New serological platform for detecting antibodies against *Mycobacterium tuberculosis* complex in European badgers

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Abstract

European badgers (*Meles meles*) have been identified as wildlife reservoirs for *Mycobacterium bovis* in the UK and Ireland, and may also have a role in the epidemiology of animal tuberculosis in other European regions. Thus, detection of *M. bovis*-infected badgers may be required for the purposes of surveillance and monitoring of disease levels in infected populations. Current serological assays to detect *M. bovis* infection in live badgers, while rapid and inexpensive, show limited diagnostic sensitivity. Here we describe and evaluate new ELISA platforms for the recognition of the P22 multiprotein complex derived from the purified protein derivative (PPD) of *M. bovis*. The recognition of IgG against P22 multiprotein complex derived from PPD-B was tested by ELISA in the serum of badgers from the UK, Ireland and Spain. TB infection in the badgers was indicated by the presence of *M. bovis* in tissues by culture and histology at post-mortem examination and TB-free status was established by repeated negativity in the interferon γ release assay (IGRA). In experimentally infected badgers, humoral antibody responses against P22 developed within 45 days post-infection. The ELISA tests showed estimated sensitivity levels of 74–82% in experimentally and naturally infected badgers with specificities ranging from 75% to 100% depending on the badger population tested. The P22 multi-antigen based ELISAs provide a sensitive and specific test platform for improved tuberculosis surveillance in badgers.

Keywords: Badgers, diagnosis, P22 ELISA, tuberculosis, wildlife reservoir.

Introduction

Tuberculosis caused by *M. bovis* and closely related members of the *Mycobacterium tuberculosis* complex remains one of the most important infectious diseases in cattle. Tuberculosis in cattle is controlled in Europe through routine test and slaughter policies. However, eradication of the disease can be hindered by the persistence of infection in wild animal reservoirs (Corner 2006) and European badgers (*Meles meles*) play an important role in the transmission cycle of bovine tuberculosis in the UK and Ireland (Donnelly *et al.* 2007; More 2009; Murphy *et al.* 2010) and possibly in other European regions (Balseiro *et al.* 2013; Bouchez-Zacria *et al.* 2017).

The most sensitive tests for diagnosing tuberculosis in live badgers are currently based on cell-mediated responses (Dalley *et al.* 2008; Balseiro *et al.* 2011). These tests can be difficult to implement...
on a large scale because they (1) require sufficient volumes of fresh blood, which need to be processed within a few hours after collection; (2) require more sophisticated expertise and laboratory equipment than ELISAs; (3) are more expensive than ELISAs; (4) require longer processing time and (5) cannot be used with post-mortem samples. Antibody detection tests, in contrast, can be used to screen large numbers of animals because they are simple to perform on fresh and stored samples, require small sample volume, are relatively inexpensive and can be completed within 24 h. They can be performed and interpreted in parallel with cellular immune assays to increase overall sensitivity across the full spectrum of disease including when cellular anergy occurs (Green et al. 2009; Whelan et al. 2010; Chambers et al. 2011; Chambers 2013; Waters et al. 2015; Casal et al. 2017). However, the serological tests currently available for diagnosing TB in badgers, the DPP® VetTB (lateral-flow antibody detection test by Chembio, NY, USA) (Lyashchenko et al. 2013; Schaftenaar et al. 2013; Che’ et al. 2015; Vogelnest et al. 2015) that has recently replaced the BrockTB STAT-PAK (Chembio, NY, USA) and a chemiluminescent multiplex immunoassay from Enfer Scientific (Co. Kildare, Ireland) (Aznar et al. 2017) are relatively insensitive: approximately 55% (S. Lesellier, personal communication) and 25.3% (Aznar et al. 2017) respectively in naturally infected badgers, depending on the cut-off values applied. Both assays primarily target MPB83, MPB70, ESAT-6 and CFP-10.

In order to provide an expanded range of tests for large-scale testing of M. tuberculosis complex infection in badgers, we have developed ELISA systems targeting antibodies against the recently described P22 multiprotein antigen (Infantes-Lorenzo et al. 2017), which is affinity-purified from bovine purified protein derivative (PPD) of M. bovis and has been shown to provide greater diagnostic sensitivity than conventional ELISAs in other host species in a high prevalence setting (Casal et al. 2017). In this study the P22-based ELISAs were evaluated using serum samples from badgers infected naturally and experimentally with M. bovis and uninfected captive and wild badgers from three different European locations.

