Expression analysis and biological characterization of *Babesia* sp. BQ1 (Lintan) (*Babesia motasi*-like) rhoptry-associated protein 1 and its potential use in serodiagnosis via ELISA

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**Abstract**

**Background:** In China, ovine babesiosis is one of the most important tick-borne haemoparasitic diseases of small ruminants. It has a significant economic impact, and several *Babesia motasi*-like isolates have been recently shown to be responsible for ovine babesiosis in this country.

**Methods:** Full-length and C-terminal-truncated forms of the *rap-1a61-1* gene of *Babesia* sp. BQ1 (Lintan) were cloned into the pET-30a plasmid and subsequently expressed as His-fusion proteins. The resulting recombinant RAP-1a proteins (rRAP-1a61-1 and rRAP-1a61-1/CT) were purified and evaluated as diagnostic antigens using Western blot analysis and ELISA. The native *Babesia* sp. BQ1 (Lintan) RAP-1 protein was recognized using Western blots and IFAT by antibodies that were raised in rabbits against rRAP-1a61-1/CT. The specificity, sensitivity and positive threshold values for rRAP-1a61-1/CT in ELISA were evaluated.

**Results:** Cross-reactivity was observed between rRAP-1a61-1/CT and positive sera for *Babesia* sp. BQ1 (Lintan), *Babesia* sp. BQ1 (Ningxian) and *Babesia* sp. Tianzhu isolates obtained from infected sheep. At one week post-inoculation, a significant increase was observed in the amount of antibodies produced against RAP-1a, and high levels of antibodies against RAP-1a were observed for 3 months (at 84 days p.i.). A total of 3198 serum samples were collected from small ruminants in 54 different regions in 23 provinces of China. These samples were tested using ELISA based on the rRAP-1a61-1/CT protein. The results indicated that the average positive rate was 36.02%.

**Conclusions:** The present study suggests that rRAP-1a61-1/CT might be a potential diagnostic antigen for detecting several isolates of *B. motasi*-like parasites infection.

**Keywords:** *Babesia* sp. BQ1 (Lintan), Rhoptry-associated protein, Biological characterization, Diagnosis

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Background

Ovine babesiosis is one of the most important tick-borne diseases of small ruminants. This disease has been described in many countries throughout Asia, Europe and Africa [1]. A phylogenetic tree based on the sequence of the 18S rRNA gene showed that Babesia species belonging to the following four groups are responsible for babesiosis in small ruminants: B. ovis, B. motasi which are a European and a Chinese clade (B. motasi-like), and the Babesia sp. Xinjiang and B. crassa [2–7]. In the B. motasi Chinese clade (B. motasi-like), six Babesia isolates (Babesia sp. BQ1 (Lintan), Babesia sp. BQ1 (Ningxian), Babesia sp. Tianzhu, Babesia sp. Hebei, Babesia sp. Madang and Babesia sp. Liaoning) have been recently reported in the Gansu, Hebei and Liaoning provinces of China. These isolates can be classified into one or more sub-groups according to their phylogenetic relationships, which have been reconstructed based on several (18S rRNA, ITS, 28S, rap-1) phylogenetic markers [7–12].

Currently, the strategy for controlling babesiosis is based on chemotherapy against either the vector or the parasite. Several compounds are used, but mainly ivermectin is used against ticks and imidocarb against Babesia. Several new anti-Babesia drugs (Triclosan, Nerolidol, Artesunate and Atovaquone) are also used to control and treat babesiosis [13]. However, this strategy has many drawbacks, such as drug-resistance, the presence of drug metabolites in the milk and meat, and high toxicity to the animals themselves. Optimal treatment methods, including those with biological activity against ticks and vaccines that target either the ticks or the parasites, have been extensively studied. But only live attenuated vaccines are currently used in some countries, and no producing or testing recombinant or subunit vaccines have been available to prevent babesiosis so far [13–16].

Clinical symptoms of babesiosis usually include fever, anemia, and hemoglobinuria and can lead to death in severe cases, which become apparent when merozoites from the Babesia invade and replicate within host erythrocytes, resulting in detectable parasitemia [17]. During the process of erythrocytic invasion, some of the proteins, including apical membrane antigen-1 (AMA-1), rhoptry-associated-protein-1 (RAP-1), and spherical body proteins (SBP-1, 2, 3), are secreted by the apical organelles (rhoptries, micronemes and dense granules). Although the functions and implications of these proteins for vaccines designed against Babesia infections have been described mainly in B. bigemina and B. bovis, these data may form a basis for developing a recombinant vaccine against other Babesia infections [17].

RAP-1 is an immunogenic protein that is approximately 40–60 kDa in size. It is localized in rhoptries and has been characterized in all Babesia species examined so far (B. bovis, B. bigemina, B. divergens, B. canis, B. ovis, B. motasi-like, B. orientalis and B. gibsoni) [12, 18–23]. RAP-1 was detected at all asexual growth stages [24] and was expressed during the sporozoite stage by a RAP-1 specific antisera that was obtained from rabbits immunized to neutralize the binding of sporozoites of B. bovis to erythrocytes [25]. Moreover, the role of RAP-1 in the erythrocyte invasion process has been studied in vitro during parasite adhesion to erythrocytes, and experiments have been performed to study the inhibitory effect of RAP-1-specific antibodies on growth and invasion [24–26].

Immunization with native and recombinant RAP-1 protein from B. bovis and B. bigemina also induced immunity and protected animals upon challenge [27–29]. These studies strongly support the notion that RAP-1 plays a functional role in the biology and development of Babesia parasites and represents a potential candidate for the development of recombinant vaccines against babesiosis [30–33].

