous genes. All species in the genus Anaplasma shared 643 genes in common, and the 2 A. capra strains together with other intraerythrocytic Anaplasma species (A. ovis, A. centrale, and A. marginale) shared 75 genes that are not present in the other 2 species, A. phagocytophilum and A. platys. Compared with other members of the genus Anaplasma, 14 genes were not possessed by A. capra. Of note, a total of 54 genes were only shared by the 2 A. capra strains, which had other 14 distinct genes in BIME1 and 10 in



Figure 1. Anaplasma capra in the erythrocytes of an infected goat detected by fluorescence in situ hvbridization (FISH) in study of emerging intraerythrocytic A. capra and high prevalence in goats, China. Glowing red indicates A. capra; blue indicates leukocyte nucleus stained with fluorescent antibody blocker containing DAPI. A) FISH results under fluorescence microscope of A. capra. B) FISH results of A. capra-negative blood smear. C) FISH results showing different shapes and sizes of A. capra in erythrocytes.

BIME2 (Figure 3, panel B). In addition, we identified 25 virulent genes in the 2 *A. capra* strains that were shared by all the species in the genus of *Anaplasma*, including *virB2* gene family, *virB6* gene family, *virB4* gene family, *virB8* gene family, *virB9* gene family, and *virB3*, *virB7*, *virB10*, *virB11*, *virD4*, and Ats-1 genes that encode the type 4 secretion system and membrane protein-encoding genes (Appendix Table 5).

Among the 54 unique genes of *A. capra*, a total of 37 were unclassified, none of which was assigned to any KEGG category. Six of the remaining 17 genes were associated with metabolic processing, 5 genes were related to genetic information processing, and 6 were involved in signaling and cellular processing (Appendix Table 6). Among them, the most noteworthy of genes were *RSF1*, a gene related to the repair of DNA double-strand breaks (44), and *desk*, which

encodes a protein acting as a kinase at cold temperatures in *Bacillus subtilis* (45).

We classified the coding proteins of the 2 *A. capra* strains (BIME1 and BIME2) into functional clusters of orthologous group (COG) categories and compared them with those of representative species strains in the genus *Anaplasma* (Appendix Table 7). Most proteins were involved in translation, ribosomal structure and biogenesis, energy production and conversion, and nutrient (including amino acid, nucleotide, carbohydrate, coenzyme, and lipid) transport and metabolism, all of which were essential for bacterial survival. Of note, the number of genes encoding cell wall and membrane in *A. platys* was substantially lower than those of other *Anaplasma* species. In addition, ~10% of the proteins did not assign to any COG category and were classified as function unknown in each species.

RESEARCH



Figure 2. Circular map of *Anaplasma capra* strains BIME1 and BIME2 genomes in study of emerging intraerythrocytic *A. capra* and high prevalence in goats, China. The outermost ring shows the genome size in 100-kb increments. Moving inward, the blue-green and red marks indicate the coding sequences on the reverse and forward strands. The fourth ring represents the sequencing depth. The fifth ring shows the G+C skew, and the sixth rings show and G+C content. The location of *groEL and gltA* genes and the complete ribosomal RNA genes (5S rRNA, 16S rRNA, and 23S rRNA) within the genome are indicated.

We screened blood samples from 3 flocks of 54 goats in Shandong Province and a flock of 18 goats in Guizhou Province (Appendix Figure 1) by using nested PCR and qPCR targeting different regions of the gltA gene (Appendix Table 1). The overall positive rate was 59.7% (95% CI 48.4%-71.0%), and the positive rate was significantly higher among goats in Guizhou Province than in Shandong Province (77.8% vs. 53.7%; p<0.001). Accordingly, among the *H. longicornis* ticks collected from the same sites of the positive goats, the overall positive rate was 8.0% (95% CI 4.2%-11.8%), and the A. capra infection rate was significantly higher among ticks in Guizhou Province than that in Shandong Province (15.8% vs. 4.9%; p<0.001) (Appendix Table 8). To understand the genetic diversity, we amplified A. capra 16S rRNA (1,500 bp), groEL (1264 bp), and *msp4* (799 bp) genes from those positive samples. We compared the nucleotide identities for each gene sequence and (Appendix Figures 2-5; GenBank accession numbers are provided).

