An improved ELISA for the detection of antibodies against *Babesia bovis* using either a native or a recombinant *B. bovis* antigen

R. Böse 1, R.H. Jacobson 2, K.R. Gale 1, D.J. Waltisbuhl 1, and I.G. Wright 1

1 CSIRO Division of Tropical Animal Production, Long Pocket Laboratories, Private Mail Bag 3, Indooroopilly, Queensland 4068, Australia
2 College of Veterinary Medicine, Cornell University Ithaca, NY 14853, USA

Accepted April 5, 1990

Abstract. Two new enzyme-linked immunosorbent assays (ELISA) for the diagnosis of *Babesia bovis* in cattle are described. The ELISA using a native antigen is more sensitive and less laborious than the assays described previously, because it does not require adsorption of sera with bovine erythrocytes. The second ELISA, using a recombinant *B. bovis* antigen expressed in *Escherichia coli*, was both sensitive and specific. It is suitable to replace the native antigen, thus avoiding large batch-to-batch variations in antigen preparations and the need to sacrifice experimental cattle.

**Materials and methods**

**Antigen preparations**

**Native antigen.** The oxy-haemoglobin free antigen was prepared from infected blood essentially as described previously (Mahoney et al. 1981). The protein concentration of the final preparation was 1050 μg/ml as estimated by the method of Bradford (1976), using bovine serum albumin as a standard and a protein assay reagent obtained from Bio-Rad Laboratories (Richmond, Va., USA).

**Recombinant antigen.** A fraction of a *B. bovis*-haemagglutinating antigen conferred protective immunity to cattle (Goodger et al. 1985). A monoclonal antibody designated W11C5 reacted with an antigen contained in this fraction, and the native W11C5-affinity-purified *B. bovis* antigen(s) induced immunity (Gale et al., personal communication).

A λ-GT11 cDNA expression library made from *B. bovis* ("Samford" strain) poly A + RNA was screened with the W11C5 monoclonal antibody. A cDNA clone was identified that expressed an approximately 160-kDa *B. bovis* antigen fused with *Escherichia coli* β-galactosidase (120 kDa) (Gale et al., personal communication) that was reactive with the W11C5 monoclonal antibody. The cDNA insert from this clone was subcloned into the plasmid expression vector pGEX-I (Smith and Johnson 1988) to facilitate the single-step purification of the resulting, approximately 180-kDa antigen-glutathione-S-transferase fusion protein. When supplemented with isopropylthiogalactoside, *E. coli* containing the pGEX-W11C5 construct accumulated soluble fusion protein to 20%-30% of cell protein as estimated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) (Neville 1971) using Coomassie blue stain. The fusion protein was affinity-purified using glutathione bound to epoxy activated beaded agarose (Sigma; St. Louis, Mo., USA) essentially as described by Smith and Johnson (1988). The yield was comparatively poor, as only a small proportion of the fusion protein bound to the affinity gel. However, the preparation obtained, containing 16 μg/ml protein, was pure as judged by SDS-PAGE using silver staining (Merril et al. 1981).

**Antisera**

**Standard sera.** Sera of high and low titre were obtained from cattle experimentally infected with *B. bovis* by blood inoculation and from those vaccinated with the recombinant antigen. Negative sera originated from cattle obtained from an area free from the tick vector *Boophilus microplus*. These cattle tested negative for haemoparasites by thick blood-film examination (Mahoney and Saal 1961) and for antibodies to *B. bovis* by IFAT (Johnston et al. 1973) at a serum dilution of 1/50 and higher. Sera of at least 12 cattle in each of 3 categories (high-positive, low-positive, and negative), as determined in preliminary ELISA studies, were pooled to obtain
(1) a high-positive, (2) a low-positive, and (3) a negative standard serum. Standard sera were stored frozen in aliquots at -80°C, thawed as needed, stored at 4°C, and used for no longer than 5 days after thawing.

Field sera. Sera obtained from *Bos taurus*, *Bos indicus*, and cross-bred cattle 4 months to 8 years of age that came from *Babesia*-endemic areas (although mostly of unknown parasitological status) were screened using both assays.