RAP-1 was characterized and found to be present as multiple gene copies that are arranged in a tandem, head to tail manner in all Babesia species. In sheep Babesia species, the rap-1 locus of the B. motasi-like gene is similar to the locus of B. bigemina: in both species, three different isoforms (rap-1a, rap-1b and rap-1c), 5 identical rap-1b copies and a single rap-1c located at end of locus have been described [12, 34]. Two rap-1a classes named rap-1a61 and rap-1a67 have been described, and each class is represented by two close variants (rap-1a61-1 and rap-1a61-2; rap-1a67-1 and rap-1a67-2) in Babesia sp. BQ1 (Lintan) [34]. Polymorphisms were limited to three nucleotide positions in the rap-1a61 copies (rap-1a61-1, -2, -3) and were found in the rap-1 locus of Babesia sp. BQ1 (Lintan) (rap-1a61-1 and -2), Babesia sp. BQ1 (Ningxian) and Babesia sp. Tianzhu (rap-1a61-2 and -3) [12, 34]. According to a phylogenetic tree that was constructed based on the rap-1 gene sequences, the B. motasi-like group can be separated into two subgroups: one that consists Babesia sp. BQ1 (Lintan and Ningxian) and Babesia sp. Tianzhu and another that contains Babesia sp. Hebei [12].

The rap-1a gene has been found in all Babesia species and shows highly conserved protein features (30–45 % identity among species), suggesting that this gene plays an important role in the RBC invasion process [35]. Moreover, RAP-1a, in Babesia has been widely used as a vaccine candidate and diagnostic antigen [21, 29, 36, 37]. The expression, biological characterization and immunogenicity of RAP-1a in sheep Babesia species have not been extensively studied. In the present study, we cloned and expressed RAP-1a and then detected the presence of antibodies directed against native and recombinant forms of RAP-1a during the course of infection. To evaluate the potential of using recombinant RAP-1a as a diagnostic antigen, we compared the diagnostic potential
of full-length and truncated C-terminal recombinant \textit{Babesia} sp. BQ1 (Lintan) RAP-1a proteins using via Western blot analysis. We discuss our results in combination with the outcomes of previous studies.

\textbf{Methods}

\textbf{Parasites}

The original isolates of the \textit{B. motasi}-like group, including \textit{Babesia} sp. BQ1 (Lintan), \textit{Babesia} sp. BQ1 (Ningxian), \textit{Babesia} sp. Tianzhu and \textit{Babesia} sp. Hebei, were obtained from infected sheep and then cryopreserved in liquid nitrogen at the Vector and Vector-borne Disease (VVBD) Laboratory of the Lanzhou Veterinary Research Institute (LVRI) (CAAS Lanzhou, China) [3, 4, 6].

\textbf{Sera}

Positive sera were obtained from five different breeds of sheep (two “Tan mutton” Chinese sheep, including No. 3216 and 2007, and three “Vendean” French sheep, including No. 3309, 3533 and 3446) that were infected with \textit{Babesia} sp. BQ1 (Lintan). Immune-positive sera were obtained from sheep Nos. 08026, 08040 and 08020, which were infected by \textit{Babesia} sp. BQ1 (Ningxian), \textit{Babesia} sp. Tianzhu and \textit{Babesia} sp. Hebei, respectively, and immune-positive sera were obtained from sheep No. 3201, which was infected with \textit{Babesia} sp. Xinjiang, as previously described [38]. These serum samples were used to evaluate cross-infection and antibody kinetics. Positive sera from sheep that were experimentally infected with \textit{Theileria luwenshuni}, \textit{T. uilenbergi} and \textit{Anaplasma ovis} or bovine \textit{B. bovis} and \textit{B. bigemina} were used as the controls to evaluate cross-reactivity.

Fifty serum samples were collected from five sheep (3–12 wpi) that were experimentally infected with \textit{Babesia} sp. BQ1 (Lintan). In total, 141 sera and anticoagulated blood samples were collected from sheep in the Xinjiang, Henan and Gansu provinces of China. Genomic DNA was purified from blood samples and tested using specific PCR primers, and 141 genomic DNA samples were found to be specifically positive for \textit{Babesia} sp. BQ1 (Lintan) in the sequencing analysis (data not shown). The sera samples \((n = 141)\) that corresponded to these genomic DNA samples were selected as the positive serum samples. A total of 191 sera were used as positive samples to evaluate the sensitivity of the ELISA, and a mixture of these sera was used as the positive control serum.

Sera were collected from 492 lambs that were purchased from a \textit{Babesia}-free region in Jingtai country (Gansu province of China) from 2010 to 2015. After nested PCR tests were performed, none of the samples tested positive, and a mixture of these serum samples was used as the negative control serum to evaluate the specificity of the ELISA.

A total of 3198 serum samples were randomly collected from clinically healthy sheep from 54 different locations in 23 Chinese provinces between 2010 and 2015 (Fig. 1). All blood samples were collected in non-anticoagulation tubes and transported to the laboratory while on ice. The serum was then separated and stored at -20 °C until further use.

\textbf{Ethical approval}

This study was approved by the Animal Ethics Committee of the Lanzhou Veterinary Research Institute at the Chinese Academy of Agricultural Sciences. All animals used in this study were handled in accordance with the Animal Ethics Procedures and Guidelines of the People’s Republic of China.