The *gltA* genes amplified from either goats or ticks in this study had 99.7%–100% identity with each other and with the strain that infected humans (Appendix Figure 2). The phylogenetic analysis based on *gltA* gene revealed that the *A. capra* sequences in this study were in an independent cluster from those previously reported in various animals from China and South Korea but distinct from those detected in wild and domestic animals from Europe and Kyrgystan. The South Korea water deer seemed to be capable of carrying both variants of A. capra (Figure 4, panel A). No A. capra groEL gene was acquired from tick samples, and the sequences from goats shared 99.4%-100% identity with each other and 99.8%-100% with sequences from humans (Appendix Figure 3). Similarly, the phylogenetic analyses based on the groEL gene revealed that A. capra strains of this study clustered with those from humans, dogs, and domestic ruminants in Asia but were distinguished from those in Europe (Figure 4, panel B). The entire 16S rRNA gene sequences (1,500 bp) of A. capra detected in goats and *H. longicornis* ticks from either Shandong or Guizhou Province shared average similarity of >99.7% from each other and from the sequence detected in humans (Appendix Figure 4). The phylogenetic tree based on 16S rRNA gene sequences indicated that all the A. capra strains detected in this study were in the same clade with previously reported strains in Asia (Figure 4, panel C). The A. capra msp4 gene sequences were also relatively conserved (Appendix Figure 5) among the goats and ticks, and the topology of phylogenetic tree based on *msp4* gene were similar to that based on the 16S rRNA gene, in which all A. capra sequences clustered in the clade different from other members of Ana*plasma* species (Figure 4, panel D).

Discussion

Whole-genome assembly of obligate intracellular bacteria has usually been hindered by the DNA presence of host cells. In this study, we first assembled 2 complete genomes of A. capra from the red blood cells of infected goats by using the metagenomic sequencing strategy. Because A. capra is an intraerythrocytic pathogen (1,29), we separated erythrocytes from the periphery blood of the infected goats and then lysed them for maximum removal of goat DNA. After metagenomic next-generation sequencing, we discarded the remaining goat genomic sequences and successfully assembled the A. capra genomes from 2 infected goats. The high percentage of reads from goat could be attributable to the low abundance of A. capra in erythrocytes or the fact that all other host cells rather than erythrocytes were not totally removed during the isolation of erythrocytes. In any case, the completeness of the 2 A. capra genomes are up to 99.79% for BIME1 and 99.36% for BIME2. The genome sizes obtained in this study reach 1,066,874 bp for BIME1 and 1,059,758 bp for BIME2. Therefore, their predicted sizes are ≈1.07 Mbp, which remain the smallest genome in the genus of Anaplasma. The phylogenetic analysis based on genome sequences and the comparative analyses of genomic characteristics provide the evidence that A. capra is closely related to other intraerythrocytic Anaplasma species, including A. ovis, A. centrale, and A. marginale.

The genome of *A. capra* consists of a single circular chromosome with a total size of 1.07 Mbp and has

862 protein-coding genes, which is smaller than other Anaplasma species. In fact, all the Anaplasma genomes sequenced so far are relatively small compared with free-living bacteria. The small genome size might be because a part of the intracellular bacterial functions has been compensated by the host cells, a process of reductive evolution that has occurred in the order Rickettsiales because of long-term intracellular association with eukaryotic hosts (46). This reductive evolution is associated with the frequent formation of pseudogenes, affecting distinct loci in different species (47). Moreover, we found that the G+C content of A. capra is close to that of A. ovis, A. marginale, and A. centrale. Of note, their relatedness also seems to be closest according to the phylogenetic analysis. The common invasiveness of erythrocytes also accounts for their high similarity.

A limitation of this study is that both the *A. capra* genomes were directly derived from the blood samples of infected goats through metagenomic next-generation sequencing. Unfortunately, we did not obtain the genomes at chromosome level, which usually relies on 3rd-generation sequencing of an isolate. In any case, this study reveals the genomic characteristics of *A. capra* and sheds light on its genetic diversity.

