Evaluation of a Rapid Serological Test for the Determination of *Mycobacterium bovis* Infection in Badgers (*Meles meles*) Found Dead

Mark A. Chambers  
*TB Research Group, Department of Statutory and Exotic Bacterial Diseases, Veterinary Laboratories Agency Weybridge, New Haw, Surrey KT15 3NB, United Kingdom*

Konstantin P. Lyashchenko  
*Chembio Diagnostic Systems, Inc., Medford, New York*

Rena Greenwald  
*Chembio Diagnostic Systems, Inc., Medford, New York*

Javan Esfandiari  
*Chembio Diagnostic Systems, Inc., Medford, New York*

Eurig James  
*Veterinary Laboratories Agency Carmarthen, Johnstown, Carmarthen SA31 3EZ, United Kingdom*

*See next page for additional authors*

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Mark A. Chambers, Konstantin P. Lyashchenko, Rena Greenwald, Javan Esfandiari, Eurig James, Leslie Barker, Jeff Jones, Gavin Watkins, and Simon Rolfe

*TB Research Group, Department of Statutory and Exotic Bacterial Diseases, Veterinary Laboratories Agency Weybridge, New Haw, Surrey KT15 3NB, United Kingdom; Chembio Diagnostic Systems, Inc., Medford, New York 11763; Veterinary Laboratories Agency Carmarthen, Johnstown, Carmarthen SA31 3EZ, United Kingdom; Veterinary Laboratories Agency Shrewsbury, Harlescott, Shrewsbury SY4 4HD, United Kingdom; and Office of the Chief Veterinary Officer, Welsh Assembly Government, Cathays Park, Cardiff CF10 3NQ, United Kingdom*

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Between October 2005 and May 2006, a total of 727 badgers found dead in Wales were reported, and 550 were delivered to the Regional Laboratories of the Veterinary Laboratories Agency (VLA). Of the 459 carcasses suitable for examination, 55 were deemed to be infected with *Mycobacterium bovis* on the basis of culture, spoligotyping, and variable-number tandem repeat typing. Acid-fast bacteria were observed histologically in a further six badgers, but these bacteria were not confirmed as *M. bovis* by culture. A rapid serological test (BrockTB Stat-Pak) performed on thoracic blood showed a sensitivity of 35% and a specificity of 99%. Presence of *M. bovis* infection was 45 times more likely to be confirmed postmortem by culture in BrockTB Stat-Pak-reactive animals than in seronegative ones. Using visible carcass lesions as a marker of bovine tuberculosis (bTB) infection had a similar sensitivity (38%) but was significantly less specific (84%) than serology. The overall accuracy of the antibody detection was 93% (346 correct results from 374 tests), whereas the accuracy of regarding visible lesions as a marker for bTB infection was 78% (354 correct from 453 carcasses examined). Culture remains the gold standard method for detecting *M. bovis* infection in badgers. However, where resources are limited and/or an instant result is preferred, the BrockTB Stat-Pak could be used in field surveillance efforts to identify animals which should be examined further by only submitting test-negative animals to more detailed postmortem examination and culture.

*Mycobacterium bovis* infection is the cause of bovine tuberculosis (bTB) in a wide range of mammal species, including domestic livestock and captive and free-ranging wildlife. Bovine TB remains an important zoonotic disease with significant impacts on the economy in many countries (6, 22, 23). Eurasian badgers (*Meles meles*) are a wildlife maintenance host of bTB in Great Britain and Ireland (5, 15) and are implicated in the maintenance and onward transmission of *M. bovis* infection to cattle (10, 19).

Surveillance of wildlife vectors of disease for prevalence estimates of infection may be valuable in disease control strategies and for the assessment of risk of transmission to livestock. Diagnosis of bTB in live badgers has been demonstrated using assays of both serological (4, 20) and cell-mediated (8, 9) immunity. While isolation of *M. bovis* from clinical samples is definitive, it is too insensitive for badgers, as infected animals yield positive samples infrequently and intermittently (3). A rapid serological test (BrockTB Stat-Pak; Chembio Diagnostic Systems, Inc.) has recently been developed for the diagnosis of bTB in multiple wildlife species (20). The test has modest sensitivity (46 to 55%) for antibody detection in live, infected badgers, but it has the advantages of being simple, rapid, inexpensive, and suitable for field application. Its utility as an animal-side test for badgers, however, is limited by the difficulties associated with obtaining a blood sample from a nonanesthetized animal.