\textbf{Cloning of the full-length and the C-terminal truncated RAP-1a61-1 genes}

Genomic DNA was extracted from infected blood samples obtained from sheep infected with \textit{Babesia} sp. BQ1 (Lintan) using a QiAamp DNA Blood Kit (QIAGEN, Maryland, USA) according to the manufacturer’s instructions. These DNA samples were used as the templates for PCR. The oligonucleotide primers that were designed based on the DNA sequence of \textit{Babesia} sp. BQ1 (Lintan) RAP-1a61-1 (GenBank accession number KC953701) and modified to include either \textit{Xba} I and \textit{BamH} I or \textit{Nde} I and \textit{Hind} III (New England BioLabs) enzyme restriction sites (underlined below) at their 5’ end. The following primers were used: RAP-1a61-full-F: 5’-CGC TCT AGA -GT ACG CCA TTA CCA CCG TTC-3’, RAP-1a61-full-R: 5’-GCC GGA TCC GAG CCA CGA ATC AGC TTC-3’, RAP-1a61/CT-F: 5’-CGC CAT ATG ACC ACG GTG ATG CCC GAG-3’, and RAP-1a61/CT-R: 5’-GCC AAG CTT GAG CCA CGA ATC TTC GG AAT GGA TCC-3’. The PCR products were cloned into the pUC57 vector according to the manufacturer’s instructions (Genscript). The inserts were sequenced using vector primers. Representative plasmids were selected and digested using either \textit{Xba} I and \textit{BamH} I or \textit{Nde} I and \textit{Hind} III restriction enzymes and subsequently cloned into digested pET-30a expression vectors (Genscript) according to the manufacturer’s instructions. This resulted in the generation of the recombinant plasmids pET-30a-RAP-1a61-1 and pET-30a-RAP-1a61-1/CT, which contained the full-length gene (1359 bp, except for signal peptide sequence) and the C-terminal-truncated fragment (450 bp), respectively. Both of the cloned RAP-1a proteins were tagged with histidine.

\textbf{Expression and purification of the recombinant proteins}

To express the recombinant proteins, both pET-30a-RAP-1a61-1 and pET-30a-RAP-1a61-1/CT were transformed
into BL21 E. coli (DE3 strain) cells. The expression of the His-tagged fusion proteins rRAP-1a61-1 and rRAP-1a61-1/CT was induced in the presence of kanamycin (50 μg/ml) by adding 1 mM IPTG and then culturing the cells at 15 °C for 16 h. The bacterial cultures were harvested and lysed using ultrasonication in binding buffer (50 mM Tris, 6 M Gu-HCl; pH 8.0) containing phenylmethylsulfonyl fluoride (PMSF) and then purified as inclusion bodies from the E. coli cells. The target protein was then eluted using urea followed by a series of buffers containing increasing concentrations of imidazole (200 mM, 500 mM and 2500 mM). After the proteins were renatured, they were suspended in 1× SDS-PAGE sample loading buffer (62.5 mM Tris-HCl [pH 6.8], 2 % SDS, 5 % β-mercaptoethanol, 10 % glycerol, and 0.02 % bromophenol blue) and boiled for 5 min. The samples were briefly centrifuged prior to being separated using SDS-PAGE. Recombinant protein expression was then analyzed using Western blot analysis.

**Preparation of anti-rRAP-1/CT specific rabbit immune serum**

New Zealand Rabbits (2 kg each) were subcutaneously injected with 400 μg of purified rRAP-1a61-1/CT protein that had been emulsified in Freund’s complete adjuvant (FIA, Sigma). These injections were administered on days 15, 20 and 28. The sera were collected from the immunized rabbits 15 days after the last injection, purified and stored at -20 °C until further use.

**Western blot analysis**

Approximately 20 μg of purified Babesia sp. BQ1 (Lintan) recombinant proteins (rRAP-1a61-1 and rRAP-1a61-1/CT) were separated using SDS-PAGE in 18 % polyacrylamide gels. The proteins were transferred to nitrocellulose (NC) membranes with a 0.45 μm pore size (BioRad) at 24 V and 50 W for 35 min. The membrane was then cut into 0.25 cm strips and incubated in a blocking solution [5 % skimmed milk powder in Tris-buffered saline (pH 7.6) with 0.1 % Tween-20 (TBST)] for approximately 3 h at 4 °C on a shaker. After the NC strips had been washed three times, they were incubated for 1 h with each tested serum (diluted at 1:100 in TBST). After the strips were washed three times in TBST, they were incubated for 2 h with monoclonal anti-goat/sheep IgG-alkaline phosphatase conjugates (Sigma, A8062, dilution: 1:5000). The strips were then washed three times with TBST and incubated in a 5-bromo-4-chloro-3-indolyl phosphate/nitro blue tetrazolium (BCIP/NBT) liquid substrate system (B1911-100ML, Sigma) for 15 to 20 min. The approximate molecular weights of the revealed proteins

*Fig. 1* The geographical distribution of the B. motasi-like sp. that was identified in this study and the distribution of *Haemaphysalis* spp. in China. The positive rates exceed 50 % indicated with deep red, and the positive rates ranged from 20 to 40 % indicated with light red, less than 10 % infected rates indicated without color.
were evaluated by comparing their migration to a standard molecular weight marker (SM0671, 10–170 kDa, PageRuler Prestained Protein Ladder; SM1861, 1.7–40 kDa, Spectra Multicolor Low Range Protein Ladder, Thermo Scientific).