The high prevalence of *A. capra* in goats from Shandong and Guizhou Provinces in this study



Figure 3. Phylogenetic tree and genomic comparison among *Anaplasma* species in study of emerging intraerythrocytic *A. capra* and high prevalence in goats, China. A) Phylogenetic tree of *Anaplasma* species based on all the genomic sequences deposited in GenBank, constructed by using maximum-likelihood method with *Ehrlichia chaffeensis* as an outgroup. The percentages of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) are shown next to the branches. B) Differences in gene contents among *Anaplasma* species strains. Venn diagrams show the distribution of shared and unique gene clusters among representative *Anaplasma* species.

RESEARCH

further indicate that domestic ruminants might be the main animal hosts, as suggested by previous studies (2–5). *H. longicornis* ticks collected from the same sites of the positive goats either in Shandong Province or Guizhou Province are naturally infected with *A. capra*,

implying the role of the tick species in transmission of the pathogen. Phylogenetic analyses based on the *gltA* and *groEL* genes demonstrate that *A. capra* strains detected from goats and *H. longicornis* ticks in this study are clustered in the same clade with those



Figure 4. Phylogenetic analysis of *Anaplasma capra* based on nucleotide sequences of 4 genes in study of emerging intraerythrocytic *A. capra* and high prevalence in goats, China. A) Phylogenetic tree based on 536 bp nucleotide sequence of *gltA*. B) Phylogenetic tree based on 620 bp nucleotide sequence of *groEL*. C) Phylogenetic tree based on 860 bp nucleotide sequence of 16S rRNA. D) Phylogenetic tree based on 642 bp nucleotide sequence of *msp4*. We performed bootstrap analysis of 1,000 replicates to assess the reliability of the reconstructed phylogeneties. GenBank accession numbers are provided. Scale bars show estimated evolutionary distance.

from humans, domestic ruminants, dogs, and Korean water deer (2,3,5,10). Of note, another clade of *A. cap-ra* strains is mainly found in the wild and domestic animals from Europe and Kyrgyzstan (6,10,48). Those findings suggest that the enzootic cycles in various regions of the world might be different. Public health professionals should pay enough attention and formulate prevention and control strategies to reduce the health threat of the emerging tickborne pathogen to humans in other countries besides China.

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Genomic Characteristics of Emerging Intraerythrocytic *Anaplasma capra* and High Prevalence in Goats, China

Appendix

Appendix Table	1 Nucleotide sec	wence of primers	used in the study
Appendix Table	1. Nucleotide 360	fuction of primers	used in the study

	Primer		Annealing		
Target	name	Primer sequence (5'-3')	temperature (°C)	Amplicon size (bp)	References
rrs	Eh-out1	TTGAGAGTTTGATCCTGGCTCAGAACG	50	1500	(1)
	3–17U	WAAGGWGGTAATCCAGC			
gltA	Outer-f	GCGATTTTAGAGTGYGGAGATTG	55	1076	(2)
	Outer-r	TACAATACCGGAGTAAAAGTCAA			
	Inner-f	GGGTTCMTGTCYACTGCTGCGTG	51	792	(2)
	Inner-r	TTGGATCGTARTTCTTGTAGACC			
groEL	Forward	GCGAGGCGTTAGACAAGTCCATT	56	1264	(2)
	Reverse	TCCAGAGATGCAAGCGTGTATAG			
msp4	Forward	CAGTCTGCGCCTGCTCCCTAC	55	799	(2)
	Reverse	AGGAATCTTGCTCCAAGGTTA			
msp2	Forward	GCGTGTTGATGGCTCTGGT	51	1139	(2)
	Reverse	ACCAGTATCCTTATTTTTACC			
gltA*	Forward	CGAATCTATTTGCCTGCTT	60	200	This study
	Reverse	ATCGTAATTCTTGTAGACCCT			-

*This pair of primers is used for the quantitative PCR.