Enzyme-antibody conjugate

The conjugate used for most experiments was a pool of three monoclonal antibodies specific for bovine and/or ovine immunoglobulin (Ig) G1 and IgG2 coupled to hors eradish peroxidase (HRP) (Australian Monoclonal Development P/L; Artarmon, New South Wales, Australia).

Substrate solution

Substrate (5-aminosalicylic acid) recrystallised from a commercial-grade product (Sigma Chemical Co.) by the method of Ellins and Gielkens (1980) was dissolved in 0.1 M phosphate buffer (pH 6.0) at a concentration of 1 μg/ml heating to 50°C for 10 min, cooling to room temperature, and readjusting the pH 6.0 to 1 N NaOH. Substrate was prepared daily and H2O2 was added to a final concentration of 6 mM immediately before use. The concentration of a 30% H2O2 solution was determined and occasionally checked by spectrophotometry at 240 nm (Saunders et al. 1978); use of stock solution was discontinued if the concentration was < 8 M.

Assay procedure

The assays were designed as direct non-competitive ELISAs using microtitre plates (Greiner 655061, batch 192850) as a solid phase. Antigens were diluted in 0.1 M carbonate-bicarbonate buffer (pH 9.6) and 200 μl was added to each well and incubated overnight at 4°C, followed by three washes with PBS containing 0.05% (v/v) Tween 20 (PBS-T). To reduce non-specific binding of sera to the polystyrene surface, wells were blocked for 1 h at 37°C with 225 μl/well of PBS containing 2% horse serum. Blocking agent was removed and 200 μl bovine serum diluted in PBS containing 1% horse serum was added, followed by a 2 h-incubation at 37°C. Standard serum pools were applied in quadruplicate on each microtitre plate, whereas test sera were run in duplicate. Three washes with PBS-T were followed by a 1-h incubation at 37°C with 200 μl conjugate diluted 1/250 (2 μg/ml) in PBS containing 1% horse serum. Microtitre plates were washed again and the last washing solution was left in the wells until substrate addition. After substrate addition, microtitre plates were agitated on a shaking device (DSG Titertek/4, Flow Laboratories).

Data generation and evaluation

Absorbance values were measured using an ELISA reader (Titertek Multiscan Plus MKII, Flow Laboratories) interfaced with an IBM-compatible personal computer. Data was processed using the computer-based Kinetics Linked Immunosorbent assay program (KELA) essentially as described previously (Barlough et al. 1983). Briefly, the rate of peroxidase substrate reaction was calculated on the basis of three data points obtained by recording three absorbance readings (492 nm) at 2, 4, and 6 min after substrate addition. The regression coefficient or KELA slope value was calculated as the linear relationship between the rate of substrate conversion by enzyme and time. From 20 daily runs, the average value to be expected for the 3 standard serum pools (high- and low-positive and negative) was calculated. A nomograph was established daily, the obtained values were compared with the expected values, and the correlation was calculated. Values for all samples were normalised on the basis of the daily established nomograph to allow for day-to-day and plate-to-plate variation.

To allow for comparison of data generated using a single absorbance value, the correlation between KELA slopes (× 10^3) and the absolute absorbance values obtained 6 min after substrate addition was calculated on the basis of 600 individual readings [absorbance (492 nm, 6 min) = 0.035 + (0.006524 × KELA slopes (× 10^3)].

Results and discussion

Preliminary studies

Native antigen. Initially, the protocol of Waltisbuhl et al. (1987) was followed, except that adsorption of sera was omitted and the assay was analyzed by simultaneous testing of different concentrations of antigen, sera (the three different standards; ten individual negative sera), and conjugate [goat anti-bovine IgG (H+L) HRP, 100 μg protein/ml]. Analysis of data by KELA showed that the high dilutions (1/1000) of sera and conjugate gave comparatively low KELA slope values; therefore, further studies included log2 dilutions starting with 1/100 approx. 10.5 μg/ml for the antigen, 1/50 for the serum, and 1/125 (approx. 0.8 μg/ml) for the conjugate.