**IFAT and confocal laser microscopic observation**

An immunofluorescence analysis was performed using a procedure previously described by Moitra et al. [39] with rabbit antiserum directed against rRAP-1a61-1/CT. The iRBC membrane was stained using a PKH26 red fluorescent cell membrane labeling kit (MINI26, Sigma-Aldrich) according to the manufacturer’s instructions. The iRBCs were fixed in a buffer containing 4 % paraformaldehyde and 10 mM piperazine-N, N-bis (2-ethanesulfonic acid) in PBS (PIPES) at pH 6.4 on 22 mm² poly-L-lysine-coated coverslips for 30 min at room temperature. The coverslips were washed in PBS and then blocked for 1 h in 3 % bovine serum albumin (BSA) in PBS containing 0.25 % Triton X-100. The coverslips were stained overnight at 4 °C with either pre-immune serum or purified anti-rRAP-1a61-1/CT serum (1:50) diluted in blocking buffer without Triton X-100. The coverslips were washed 3 times in PBS, stained for 1 h with FITC-conjugated anti-rabbit IgG (whole molecule) secondary antibodies (Sigma: F0382; 1:80), and then washed 3 times in PBS. The coverslips were stained with 1 μg/ml 4,6-diamidino-2-phenylindole (DAPI) (Sigma: D8417) for 10 min and then examined for reactivity using confocal microscopy.

**ELISA**

The optimum concentration of coating buffer (0.1 M carbonate/bicarbonate, pH 9.6) was determined for rRAP-1a61-1/CT (100 μl at 2 μg/ml). It was then distributed and adsorbed in 96-well flat-bottom microplates at 4 °C overnight. After three washes with PBST (0.1 % Tween-20), the plates were blocked with 200 μl of 2 % gelatin in PBST at 37 °C for 1 h. After the plates were washed, the positive and negative sera (dilution: 1/100) were distributed in duplicate, and the plates were incubated for 1 h at 37 °C. After the plates were washed 3 times, 100 μl of monoclonal antibody (IgG, sheep peroxidase) (Sigma, A9452, dilution: 1:1000) was added to each well, and the wells were incubated for 1 h at 37 °C. After the plates were washed another 3 times, 100 μl of 1-Step™ Ultra TMB-ELISA (34028–250 ml, Thermo Scientific) was added to each well, and the plates were incubated for 15 to 25 min at room temperature. The reaction was stopped by adding 100 μl of 2 M H2SO4, and the plates were read at 450 nm using an ELISA automat (Bio-Rad Model 680 microplate reader, USA).

**Results**

**Cloning and expression of RAP-1a61-1 and C-terminal-truncated RAP-1a61-1**

The genes encoding RAP-1a61-1 (aa 22–474) and C-terminal RAP-1a61-1/CT (aa 325–474) (aa sequence corresponding to GenBank accession number AGV15809) were successfully amplified using PCR and inserted into the cloning vector pUC57. The pET-30a-RAP-1a61-1 and pET-30a-RAP-1a61-1/CT recombinant plasmids were identified using enzyme restriction analysis (Sac I/Sal I and Mlu I/Hind III, respectively) and confirmed by sequencing using specific primers. The insert sequences were also verified (data not shown). After the expression of each recombinant protein was induced using IPTG, the E. coli lysates were analyzed using SDS-PAGE and Western blot analysis of an anti-His serum. The results of SDS-PAGE showed that rRAP-1a61-1 and rRAP-1a61-1/CT were expressed as proteins that were approximately 52 kDa and 20 kDa in size, respectively (Fig. 2).

**Evaluation of the specificity of rRAP-1a61-1 and rRAP-1a61-1/CT using Western blot analysis**

The reactivity of each positive serum that was obtained from infected sheep with rRAP-1a61-1 and rRAP-1a61-1/CT was analyzed using Western blot analysis. Strong reactions showing an expected size of rRAP-1a61-1/CT (approximately 20 kDa) were observed in the positive sera obtained from sheep infected with Babesia sp. BQ1 (Lintan) (No. 3216), Babesia sp. BQ1 (Ningxian) (No. 08026) and Babesia sp. Tianzhu (No. 08040). In contrast, no reaction was observed between rRAP-1a61-1/CT and sera obtained from sheep infected with Babesia sp. Hebei (No. 08020) and Babesia sp. Xinjiang (No. 3201) (Fig. 3). Furthermore, cross-reactivity between the full-length rRAP-1a61-1 protein and sera from Babesia sp. Xinjiang-infected sheep and B. bigemina-infected cattle was also observed (data not shown).

**Identification of native RAP-1 proteins using Western blot analysis and IFAT**

To identify native RAP-1 in Babesia sp. BQ1 (Lintan) merozoites, antibodies against the C-terminal-truncated recombinant RAP-1a61-1/CT were obtained from rabbits and the positive serum obtained from animals infected with Babesia sp. BQ1 (Lintan) (No. 3216), and the RAP-1a61-1 protein was detected at approximately 52 kDa in the lysates of Babesia sp. BQ1 (Lintan)-infected sheep erythrocytes (Fig. 4). No specific reaction was observed in Western blot analyses between the sera with the lysates of uninfected sheep erythrocytes (data not shown).

The IFAT results showed that the merozoite nucleus of parasites were stained with DAPI and FITC-conjugated anti-rabbit IgG is able to detect anti-rRAP-1a61-1/CT.
antiserum. The immunofluorescence signals are the result of the interaction between the serum from a rabbit that was immunized by rRAP-1a61-1/CT and *Babesia* sp. BQ1 (Lintan) merozoites. Specific fluorescent signaling was not detected using pre-immune rabbit serum (Fig. 5).