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Appendix Ta	ble 2.	Probe sec	uences of fluorescence	in situ h	ybridization	(FISH)
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Probe sequence (5' to 3')	Probe sequence name
TTCTGAGCCAGGATCAAACT	AC16S-1
TCGACTTGCATGTGTTAAGC	AC16S-2
AGCAAGCTACAGATTTGGTC	AC16S-3
CGTCTGCCACTAACCAAATC	AC16S-4
AGATTCCTATGCATTACTCA	AC16S-5
TGGCTATCCCATACTACTAG	AC16S-6
GGATTATACGGTATTACCCA	AC16S-7
ATAGCGATAAATCTTTCCCC	AC16S-8
CCAACTAGCTAATCCGACAT	AC16S-9
ACAGATCACTGCCTTGGTAG	AC16S-10
TGATCATCCTCTCAGACCAG	AC16S-11
CATTGTCCAATATTCCCCCAC	AC16S-12
CATAGCTGGATCAGGCTTGC	AC16S-13
TTTTACAACCCTAAGGCCTT	AC16S-14
TCATTATCTTCCCTACTGAA	AC16S-15
GGGACTTCTTCTGTAGGTAC	AC16S-16
CGCCCAATAATTCCGAACAA	AC165-10
ΤΤΑΛΟΤΤΑΟΟΑΛΑΛΟΟΑ	AC165-18
GTTAAGCCCTGGTATTTCAC	AC165-10
	AC165-20
	AC165-20 AC165-21
	AC165 22
	AC105-22
	AC105-23
	AC105-24
	AC105-25
	AC105-20
	AC105-27
	AC105-20
	AC105-29
	AC 105-30
ATCTAACCTCCATGTCAAGA	AC165-31
AACIGCGCCTTTCTGTTAAG	AC165-32
	AC165-33
	AC16S-34
AIGAGGGIIACGCICGIIGC	AC16S-35
	AC16S-36
CACCGGCAGTTTCCTTAAAG	AC16S-37
CGTGCTGACTTGACATCATC	AC16S-38
CATTGTAGCACGTGTGTAGC	AC16S-39
CGACGTTGCAACCTATTGTG	AC16S-40
CTTTTACGGATTAGCTCAGC	AC16S-41
CTCGAGTTGCAGAGGACAAT	AC16S-42
TCCACGATTACTAGCGATTC	AC16S-43
CGAGAACGTATTCACCGTGG	AC16S-44
TGACGGGCAGTGTGTACAAG	AC16S-45
TTTGAGTTAAGCCAATTCCC	AC16S-46
CACCGACCCAACCTTAAATG	AC16S-47
TACAGCTACCTTGTTACGAC	AC16S-48

	Strain (GenBank accession no.)								
	A. marginale str.								
	A. capra str. BIME1	A. capra str. BIME2	A. ovis str. Haibei	A. centrale str. Israel	Florida	A. platys str. S3	A. phagocytophilum		
Characteristic	(GCA_025628785.1)	(GCA_025628805.1)	(NZ_CP015994.1)	(NC_013532.1)	(NC_012026.1)	(NZ_CP046391.1)	str. JM (NC_021880)		
Size (bp)	1,066,874	1,059,758	1,214,674	1,206,806	1,202,435	1,196,811	1,481,598		
GC rate (%)	48.32	48.32	48.9	50.0	49.8	45.5	41.6		
Gene counts (n)	929	932	1021	993	992	940	1155		
CDS counts (n)	862	863	945	922	913	882	997		
Pseudogenes (n)	27	29	32	27	35	17	114		
rRNAs (n)	3	3	3	3	3	3	3		
tRNAs (n)	37	37	37	37	37	34	37		
Completeness (%)	99.79	99.36	NA	NA	NA	NA	NA		

Appendix Table 3. Genomic characteristics of Anaplasma capra strains BIME1 and BIME2 compared with that of representative Anaplasma species strains*

*bp, base pair; CDS, coding sequence; rRNA, ribosomal ribonucleic acid; tRNA, transfer ribonucleic acid; NA, not applicable.