**Material and methods**

**Serum sample collection**

All serum samples were originally obtained by jugular venipuncture of anaesthetised badgers into serum separation vacutainer tubes (SST, Becton Dickinson™, New Jersey, USA). The general anaesthesia of the badgers was induced by intramuscular injection of ketamine hydrochloride (approximately 10 mg/kg, Vetalar, Boehringer Ingelheim) and medetomidine hydrochloride (approximately 0.1 mg/kg, Domitor, Pfizer) in Ireland, supplemented with butorphanol (approximately 0.1 mg/kg, Torbugesic R, ZoetisUKLtd, Tadworth, Surrey, UK) in the UK and Spain.

**Serum samples from uninfected and experimentally infected badgers from the UK**

In the UK, a total of 36 captive badgers were infected with an average of $3 \times 10^3$ CFU/mL viable M. bovis (spoligotype 9, strain 74/0449/97) by endobronchial instillation as described previously (Lesellier et al. 2011). Serum samples were obtained before the start of the study and at 0 (pre-challenge), 15, 30, 45, 60 and 75 days post-infection (dpi) (Table 1). All animals presented with TB visible lesions 75 days post-infection (dpi) and M. bovis was cultured from the animal tissues (S. Lesellier, in preparation).

**Serum samples from naturally infected and uninfected badgers from Ireland**

Serum samples were originally collected from 53 badgers trapped at the end of a vaccine field study in Ireland (Gormley et al. 2017), and examined at post-mortem. Animals were classified as tuberculosis-positive (n = 25), if lesioned tissue sections showed the presence of acid-fast bacilli based on Ziehl-Neelsen staining or if at least one tissue gave a positive result in the M. bovis culture test. The remaining animals (n = 28) were classified as tuberculosis-negative (Table 1).
Thirty-two badgers were captured in Asturias and tested negative for badger interferon gamma release assay (Table 1). These findings, together with the low local prevalence of tuberculosis in cattle and badgers (6.6% based on bacteriological culture from road-killed badgers and 0.21% in cattle, Balseiro et al. 2011), allowed us to classify these animals as tuberculosis-negative.

**Indirect P22 ELISA**

An indirect ELISA was developed to detect antibodies against the *M. tuberculosis* complex P22 complex. This complex was affinity-purified from bovine PPD [CZ Veterinaria (Porrino, Spain)] using a patented process (European Patent EP16382579) (Infantes-Lorenzo et al. 2017). The ELISA was developed using previously described methods (Casal et al. 2017), with minor modifications. Maxisorp plates were coated with 10 µg/mL P22 overnight at 4°C in phosphate-buffered saline (PBS), then blocked with 5% skimmed milk powder solution (Central Lechera Asturiana, Spain) in PBS for 1 h at room temperature. After three washes with PBS containing 0.05% Tween-20 (PBST), sera (100 µL) were added to duplicate wells at 1:100 dilution in PBS-skim milk and incubated for 60 min at 37°C. Horseradish peroxidase-conjugated CF2/HRPo anti-Badger IgG (100 µL) (Goodger et al. 1994) was diluted to 1.5 µg/mL in PBS and added to the plates, which were incubated for 30 min at room temperature. Then plates were incubated with 3,3′,5,5′-tetramethylbenzidine substrate (Perbio) for 15 min in the dark at room temperature. The reaction was stopped by adding 100 µL of 2 mmol/L H₂SO₄. Optical density was measured at 450 nm using an ELISA reader.

Negative control serum samples from UK tuberculosis-free captive badgers were included in every plate in quadruplicate. Positive controls were obtained from UK badgers experimentally infected with *M. bovis*. Sample results were expressed as an ELISA percentage E%, calculated using the following formula (Che’ et al. 2015):

\[
E% = \frac{\text{mean sample optical density}}{2} \times \frac{\text{mean of negative control optical density}}{\times 100%}
\]

Cut-off points were set-up to calculate sensitivity and specificity: E% > 100, E% > 120 and E% > 150.

**Competitive P22 ELISA**

As environmental mycobacteria can potentially interfere with tuberculosis diagnostic tests, we developed a competitive ELISA against P22 with the objective to exclude interference from cross-reactive antibodies recognising antigens from environmental mycobacteria. This assay was developed and evaluated in the same way as the indirect ELISA, except that sera were diluted in skimmed milk supplemented with avian PPD at 150 µg/mL (Casal et al. 2017). Samples were always assayed in parallel using the indirect and competitive ELISAs.
Statistical analysis

Data were analysed using SPSS 20.0 (IBM, Somers, NY, USA). A Wilson 95% confidence interval (95% CI) was calculated for each percentage.

Results

Experimentally infected badgers from UK

The proportion of sero-positive animals increased with time post-infection in the indirect and competitive ELISAs (Fig. 1). Before challenge (pre-challenge and T0), seven of the 36 badgers (19.44%) tested positive by indirect ELISA. One additional animal was only positive on one of these occasions and was not included in results shown in Table 2. As these animals were confirmed tuberculosis-free at the time of testing, this pre-challenge positivity was attributed to cross-reactivity with other antigens similar to P22, possibly antigens secreted by *M. avium* or other environmental mycobacteria. Only one of the seven badgers that tested positive in the indirect ELISA tested negative in the competitive ELISA.