Cross-reactivity with other hemoparasites using ELISA

Positive sera against *Babesia* sp. BQ1 (Lintan), *Babesia* sp. BQ1 (Ningxian), *Babesia* sp. Tianzhu, *Babesia* sp. Hebei, *Babesia* sp. Xinjiang, *T. lunwenshuni*, *T. ulilenbergi*, *A. ovis*, *B. bovis*, and *B. bigemina* were tested using ELISA to evaluate cross-reactivity with rRAP-1a61-1/CT. The tests were repeated in triplicate, and the specific antibody mean rate ($\text{AbR \%} = (\text{Sample mean OD-Negative control mean OD})/(\text{Positive control mean OD-Negative control mean OD}) \times 100$) and standard deviation were calculated using Excel 2007 [38]. The results indicated that cross-reactivity was observed only between three *B. motasi*-like isolates (Lintan, Tianzhu and Ningxian) and that no cross-reactivity was observed with the sera against *Babesia* sp. Hebei, *T. lunwenshuni*, *T. ulilenbergi*, *B. bovis*, *B. bigemina* or *A. ovis* (Fig. 6).

The kinetics of anti-*Babesia* sp. BQ1 (Lintan) antibodies were tested in experimentally infected sheep using ELISA

The kinetics of antibodies produced against RAP-1a have been studied using ELISA with rRAP-1a61-1/CT in sera collected from 5 intact infected sheep (Nos. 3216, 2007, 3390, 3533 and 3446) before (0 weeks post-infection, WPI0) and after infection (from WPI1 to WPI12). A significant increase in the amount of antibodies produced against RAP-1a was observed after the sheep were infected, but varying kinetics were observed between individual sheep. In two of the Chinese sheep (Nos. 3216 and 2007), antibody production against RAP-1a was increased during the early phase of infection, at 1–3 weeks (WPI1-3), after which it decreased. In contrast, a gradual increase in the number of antibodies produced against RAP-1a was observed in the three French sheep (Nos. 3390, 3533 and 3446). Higher antibody levels were observed at 6–7 weeks post-infection (WPI6-7) in sheep Nos. 3390 and 3446 and at 9 weeks (WPI9) in sheep No. 3533. In general, antibodies were produced at 1 week post-inoculation, continued to increase through 6 weeks (WPI6), and remained stable.

Cross-reactivity with other hemoparasites using ELISA

Positive sera against *Babesia* sp. BQ1 (Lintan), *Babesia* sp. BQ1 (Ningxian), *Babesia* sp. Tianzhu, *Babesia* sp. Hebei, *Babesia* sp. Xinjiang, *T. lunwenshuni*, *T. ulilenbergi*, *A. ovis*, *B. bovis*, and *B. bigemina* were tested using ELISA to evaluate cross-reactivity with rRAP-1a61-1/CT. The tests were repeated in triplicate, and the specific antibody mean rate ($\text{AbR \%} = (\text{Sample mean OD-Negative control mean OD})/(\text{Positive control mean OD-Negative control mean OD}) \times 100$) and standard deviation were calculated using Excel 2007 [38]. The results indicated that cross-reactivity was observed only between three *B. motasi*-like isolates (Lintan, Tianzhu and Ningxian) and that no cross-reactivity was observed with the sera against *Babesia* sp. Hebei, *T. lunwenshuni*, *T. ulilenbergi*, *B. bovis*, *B. bigemina* or *A. ovis* (Fig. 6).

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until 9 weeks (WPI9), after which slight decreases were observed through 12 weeks (WPI9-12). High levels of antibodies against RAP-1a usually lasted 3 months (84 days p.i.), despite the decrease in production that was observed after 6 weeks (42 days p.i.) (Fig. 7).

Evaluation of specificity, sensitivity and positive threshold values in ELISA to detect rRAP-1/CT
MedCalc statistical software was used to evaluate the sensitivity, specificity, and positive threshold value in ELISAs performed to test 191 positive sera and 492 negative sera, as previously described [40]. A percentage corresponding to the specific antibody mean rate (AbR %) was calculated for each serum sample. The results indicated that the threshold value was 18.84 %, which corresponded to 94.8 % sensitivity (95 % CI = 82.4–98.4) and 96.1 % specificity (95 % CI = 85.1–99.4). The numbers of false positive and false negative sera were 10 and 15, respectively (Fig. 8).

ELISA identification of the rRAP-1a61-1/CT protein as a potential antigen for use in serological epidemiology in B. motasi-like sub-group (Lintan, Ningxian and Tianzhu isolates) infections
Field serum samples were extracted from a total of 3198 sheep and goats from 54 different regions of 23 Chinese provinces. The samples were analyzed using ELISA to evaluate the prevalence of infection with B. motasi-like
(Lintan, Ningxian and Tianzhu) isolates in these regions. The results of ELISA detection to identify positive samples are summarized in Table 1. Sero-positive with *B. motasi*-like were identified in 51 regions covering all 23 surveyed provinces. The total positive rate in these regions was 36.02 % (95 % CI = 33.7 – 38.4). The sero-prevalence ranged from 2.22 to 81.11 %. The significantly high positive rates exceeded 50 % in 7 of the 23 surveyed provinces, including Liaoning (50 %), Inner Mongolia (61.01 %), Shaanxi (75.68 %), Hunan (52.73 %), Sichuan (59.72 %), Anhui (81.11 %) and Shanxi (58 %). Three additional provinces, including Qinghai (45.35 %), Ningxia (49.38 %) and Guangdong (41.89 %), had positive rates exceeding 40 %. Lower sero-prevalence was observed in Tibet (2.22 %), Jilin (7.55 %) and Guangxi provinces (9.74 %). Positive rates that were greater than 80 % were observed in Chifeng (86.15 %) in Inner Mongolia and Lintan (83.33 %) in the Gansu provinces (Table 1, Fig. 1).