Appendix Table 4. The estimated values of average nucleotide identity (ANI) and DNA-DNA hybridization (DDH) between
Anaplasma capra and the other Anaplasma species

_	A	NI	DDH		
Species	A. capra str. BIME1	A. capra str. BIME2	A. capra str. BIME1	A. capra str. BIME2	
A. ovis	78.0783	78.0878	17.4	17.5	
	(GCA_002849345.1)	(GCA_002214625.1)	(GCA_002849345.1)	(GCA_002849345.1)	
A. marginale	78.2897	77.9471	17.9	17.9	
	(GCA_008801305.1)	(GCA_000020305.1)	(GCA_000172475.1)	(GCA_000172475.1)	
A. centrale	77.9688	77.8613	17.4	17.4	
	(GCA_000024505.1)	(GCA_000024505.1)	(GCA_000024505.1)	(GCA_000024505.1)	
A. phagocytophilum	(-)	(-)	13.0	13.0	
			(GCA_023476575.1)	(GCA_023278635.1)	
A. platys	(-)	(-)	13.1	13.1	
			(GCA_012790675.1)	(GCA_012790675.1)	

Appendix Table 5. Virulence genes in Anaplasma capra str. BIME1 and BIME2

		GenBank accession number		
Gene	Description	A. capra str. BIME1	A. capra str. BIME2	
virB2	type IV secretion system protein VirB2 family	MCU7611221.1	MCU7612774.1	
		MCU7611222.1	MCU7612775.1	
		MCU7611775.1	MCU7612776.1	
		MCU7611780.1		
		MCU7611781.1		
		MCU7611782.1		
virB3	type IV secretion system protein VirB3	MCU7611541.1	MCU7612020.1	
virB4	type IV secretion system protein VirB4 family	MCU7611542.1	MCU7612019.1	
		MCU7611779.1	MCU7612773.1	
virB6	type IV secretion system protein VirB6 family	MCU7611543.1	MCU7612018.1	
		MCU7611544.1	MCU7612017.1	
		MCU7611545.1	MCU7612016.1	
		MCU7611546.1	MCU7612015.1	
virB7	type IV secretion system protein VirB7	MCU7611364.1	MCU7612438.1	
virB8	type IV secretion system protein VirB8 family	MCU7611203.1	MCU7612293.1	
		MCU7611581.1	MCU7611980.1	
virB9	type IV secretion system protein VirB9 family	MCU7611202.1	MCU7612294.1	
		MCU7611762.1	MCU7612488.1	
virB10	type IV secretion system protein VirB10	MCU7611201.1	MCU7612295.1	
virB11	type IV secretion system ATPase VirB11	MCU7611200.1	MCU7612296.1	
virD4	type IV secretion system component VirD4	MCU7611199.1	MCU7612297.1	
Ats-1	Anaplasma T4SS translocated substrate-1	MCU7611426.1	MCU7612135.1	
ompA	outer membrane protein OmpA	MCU7611514.1	MCU7612047.1	
Asp14	14-kDa <i>Anaplasma</i> surface protein Asp14	MCU7611843.1	MCU7612563.1	

Appendix Table 6. Genes predicted to be unique in Anaplasma capra str. BIME1 and BIME2

Gene	Protein	F	unction
menA	1,4-dihydroxy-2-naphthoate polyprenyltransferase	Metabolic processing	Menaquinone (vitamin K2) biosynthesis
unknown <i>MKK</i> 9	Glycosyltransferase 2 family protein Mitogen-activated protein kinase kinase 9		Glycan metabolism Ethylene and camalexin biosynthesis
MqnX CPS1 atuF Zbtb46	Aminodeoxyfutalosine deaminase Peregrinol diphosphate synthase Geranyl-CoA carboxylase α subunit Zinc finger and BTB domain-containing protein	Genetic information	Menaquinone Biosynthesis Metabolism Geraniol degradation Transcription factor
UFL1 Hmbox1 RSF1 Ara54	46 E3 UFM1-protein ligase 1 Homeobox-containing protein 1 Remodeling and spacing factor 1 E3 ubiquitin-protein ligase	processing	Cellular regulation Transcription factor DNA repair Cellular regulation

Gene	Protein		Function
MACC1	Metastasis-associated in colon cancer protein 1	Signaling and cellular processing	Signaling regulator
dia	Diaphanous protein		Cytokinesis
desK	Membrane-associated kinase DesK		Membrane-associated kinase
hbhA	Heparin binding hemagglutinin		Virulence factor
Mrgprg	Mas-related G protein-coupled receptor member G		G protein-coupled receptor
unknown	Membrane protein		Protein with domain of unknown function
unknown*	unclassified protein	Func	tion unknown
*Including 37 ur	nclassified genes.		