These experiments showed that the main problems involved a generally high background activity and, in particular, unacceptably high reactions for some of the individual negative sera. However, these phenomena directly correlated with the serum and conjugate concentrations and were largely independent of the antigen concentration. Moreover, negative sera reacted even without the prior addition of antigen to the microtitre plate. This was not related to haemolysis. Obviously, serum components recognised by the affinity-purified conjugate were adhering to the polystyrene surface. Different blocking agents, such as gelatin, hen-egg albumin, and horse serum, and inclusion of Tween 20 in the diluting buffer were evaluated for their ability to prevent this. Horse serum was best, but it produced only partial improvement. Because the non-specific reactions were also conjugate-dependent, a variety of different conjugates, conventional or affinity-purified, were tested. Although some conjugates were more suitable than others, none was entirely satisfactory.

Non-specific reactions were virtually abolished when the IgG-specific monoclonal antibody conjugate was used. Therefore, the serum components binding avidly to polystyrene are probably not IgG. It is possible that IgM was causing the non-specific reactions, as conventional conjugates against bovine IgG also react with IgM. The monoclonal antibody conjugate was used exclusively for all further studies, although it was recognised that IgM antibodies are not detected in the early stage of infection.

Recombinant antigen. Prior to the expression of the antigen as a G-S-T fusion protein, preparations of a β-galac-
tosidase fusion protein were purified by various means (e.g., antibody affinity chromatography, gel filtration). All preparations obtained, even if contaminated with only minor amounts of *E. coli* proteins, proved unsatisfactory and gave poor discrimination, i.e., unacceptably high reactions with negative sera that were presumably due to anti-*E. coli* antibodies present in most bovine sera. To assess critically whether the antigen preparation judged to be pure by SDS-PAGE would be suitable, sera from cattle vaccinated with crude *E. coli* lysate were tested by ELISA. Reactions occurred in the range of negative sera, and adsorption of sera with *E. coli* lysate lowered the responses only marginally.

**Optimization of the assays**

Various concentrations of all reagents used were tested simultaneously and data were evaluated by the KELA program. Both assays gave a good discrimination over a wide range of antigen dilutions, i.e., 1/100–1/1600 for the native antigen and 1/10–1/3200 for the recombinant antigen. Within the given range, the reactions for positive sera increased with antigen concentration, whereas negative sera gave very low and almost identical reactions regardless of the antigen concentration. To conserve antigen, the concentration giving about 90% of the maximal reading observed for the high-positive standard serum pools was chosen as the working dilution. In summary, dilution conditions for optimal discrimination were: 1/400 for the native and 1/200 (approx. 0.08 µg/ml) for the recombinant antigen; 1/100 for bovine sera; and 1/250 (approx. 2 µg/ml) for the monoclonal antibody conjugate. One blocking step using PBS containing 2% horse serum before the addition of bovine sera and the dilution of sera in PBS containing 1% horse serum were optimal.

**Assay evaluation under optimised conditions**

**Calculation of the threshold.** Sera from 72 cattle that originated from an area free of *Boophilus microplus* and tested negative for haemoparasites by thick blood-film analysis (Mahoney and Saal 1961) as well as for antibodies against *B. boris* by IFAT (Johnston et al. 1973) were used to determine the threshold, calculated as the average of KELA slope values (×10²) plus 3 standard deviations. Values were calculated for native antigen [average, 4.0 (range, 2.1–6.0); SD, 0.9; threshold, 6.7] and for recombinant antigen [average, 1.3 (range, 0–3.5); SD, 0.8; threshold, 3.7].

**Validation of assays.** Both assays were sensitive and specific and compared well with previously published results (Table 1). The native antigen ELISA was more sensitive than that previously reported by Waltisbuhl et al. (1987), probably due to several factors. First, in the present study the threshold was reduced to only about 4% of the high-positive standard, whereas Waltisbuhl et al. (1987) obtained a threshold equalling 20% of the reaction for the positive control. Second, we used a complete antigen, whereas Waltisbuhl et al. (1987) employed a fractionated antigen preparation. Finally, our serum concentration (1/100) was considerably higher than that of the previous study (1/1000).

**Adsorption of sera with bovine erythrocytes.** The haemagglutination assay (HA) and IFAT require the adsorption of sera, as isoantibodies against erythrocytes cause false-positive reactions. Isoantibodies apparently also interfered with the ELISA; adsorption reduced the number of false-positives in this test as well. However, when the new protocol was adopted, we found that adsorption had practically no effect and could be omitted (Fig. 1). This offers further support for our observation in preliminary studies that false-positive reactions are due to conjugate non-specificity rather than to antigen impurities.