**Discussion**

In the genus *Babesia*, RAP-1 was initially described according to its recognition by monoclonal antibodies to merozoite antigens in *B. bigemina*. It was designated p58 at that time [41], and it was subsequently identified in all *Babesia* species that were examined. The sequence of the full-length *rap-1* gene was first published in *B. bovis* (named pBv60) [18], and the sequence and multigene nature of p58 (later named *rap-1a*) were described in *B.
The name RAP-1 was first applied when *B. bovis*-like isolates and *B. bovis* and homologue, *Babesia* locus was initially defined in *Babesia bigemina*. *B. motasi* were initially described in *et al. Parasites & Vectors* B. bigemina B. motasi B. sp. Xinjiang, another genes obtained from rap-1 isolates at both the nu-
rap-1 Babesia Babesia Babesia sp. BQ1 (Lintan) B. motasi B. sp. Tianzhu and at the B-cell epitopes pri-
rap-1 Babesia Babesia Babesia B. motasi Babesia sp. BQ1 (Ningxian) in both Western blot ana-
sequences were almost – sp. BQ1 (Ningxian) and *B. bigemina* because of the conserved features in the 
carrier between RAP-1a and RAP-1a67 are more conserved at the C-terminal region than the N-terminal region in *B. motasi*-like [34], which is not consistent with the common features described above. Moreover, the sequences of RAP-1a and the full rap-1 locus, including the intergenic regions in three isolates (Lintan, Ningxian and Tianzhu) of *B. motasi*-like, were almost identical [12]. The features of the more conserved C-
terminal region of the RAP-1a protein in *B. motasi*-like could indicate that this form is an ideal candidate for diagnostics aimed at identifying several *B. motasi*-like isolates.

The sera were able to recognize the native RAP-1 protein in Western blot analysis and IFAT, suggesting that the sheep had been immunized against RAP-1 during infection. Significant cross-reactivity was observed between rRAP-1a61-1/CT of *Babesia* sp. BQ1 (Lintan) and the positive sera obtained from *Babesia* sp. Tianzhu and *Babesia* sp. BQ1 (Ningxian) in both Western blot analyses and ELISA, but not in the Hebei isolate. This is not surprising because the rap-1a sequences were almost identical in these three *Babesia* isolates at both the nucleotide and amino acid level. In contrast, the sequences of all of the rap-1 genes obtained from *Babesia* sp. Hebei were clearly different from the other 3 isolates, suggesting that this isolate might belong to a different species [12]. These findings were inconsistent with a previous study that indicated that weak cross-reactivity between *Babesia* sp. BQ1 (Ningxian) and *Babesia* sp. BQ1 (Lintan) was detected using Western blot analysis but not ELISA using a merozoite antigen (BQMA) [40]. Some novel antigens (e.g., BQP35 and BOHsp90) for *Babesia* sp. BQ1 (Lintan) were recently identified as candidate diagnostic antigens. Western blot analysis and ELISA tests that are based on BQP35 have shown that there is cross-reactivity between *Babesia* sp. BQ1 (Lintan) and *Babesia* sp. Tianzhu, which is in agreement with the results obtained when merozoite extracts were used [51]. However, rBQHSP90 could not differentiate between any *B. motasi*-like isolates and *Babesia* sp. Xinjiang, another distinct ovine *Babesia* species [52]. Differences in cross-reactivity have been observed when using the ELISA method based on BQP35, BQHsp90 or RAP-1a of *Babesia* sp. BQ1 (Lintan), and these might be caused by differences in molecular characterizations and the detection limitations and sensitivities of these antigens. These findings suggest that rRAP-1a61-1/CT could be used as a potentially diagnostic antigen to increase the sensitivity and specific-
| Province | Prefecture | No. of sera | No. of positive sera | Positive rate (%) in different regions | Positive rate (%) in each province |
|----------|------------|-------------|----------------------|-----------------------------------------|-----------------------------------|
| Qinghai  | Haibei     | 269         | 122                  | (122/269) 45.35                         |
| Xinjiang | Yili       | 229         | 82                   | 35.81                                   |
|          | Akesu      | 90          | 20                   | 22.22                                   |
|          | Habhahe    | 100         | 16                   | 16                                      |
| Gansu    | Jingtai    | 120         | 8                    | (160/447) 35.79                         |
|          | Jiayuguan  | 81          | 6                    | 7.41                                    |
|          | Tianzhu    | 81          | 56                   | 69.14                                   |
|          | Yongchang  | 75          | 15                   | 20                                      |
|          | Lintan     | 90          | 75                   | 83.33                                   |
| Tibet    | Lhasa      | 90          | 2                    | (2/90) 2.22                             |
| Liaoning | Huanren    | 29          | 14                   | 48.28                                   |
|          | Anshan     | 27          | 10                   | 37.04                                   |
|          | Fengcheng  | 28          | 18                   | 64.29                                   |
| Jilin    | Yongji     | 26          | 3                    | 11.54                                   |
|          | Jiutai     | 27          | 1                    | 3.70                                    |
| Inner Mongolia | Ordos | 63       | 21                   | (133/218) 61.01                        |
|          | Manzhouli  | 13          | 0                    | 0                                       |
|          | Baotou     | 12          | 0                    | 0                                       |
|          | Chifeng    | 130         | 112                  | 86.15                                   |
| Shaanxi  | Yulin      | 74          | 56                   | (56/74) 75.68                           |
| Ningxia  | Wuzhong    | 81          | 40                   | (40/81) 49.38                           |
| Guangxi  | Jingxi     | 35          | 3                    | 8.57                                    |
|          | Tianyang   | 56          | 10                   | 17.85                                   |
|          | Pingxiang  | 18          | 0                    | 0                                       |
|          | Guilin     | 45          | 2                    | 4.44                                    |
| Hunan    | Xinhuang   | 29          | 16                   | (29/55) 52.73                           |
|          | Xiangtan   | 26          | 13                   | 50                                      |
| Sichuan  | Panzhihua  | 36          | 25                   | 69.44                                   |
|          | Hejiang    | 36          | 18                   | 50                                      |
| Yunnan   | Fuyuan     | 35          | 2                    | 5.71                                    |
|          | Honghe     | 32          | 16                   | 50                                      |
|          | Jinghong   | 20          | 5                    | 25                                      |
|          | Yanshan    | 34          | 10                   | 29.41                                   |
|          | Menghai    | 12          | 3                    | 25                                      |
|          | Ruili      | 32          | 14                   | 43.75                                   |
|          | Kunming    | 37          | 15                   | 40.54                                   |
| Anhui    | Hefei      | 90          | 73                   | (73/90) 81.11                           |
| Guangdong | Zhaoqing  | 38          | 20                   | 52.63                                   |
|          | Lianzhou   | 36          | 11                   | 30.56                                   |
| Guizhou  | Dushan     | 30          | 1                    | 3.33                                    |
|          | Ziyun      | 32          | 13                   | 40.63                                   |
|          | Yuping     | 32          | 19                   | 59.38                                   |
|          | Guiyang   | 72          | 13                   | 18.06                                   |
The kinetics analysis revealed that the humoral immune response to this parasite varied in different sheep breeds (Chinese and French sheep) during the course of infection. This result was inconsistent with a previous study that used BQMA and indicated that there is no difference in kinetics between these two sheep breeds [38]. Antibody production against RAP-1 was initially increased in the two infected Chinese sheep between WPI1 and WPI2, but it then decreased. These results were consistent with the findings of a recent study, in which a RAP-1a peptide designed from the RAP-1a N-terminal of Babesia sp. BQ1 (Lintan) was used as an antigen to perform ELISA. Antibody production against this peptide displayed kinetics that were similar to those observed against rRAP-1a61-1 (data not shown). In contrast, the production of specific antibodies against Babesia sp. BQ1 (Lintan) in the three infected French sheep was increased at three weeks post-infection and peaked between six and nine weeks post-infection. On average, RAP-1a was expressed at sequentially higher levels from week 1 to week 7 post-infection before its levels subsequently decreased. The difference in the kinetics of antibody production between these five infected sheep might result from differences in sheep breeds or individual variation. In general, our results are similar to those described in a previous study that used a merozoite antigen (BQMA) to evaluate the kinetics of antibody production during the course of Babesia sp. BQ1 (Lintan) infection. That study showed that antibody production was decreased from WPI 12 (approximately 80 DPI) [38].