Appendix Table 7. Functional Clusters of Orthologous Groups of protein-coding genes from the representative *Anaplasma* species strains

		A	A	Antia	Assertusia	A	A	A when we as the while we
	Europhian al	A. capra	A. capra	A. OVIS	A. centrale	A. marginale	A. platys	A. pnagocytopnilum
Cotogony	Functional	SUL DIVIE I	SU. DIVIEZ	str. naibei	Str. Israel	Str. Florida	Str. 55	SUL JIVI
Category	category	4	4	4		genes	4	4
A	RNA	1	1	1	1	1	1	1
	processing and							
-	modification	•	•	•	0	•	0	•
В	Chromatin	0	0	0	0	0	0	0
	structure and							
-	dynamics							
С	Energy	67	67	74	70	68	71	74
	production and							
_	conversion							
D	Cell cycle	16	15	18	17	18	11	14
	control, cell							
	division,							
	chromosome							
	partitioning							
E	Amino acid	31	31	35	34	35	30	25
	transport and							
	metabolism							
F	Nucleotide	54	54	55	54	54	52	56
	transport and							
	metabolism							
G	Carbohydrate	21	20	26	26	25	27	22
	transport and							
	metabolism							
Н	Coenzyme	65	65	67	64	66	58	68
	transport and							
	metabolism							
1	Lipid transport	28	28	30	29	28	29	29
	and metabolism							
J	Translation	126	127	131	130	130	128	137
•	ribosomal							
	structure and							
	hiogenesis							
к	Transcription	18	17	21	20	21	10	21
	Replication	52	53	53	54	52	51	57
L	recombination	52	00	00	54	52	51	51
	and repair							
M	Coll	19	47	62	59	60	35	62
IVI	wall/mombrane/	40	47	02	50	00	35	02
	biogonogia							
N	Diogenesis	0	0	0	0	0	0	0
N		2	2	2	2	2	2	2
0		44	44	43	43	43	43	45
	al modification,							
	protein							
	turnover,							
	chaperones							

		A. capra	A. capra	A. ovis	A. centrale	A. marginale	A. platys	A. phagocytophilum	
	Functional	str. BIME1	str. BIME2	str. Haibei	str. Israel	str. Florida	str. S3	str. JM	
Category	category		Number of genes						
Р	Inorganic ion	31	31	36	35	34	35	34	
	transport and								
	metabolism								
Q	Secondary	12	12	12	12	12	11	12	
	metabolites								
	biosynthesis,								
	transport and								
_	catabolism								
R	General	0	0	0	0	0	0	0	
	function								
•	prediction only								
S	Function	90	91	89	91	92	83	88	
-	unknown	•	•	10	40	40	10	•	
I	Signal	9	9	10	10	10	10	9	
	transduction								
	mecnanisms	00	00	00	00	00	40	40	
U	Intracellular	38	38	38	38	38	40	40	
	tranicking,								
	secretion, and								
	transport								
V	Dofonso	4	1	1	1	٨	4	Λ	
v	mochanisms	4	4	4	4	4	4	4	
\ M /	Extracellular	0	0	0	0	0	0	0	
••	structures	0	0	0	0	0	0	U	
x	Mohilome	0	0	0	0	0	0	0	
~	nronhages	0	0	0	0	0	0	0	
	transposons								
Y	Nuclear	0	0	0	0	0	0	0	
	structure	Ŭ	Ũ	ů,	č	č	Ŭ	Ŭ	
Z	Cytoskeleton	0	0	0	0	0	0	0	