The indirect and competitive ELISAs showed similar sensitivity of detection (82% and 79% respectively) of infected animals at the end of the study, day 75 (Table 2). At 60 dpi, 73.52% were found positive by the indirect ELISA (Fig. 1). As infection progressed, the mean E% increased from 127.30% before challenge to 336.26% at 75 dpi (Kruskal-Wallis test, d.f. = 6, *P* < 0.001). The largest increase in the proportion of positive animals occurred approximately 1 month after infection, when the level of circulating antibodies was expected to rise. The rate of positive test responses for the period from 0 to 30 dpi differed significantly from the rate for the period from 45 to 75 dpi (Wilcoxon signed-rank test, *P* < 0.001).

Naturally infected and uninfected badgers from Ireland

Of 25 naturally infected badgers, 20 tested positive by the indirect ELISA and 19 by the competitive ELISA for the cut-off point of 100%, yielding mean sensitivities of 80% and 76% respectively. The highest values tended to occur in animals with visible lesions (data not shown). The specificity of the competitive and indirect ELISAs was 85.71% when a cut-off of E% = 100% was used, and increased to 89.29% with a cut-off of 120%, but sensitivity decreased slightly (Table 2). The specificity was marginally lower in the indirect ELISA than in the competitive ELISA with both cut-offs (75% and 85% respectively) (Table 2).

Figure 2 presents the median levels of serological responses, which differed significantly between naturally infected and uninfected badgers (Mann–Whitney, *P* < 0.001).

Uninfected badgers from Spain

Of 32 tuberculosis-free Spanish badgers, only one tested positive in the indirect ELISA, giving a specificity of 96.88%. No animal tested positive in the competitive ELISA, giving a 100% specificity (Table 2).
Discussion

Our results suggest that the indirect and competitive P22 ELISA tests described here have the potential to provide greater levels of sensitivity compared to other anti-\textit{M. tuberculosis} complex antibody detection tests currently used in badgers. These inexpensive and straightforward tests require small serum volumes and their application could help improve tuberculosis diagnostics and field studies in badgers.

In experimentally infected badgers, we obtained high sensitivities ranging from 73.52\% at 60 dpi to 82.35\% at 75 dpi. When applied to samples from badgers from Ireland, the indirect ELISA showed sensitivity levels of 80\% and the competitive ELISA showed 76\% sensitivity. These are considerably higher than the sensitivities of 25.3–55\% reported for other antibody tests for badgers (Clifton-Hadley \textit{et al.} 1995; Greenwald \textit{et al.} 2003; Chambers \textit{et al.} 2010; Aznar \textit{et al.} 2017). They are also consistent with high sensitivity observed in infected cattle (Casal \textit{et al.} 2017). From a disease management point of view, the test would likely be applied in populations with high infection levels, using high levels of sensitivity in order to maximise the detection of infected animals.

The specificity of the serological assays varied between badger populations/countries of origin, possibly as the result of potential exposure to different environment mycobacterial species. It reached 100\% in a Spanish badger captive population and 90\% in uninfected badgers from the Republic of Ireland when using the cut-off point of \( E = 120\% \) in the competitive ELISA. However, lower specificity

\[ \text{Table 2. Proportion (\%) of badgers testing positive in an indirect (ID) or competitive (CP) ELISA detecting serum antibodies against the } \]

\begin{tabular}{|c|c|c|c|c|c|c|}
\hline
 & n Animals & & & & & \\
 & & Cut-off 100\% & & Cut-off 120\% & & Cut-off 150\% & \\
 & & ID & CP & ID & CP & ID & CP \\
\hline
TB infected & UK & 34 & 82 (66–92) & 79 (63–90) & 73.53 (57–85) & 73.53 (57–85) & 64.71 (48–79) & 64.71 (48–79) \\
 & Ireland* & 25 & 80 (61–91) & 76 (57–89) & 76 (57–89) & 64.71 (48–79) & 64.71 (48–79) \\
Spain & 32 & 3.13 (1–16) & 0 (0–11) & 0 (0–11) & 0 (0–11) & 0 (0–11) & 0 (0–11) \\
\hline
\end{tabular}

Intervals in parentheses indicate 95\% confidence limits. *Based on \textit{M. bovis} culture test. †No visible lesions and culture-negative.
results were recorded when we used samples from apparent tuberculosis-free badgers from the UK. This is suggestive of different levels of immune sensitisation in the different countries: *M. avium* infection in badgers has previously been reported (Balseiro *et al.* 2011). In addition, we cannot rule out that all of the UK badgers were genuinely TB free. The different specificity levels for the same interpretation criteria suggest that the UK badgers used in this study may have been previously exposed to cross-reactive antigens homologous to P22.