It is possible that isoantibodies are IgM by nature and, although not recognised by the IgG-specific monoclonal antibody, give rise to false-positive reactions when conventional conjugates are used. Alternatively, or in addition, adsorption may have removed or inactivated serum components that otherwise adhere to the polystyrene surface of the microtitre plate, which, again, were recognised by conjugates that had been used before but not by the monoclonal antibody conjugate.

**Cross-reactivity with other haemoprotezoa.** Sera from 36 cattle that had other haemoprotezoan infections detected by thick blood-film examination (Mahoney and Saal 1961) were also tested. In all, 10 cattle infected with *B. bigemina* (2–4 weeks after infection by blood inoculation), 12 infected with *Theileria orientalis* (field infec-

---

**Table 1. Comparison of the native and the recombinant antigen ELISA with previously published techniques using sera from cattle of known parasitological status**

| Time after infection | Previous results | ELISA results |
|---------------------|------------------|---------------|
|                     | Haemagglutination | Native | Recombinant |
|                     | ELISA            | antigen | antigen |

| Condition          | ELISA results |
|--------------------|---------------|
| Infected (n=22)    | 20/2          |
| Not infected (n=10)| 2/8           |
| Infected (n=21)    | 14/7          |
| Not infected (n=10)| 2/8           |

x/y = cattle sero-positive/sero-negative
Sera of 32 cattle were tested; 22 cattle were single-infected with *B. boris* and 10 were kept as negative controls (Mahoney et al. 1979)

* Goodger and Mahoney (1974)
* Waltisbuhl et al. (1987)
Correlation of the assays. Qualitatively the assays agreed to 93% (280/300) (Table 2). Statistically, only a moderate correlation between the two assays was found (Fig. 2). This moderate correlation was anticipated because the native antigen preparation contains a large number of different antigens, whereas the recombinant antigen represents only one of these \emph{B. boris} antigens. Nevertheless, the single recombinant antigen enables the reliable detection of most infected cattle. Most naturally infected animals developed antibodies against this antigen, which indicated a high degree of conservation between parasite strains. The recombinant antigen ELISA remains at least as sensitive as the assays reported thus far (Table 1) but has the slight disadvantage of lower sensitivity in some cases. This could probably be overcome by use of a cocktail of two or more recombinant antigens, and 14 infected with \emph{Anaplasma marginale} (9 days to 8 weeks after infection by blood inoculation) showed negative reactions in both assays.

Table 2. Comparison of the native and the recombinant antigen ELISA using 300 sera from cattle mostly of unknown parasitological status

| Number of sera | KELA slope (× 10^3) |
|----------------|----------------------|
|                | Range               | Average |
| Negative reactions: |                     |         |
| Native antigen  | 147                  | 1.5–6.6 | 3.9 |
| Recombinant antigen | 145                | 0–3.6  | 1.8 |
| Positive reactions: |                     |         |
| Native antigen  | 153                  | 6.9–227.8 | 55.1 |
| Recombinant antigen | 155                | 3.7–83.8 | 18.6 |
| Positive reactions by one antigen only: |                     |         |
| Native antigen  | 9^a                  | 6.9–77.4 | 24.3 |
| Recombinant antigen | 11^b                | 3.7–7.7 | 5.1 |

^a 4 cattle were infected with \emph{B. boris} as detected by thick blood-film examination, 5 were of unknown parasitological status

^b 2 cattle were infected with \emph{B. boris} as detected by thick blood-film examination, 9 were calves about 7 months old, held in a herd in which all cattle over the age of 1 year were sero-positive
antigens, which would presumably also lead to a closer correlation with the native antigen ELISA.

Interestingly, the recombinant antigen detected some cattle that were not positive by the native antigen ELISA; 2 of these 11 cattle were infected with B. bovis, whereas the remaining 9 calves were probably positive due to maternal antibodies (Table 2). Colostral antibodies against B. bovis have been detected for as long as 68 days by HA (Goodger and Mahoney 1974) and for as long as 6 months after birth by radioimmunoassay (Wright 1990). Transferred antibodies against B. bigemina can be detected by IFAT as long as 170 days after colostrum intake (Weisman et al. 1974). Given the high sensitivity of the ELISA, it is likely that the marginally positive reactions are due to maternal antibodies. Nevertheless, only the examination of a series of blood samples from calves with maternal antibodies could prove this interpretation.