Appendix Table 8. The Anaplasma capra-positive numbers of goats and Hae. Longicornis in this study by PCR toward different

gene loci

	Goats from	Goats from	Hae. longicornis from	Hae. longicornis
Gene loci	Shandong	Guizhou	Shandong	from Guizhou
No. of tested	54	18	144	57
16S rRNA、gltA、groEL、msp4(+)	14	1	0	0
16S rRNA、gltA、msp4(+)	4	4	0	3
16S rRNA、 <i>gltA、groEL</i> (+)	1	0	0	0
gltA、groEL、msp4(+)	7	2	0	0
16S rRNA、 <i>gltA</i> (+)	1	0	0	0
gltA、msp4(+)	0	5	3	2
gltA(+)	2	2	4	4



Appendix Figure 1. The sampling sites where ticks and blood samples were collected in this study. Different color and marks represent the types of samples collected in different areas. The flags indicate the locations, where the goat blood samples were collected for next-generation sequencing of *Anaplasma capra* genomes.



Appendix Figure 2. The comparison between each *gltA* gene sequences of *Anaplasma capra* this study and sequence from human. The upper right part represents the number of bases that differ from each sequence and the lower left part represents the nucleotide identity (%) between each sequence from others.

	HL	A SDI	Goat 3	Goat A	Goat SD-5	Goat SD-b	Goat	Goat 9	Goat SD-1	DGoat SD-1	SD-1	SDI	SDI	SDI	SD-19	elGoat SD-2	olGoat SD-2	1Goat SD-2	LIGORI SD-2	blooat SD-2	SD-2	SD-2	SD-20	GOat 3	GOat	GL-11G	oat
KM206275 HLJ-14		0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974521 SD-1/Goat	100		0	0	0	5	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974522 SD-3/Goat	100	100		0	0	5	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974523 SD-4/Goat	100	100	100		0	5	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974524 SD-5/Goat	100	100	100	100		5	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974525 SD-6/Goat	100	99.6	99.6	99.6	99.6		5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	5	7	5	5	
OP974526 SD-7/Goat	100	100	100	100	100	99.6		0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974527 SD-9/Goat	100	100	100	100	100	99.6	100		0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974528 SD-12/Goat	100	100	100	100	100	99.6	100	100		0	0	0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974529 SD-13/Goat	100	100	100	100	100	99.6	100	100	100		0	0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974530 SD-14/Goat	100	100	100	100	100	99.6	100	100	100	100		0	0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974531 SD-15/Goat	100	100	100	100	100	99.6	100	100	100	100	100		0	0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974532 SD-16/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100		0	0	0	0	0	0	0	0	0	0	2	0	0	
OP974533 SD-17/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100		0	0	0	0	0	0	0	0	0	2	0	0	
OP974534 SD-18/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100		0	0	0	0	0	0	0	0	2	0	0	
OP974535 SD-20/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100	100		0	0	0	0	0	0	0	2	0	0	
OP974536 SD-21/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100	100	100		0	0	0	0	0	0	2	0	0	
OP974537 SD-22/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100	100	100	100		0	0	0	0	0	2	0	0	
OP974538 SD-23/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100	100	100	100	100		0	0	0	0	2	0	0	
OP974539 SD-25/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100	100	100	100	100	100		0	0	0	2	0	0	
OP974540 SD-26/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100	100	100	100	100	100	100		0	0	2	0	0	
OP974541 SD-27/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100		0	2	0	0	
OP974542 SD-29/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100		2	0	0	
OP974582 GZ-3/Goat	99.8	99.8	99.8	99.8	99.8	99.4	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8	99.8		2	2	
OP974583 GZ-5/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100	99.8	1.1	0	
OP974584 GZ-11/Goat	100	100	100	100	100	99.6	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100	100	99.8	100		

Appendix Figure 3. The comparison between each *groEL* gene sequences of *Anaplasma capra* this study and sequence from human. The upper right part represents the number of bases that differ from each sequence and the lower left part represents the nucleotide identity (%) between each sequence from others.



Appendix Figure 4. The comparison between each 16S rRNA gene sequences of *Anaplasma capra* this study and sequence from human. The upper right part represents the number of bases that differ from each sequence and the lower left part represents the nucleotide identity (%) between each sequence from others.



Appendix Figure 5. The comparison between each *msp4* gene sequences of *Anaplasma capra* this study and sequence from human. The upper right part represents the number of bases that differ from each sequence and the lower left part represents the nucleotide identity (%) between each sequence from others.