The competitive ELISA using avian PPD antigen did not resolve the relatively low specificity measured with samples from UK badgers. Although P22 is composed mainly of MPB70 and MPB83, it shares several proteins with the *M. avium* complex that may contribute to cross-reactivity (Infantes-Lorenzo *et al.* 2017). In addition, MPB70 and MPB83 show high sequence similarity to proteins from nontuberculous mycobacteria that do not form part of the *M. avium* complex, such as proteins from *M. kansasii* (77% identity, 98% coverage) and *M. abscessus* (73% identity, 82% coverage) (www.uniprot.org, accessed 17 November 2017). For screening tests specificity is particularly important in areas of low tuberculosis prevalence, as it minimises the disclosure of false-positive test results. However, a potential problem with determining overall specificity in different environments is that the causative cross-reactive agent is often unknown, and it is likely to vary between regions. The most appropriate way to address this is to modify the cut-off points to take into account the many confounders in local environments. Specificity can be improved by adjusting the cut-off value, albeit potentially at the expense of sensitivity. With a cut-off value of 100%, the indirect ELISA showed specificity levels of 75% and the competitive ELISA showed a specificity of 85.71%. Increasing the cut-off to 120% increased specificity of the indirect assay to 85.71% and specificity of the competitive assay to 89.29%, while sensitivity remained stable at 76%.

It is possible that the true specificity of our assays is even higher than we measured, because bacterial culture is not a gold standard test, even when performed on necropsy tissues (Crawshaw *et al.* 2008). Thus, it is possible that some badgers were infected with members of *M. tuberculosis* complex but were not detected post-mortem based on mycobacterial culture or pathology. Molecular characterisation of *M. tuberculosis* complex strains isolated from animals in the UK has revealed the presence of *M. microti*, and a small proportion of the spoligotypes was associated with badgers (Smith *et al.* 2009). Although we have no evidence of *M. microti* infections in any of the badgers used in this study, we cannot rule out the possibility that any such infection could potentially impact on the specificity of the test. Despite our ELISAs disclosing positive results for 20 of 25 culture-positive, naturally infected badgers, some of the culture-negative animals may have had subclinical infection; this would affect the reliability of our measured sensitivity and specificity. The results from the different countries highlight that test performance can vary according to different environments and cut-off point needs to be established and adapted to local conditions. Nevertheless, our high specificity observed with samples from uninfected badgers from Spain suggests that both ELISAs perform well.

The rates of test positive results in each ELISA increased with disease progression. This indicates an increase in sensitivity associated with more advanced infection, as reported in experimentally infected wild boar (*Sus scrofa*) and experimentally infected red deer (*Cervus elaphus*) (Garrido *et al.* 2011; Thomas *et al.* 2017). In badgers, the sensitivity of the BrockTB STAT-PAK appeared to be higher in animals with more severe tuberculosis; those with lesions visible at post-mortem (Clifton-Hadley *et al.* 1995; Chambers *et al.* 2008). This ability to detect animals with advanced disease is particularly important because they are potentially the principal shedders of *M. bovis*, and may pose a higher risk of bacterial transmission to livestock (Gallagher *et al.* 1998; Delahay *et al.* 2013).

**Conclusion**

In conclusion, the development of novel serological tests based on the P22 multiprotein antigen may lead to more sensitive and specific ELISAs, and prove to be an effective tool to support the detection and management of tuberculosis in badgers.
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**Conflicts of interest**

The authors declare that they have no conflicts of interest.

**Ethics statement**

All animal procedures conducted for this study were approved by the Animal Welfare and Ethical Review Board at the UK Animal and Plant Health Agency (APHA) [under the Animal (Scientific Procedures) Act 1986 and Home Office authorized Project License 70/7878], the University College Dublin Animal Research Ethics Committee (AREC-P-08-26), and research licence (B100/3187) issued by the Dept of Health & Children, Ireland, and the Animal Research Ethics Committee of the Principado de Asturias, Spain (PROAE 20-2015).

**Contributions**

JIL and DD performed the experiments, analyzed and interpreted the data and wrote the first draft of the manuscript. JIL, SL, EG and JS were involved in interpreting the data. DD, PA, SL, AB and EG obtained serum samples from animals. CG and LD analyzed the data and revised the manuscript. JIL, JS, SL and MD designed the study, analyzed the data, and revised the manuscript. IM, SL, EG, LD, AB and CG critically revised the manuscript. All authors have read and approved the final manuscript.

**References**


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