We demonstrated that a single recombinant antigen can be used successfully for the sero-diagnosis of B. bovis. The major advantages were a negligible batch-to-batch variation in antigen and the absence of a need to sacrifice experimental animals for the preparation of native antigens. In combination with more defined conjugates, recombinant antigens should lead to greater comparability and reproducibility of serological results.

Acknowledgements. The senior author was supported by a research grant from the Deutsche Forschungsgemeinschaft. We would like to thank Dr. B.V. Goodger for his helpful advice during the experimental work and his critical review of the manuscript.

References

Barlough JE, Jacobson RH, Downing DR, Marcella KL, Lynch TJ, Scott FW (1983) Evaluation of a computer-assisted, kinetics based enzyme-linked immunosorbent assay for the detection of coronavirus antibodies in cats. J Clin Microbiol 17: 202-217
Bradford MM (1976) A rapid and sensitive method for the quantitation of microgram quantities of protein using the principle of dye binding. Anal Biochem 72: 248-254
Ellens DJ, Gielkens ALJ (1980) A simple method for the purification of 5-aminosalicylic acid. Application of the product as substrate, in an enzyme-linked immunosorbent assay (ELISA). J Immunol Methods 37: 325-332
Goodger BV (1971) Preparation and preliminary assessment of purified antigens in the passive haemagglutination test for bovine babesiosis. Aust Vet J 47: 251-256
Goodger BV, Mahoney DF (1974) Evaluation of the passive haemagglutination test for bovine babesiosis. Aust Vet J 50: 246-249
Goodger BV, Wright IG, Waltisbuhl DJ, Mirre GB (1985) Babesia bovis: successful vaccination against homologous challenge in splenectomised calves using a fraction of haemagglutinating antigen. Int J Parasitol 15: 175-179
Johnston LAY, Pearson RD, Leatich G (1973) Evaluation of an indirect fluorescent antibody test for detecting Babesia argentina infection in cattle. Aust Vet J 49: 373-377
Mahoney DF, Saal JR (1961) Bovine babesiosis: thick blood films for the detection of parasitaemia. Aust Vet J 37: 44-47
Mahoney DF, Wright IG, Goodger BV (1979) Immunity in cattle to Babesia bovis after single infections with parasites of various origin. Aust Vet J 55: 10-12
Mahoney DF, Wright IG, Goodger BV (1981) Bovine babesiosis: the immunization of cattle with fractions of erythrocytes infected with Babesia bovis (syn. B. argentina). Vet Immunol Immunopathol 2: 145-156
Merrill CR, Goldman D, Sedman SA, Ebert MH (1981) Ultrasensitive stain for proteins in polyacrylamide gels shows regional variation in cerebrospinal fluid proteins. Science 211: 1437-1438
Neville DM Jr (1971) Molecular weight determination of protein-dodecyl sulfate complexes by gel electrophoresis in a discontinuous buffer system. J Biol Chem 246: 6328-6334
Saunders GC, Clinard EH, Bartlett ML, Petersen PM, Sanders WM, Payne RJ, Martinez E (1978) Serologic test systems development, July 1, 1976-September 30, 1977. Progress report La-7078-PR. Los Alamos Scientific Laboratory of the University of California, Los Alamos, New Mexico
Smith DB, Johnson KS (1988) Single step purification of polypeptides expressed in Escherichia coli as fusions with glutathione S-transferase. Gene 76: 31-40
Waltisbuhl DJ, Goodger BV, Wright IG, Commins MA, Mahoney DF (1987) An enzyme-linked immunosorbent assay to diagnose Babesia bovis infection in cattle. Parasitol Res 73: 126-131
Weisman J, Goldman M, Mayer E, Pipano E (1974) Passive transfer to newborn calves of maternal antibodies against Babesia bigemina and Babesia berbera. Refu Vet 31: 108-113
Wright IG (1990) Babesiosis. In: Rött IM, Delves PF (eds) Encyclopedia of Immunology. Saunders Scientific (in press)