A serological diagnostic test (ELISA) is the most practical and economical assay for a wide range of epidemiological investigative purposes, and it has been established as a method for evaluating the lysates of Babesia-infected erythrocytes in a number of Babesia species [55]. However, studies focusing on the development of a serological diagnostic ELISA test that is based on recombinant immune antigens that can detect ovine babesiosis are rare. The development of ELISA tests requires preliminary knowledge of parasite polymorphism, the ability to successfully culture cells in vitro to produce antigens, and/or prior genetic knowledge of the targeted antigen so that specific sequences can be selected as an epitope. Some improved ELISA methods for sero-epidemiological surveys have been developed based on specific recombinant proteins (RAP-1 and SBPs) that are involved in the RBC invasion process in B. bovis, B. bigemina and B. caballi species [36, 37, 50, 56]. In this study, the results of sero-investigations using an ELISA method that was developed based on the rRAP-1a61-1/CT protein indicated that Babesia sp. BQ1 (Lintan), Babesia sp. BQ1 (Ningxian) and Babesia sp. Tianzhu infections were identified in all of the investigated provinces. Recent large-scale sero-epidemiological ELISA surveys of Babesia sp. BQ1 (Lintan) were performed in small ruminants from 22 provinces of China, and these were also regions that were tested in our study, and the results indicated that infection by this isolate was identified in all 22 investigated provinces, with an average prevalence of 43.5 %. These values are slightly higher than the values produced in our study and they showed a positive rate that was as high as 91 % in Inner Mongolia [57]. The positive rates from 8 provinces that were tested using rRAP-1a were slightly higher, whereas the positive rates from 14 of the provinces were significantly lower in our study than in a previous study [57]. This variation may result from differences in the limitations of the tested antigens. Moreover, another

| Province | Prefecture   | No. of sera | No. of positive sera | Positive rate (%) in different regions | Positive rate (%) in each province |
|----------|--------------|-------------|----------------------|----------------------------------------|----------------------------------|
| Chongqing| Jiangjin     | 30          | 11                   | 36.67                                  | (15/53) 28.30                    |
|          | Wanzhou      | 23          | 4                    | 17.39                                  |                                  |
| Shandong | Dongying     | 90          | 19                   | (19/90) 21.11                          |                                  |
| Shanxi   | Lvliang      | 50          | 29                   | (29/50) 58                             |                                  |
| Zhejiang | Jingning     | 36          | 15                   | 41.67                                  | (32/117) 27.35                   |
|          | Qintian      | 29          | 4                    | 13.79                                  |                                  |
|          | Hangzhou     | 52          | 13                   | 25                                     |                                  |
| Hubei    | Suizhou      | 81          | 27                   | (27/81) 33.33                          |                                  |
| Hebei    | Baoding      | 169         | 25                   | (25/169) 14.79                         |                                  |
| Henan    | Linzhou      | 60          | 20                   | 33.33                                  | (26/90) 28.89                    |
|          | Neihuang     | 30          | 6                    | 20                                     |                                  |
| Total    |              | 3198        | 1152                 | (1152/3198) 36.02                      |                                  |
sero-epidemiological investigation performed using ELISA to detect infections in small ruminants by *Babesia* species indicated that there was an average infection rate of 82.38 % for *Babesia* sp. BQ1 (Lintan) in 261 sera samples obtained from Lintan County in Gansu province, and these results were similar to our results (83.33 %) for this region [38].

The positive rates from 10 of the 23 provinces were comparable to those found in a recent molecular detection study that used a nested PCR assay based on the *rap1b* gene of *B. motasi*-like species, the positive rates for *Babesia* sp. BQ1 (Lintan), *Babesia* sp. BQ1 (Ningxian) and *Babesia* sp. Tianzhu were 5, 13.8 and 21.9 % in Inner Mongolia, Gansu and Xinjiang, respectively in this previous study (they were 61.01, 35.79 and 28.16 %, respectively, in our study) [58]. No *B. motasi*-like infections were detected in DNA samples obtained from the Zhejiang, Guizhou, Chongqing and Liaoning provinces when the nested PCR test was used [58]. Larger variation was observed in the numbers of positive samples that were identified in these provinces when ELISA was used in our study. A similar study was performed that compared the results of using ELISA (66.84 %) and the molecular LAMP method (14.3 %) to analyze *Babesia* sp. BQ1 (Lintan) infections in Gansu province [38]. Therefore, the difference in prevalence that is observed when using different testing methods (e.g., PCR vs ELISA or LAMP vs ELISA) or different antigens (e.g., BQMA vs rRAP-1a) or when using the same method (e.g., ELISA) can be potentially explained by the parameters that were used or the limitations of the detection methods used in these assays.

Some studies have indicated that *Haemaphysalis qinghaiensis* and *H. longicornis* are the vectors of the *Babesia* sp. BQ1 (Lintan) isolate, while *H. longicornis* has been proposed as the transmission vector for *Babesia* sp. BQ1 (Ningxian). These two tick species are widely distributed in at least 20 provinces of China (Fig. 1) [4, 59, 60]. Whereas, the higher positive rates (exceed 25 %) of *B. motasi*-like were found in Inner Mongolia, Xinjiang, Hunan, Zhejiang, Guangdong and Chongqing, as well as in Guangxi with positive rate of 9.74 % where the *H. longicornis* and *H. qinghaiensis* have not been reported yet. This has been explained by several possibilities by Wang et al. [57]. Moreover, cases of ovine babesiosis have been reported in many Chinese provinces, including Inner Mongolia, Henan, Shaanxi, Yunnan, Sichuan, Heilongjiang and Qinghai province [61–63]. Taken together, the results show that the wide prevalence of the *B. motasi*-like isolates (Lintan, Ningxian and Tianzhu) included in this study and co-infections between these three isolates is not surprising, and these isolates co-circulate in China with high prevalence. The results suggest that the development of ELISA methods that are based on the rRAP-1a61-1/CT described in our study indicate that this ELISA is a potential diagnostic method for epidemiological investigations of known related isolates in addition to other unidentified isolates in sheep *B. motasi*-like infections.

**Conclusions**

The present study demonstrates that antibodies against rRAP-1a61-1 can detect native RAP-1 in *Babesia* sp. BQ1 (Lintan) merozoite lysates in Western blot analysis and IFAT. The high conservation of RAP-1a across the three isolates from *B. motasi*-like parasites suggests that rRAP-1a61-1/CT could be a promising common diagnostic antigen that can detect geographically distinct isolates of *B. motasi*-like infections in sheep.

**Acknowledgements**

This study was financially supported by the NSFC (Nos. 31502054, 31372432, 31201899, 31272556, 31402189, and 31471967), ASTIP, Creative Research Groups of Gansu Province (No. 1210JJA06), NRCIS-CARS-38, the Special Fund for Agro-scientific Research in Public Research (Nos. 201303035 and 201303037), MOA, the 973 Program (2015CB150300), Supporting Program (2013BAD12B03, 2013BAD12B05), MOST (China), and the Jiangsu Co-innovation Center Programme for the Prevention and Control of Important Animal Infectious Diseases and Zoonoses as a State Key Laboratory of Veterinary Etiological Biology Project.

**Authors’ contributions**

QN, JY, PY and BZ performed the experiments, including cloning, expression, Western blot analysis, ELISA and IFAT. QN drafted the manuscript. ZL, EM, JL, GG and HY supervised all work. All authors read and approved the final version of the manuscript.

**Competing interests**

The authors declare that they have no competing interests.

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**Received:** 2 March 2016 **Accepted:** 5 May 2016

**Published online:** 31 May 2016

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