Where carcasses are recovered and submitted for mycobacterial culture, the sensitivity of diagnosis depends on the effort taken for careful examination and on the number of tissue samples submitted for culture testing and histopathology (7), as well as on the condition of the carcass. In many cases, the cost involved may prove prohibitive. Reliance on the presence of visible lesions as indicative of bTB is fraught with difficulties, as infected animals may present with no visible lesions or lesions may be the result of other infections while having the appearance of bTB (reviewed in reference 13). The purpose of this study was to determine whether the BrockTB Stat-Pak test could detect *M. bovis* antibody in blood collected from the carcasses of dead badgers as an alternative means of diagnosis and decision making. Animals were obtained as part of a separate government-funded study to determine the prevalence of bTB in badgers found dead in Wales (http://new.wales.gov.uk/depc/publications/environmentandcountryside/animalhealthandwelfare/disease surveillanc econtrol/bovineb/2567889/publicationindex/2326585/badgerfounddeadreport?lang=en). Our results reveal that the BrockTB Stat-Pak test used on thoracic blood samples

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* Corresponding author. Mailing address: TB Research Group, Department of Statutory and Exotic Bacterial Diseases, Veterinary Laboratories Agency Weybridge, Woodham Lane, New Haw, Surrey KT15 3NB, United Kingdom. Phone: 44 1932 357494. Fax: 44 1932 357260. E-mail: m.a.chambers@vla.defra.gsi.gov.uk.

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was very specific (99%) but less sensitive (35%) than found previously for live badgers (2, 14). However, bTB was 45 times more likely to be confirmed in BrockTB Stat-Pak-positive animals than in BrockTB Stat-Pak-negative ones, whereas using visible carcass lesions as a marker of infection was less reliable.

MATERIALS AND METHODS

Examination of found-dead badgers. The public was asked to notify the State Veterinary Service (now Animal Health) when they saw a dead badger. Badger carcasses were collected and transported to local Veterinary Laboratories Agency (VLA) regional laboratories, usually within 24 h of collection, by animal health officers of the State Veterinary Service. Badgers were deemed unsuitable for postmortem examination if the carcass was not intact, if the carcass was distended with gas, if there was severe myiasis, or if the carcass was flattened. Carcasses were refrigerated at VLA between 2 and 8°C and examined as soon as possible after receipt and always within 72 h. Postmortem examination was conducted using a standard protocol as previously described (17). Relevant to this study, data were also collected on animal sex, weight, and tooth wear, the latter two measures being used to approximate the ages of the badgers. Cubs were considered to have tooth wear between 0 and 25% (16) and to weigh less than 8 kg (based on unpublished analyses of other badgers). The severity of lesions in each tissue examined was scored subjectively from 0 to 4, with 0 indicating no visible lesions. For confirmation of bTB, an attempt was made to collect a standard sample of one-half of each of the retropharyngeal and bronchioles, and the root of half of the mediastinal lymph nodes if available (half of the hepatic lymph node was also collected after 15 March 2006) plus any other lesions suggestive of bTB within 15 ml of 1% cetylpyridinium chloride (CPC). Any lesions suggestive of bTB available after this sampling were preserved in 10% buffered formaldehyde solution. Bite wounds were divided between a separate container of CPC and 10% buffered formaldehyde solution. The standard sample and any bite wounds were cultured for mycobacteria, usually the next day, using the methods of the standard protocol (17). Where possible, a blood sample was collected for the lateral flow immunoassay, either by using a syringe and 14-gauge needle from the thorax via the chest wall before the carcass was opened or, if this was not possible, during the necropsy. The blood sample was collected from the heart or free from within the thorax. An assessment of the condition of the blood was made on a scale of 1 to 4, with 1 being fresh with little hemolysis and 4 being severely hemolyzed. Blood samples were tested for the presence of specific antibody as previously described (21), immediately after collection. Results were read at 20 min after adding sample buffer. Any visible band in the test area of the BrockTB Stat-Pak, in addition to the control line, was considered an antibody-positive result, whereas no band in the test area in addition to the visible control line was considered a negative result.

Data analyses. Diagnostic performance was evaluated against bTB positivity (determined by culture and/or histology). The test sensitivity, specificity, and positive predictive value (PPV) were calculated using GraphPad InStat (version 3.06 for Windows; GraphPad Software, San Diego, CA) and are reported with 95% confidence interval (CI). Tests of significance between proportions (Fisher’s exact test) or between medians (Mann-Whitney test) and the calculation of odds ratio were performed using the same software.

RESULTS AND DISCUSSION

Confirmation of bTB infection and its prevalence. Between 26 October 2005 and 31 May 2006, a total of 727 badgers found dead in Wales were reported, and 550 were delivered to the Regional Laboratories of the VLA. Of the 459 carcasses that were sampled, M. bovis was isolated from 55 badgers, for which spoligotype and VNTR type were determined. Stained tissue samples from 117 culture-negative badgers revealed a further 6 culture-negative badgers considered likely to be infected with M. bovis on the basis of AFB being present. However, given that these could not be confirmed by culture they were discounted from the calculations of sensitivity and specificity, etc. Mycobacterium avium was isolated from two badgers. No acid-fast organisms were seen histologically, and no M. bovis was isolated from either of these animals.

Further details of the geographical distribution and prevalence of bTB in Welsh badgers is the subject of another paper (submitted for publication). Only four distinct molecular types of bTB were seen in the 55 badgers: spoligotype 17 (with the unique VNTR type 7555*33.1), spoligotype 22 (with VNTR type 7524*33.1), and two VNTR types of spoligotype 9. One of these shares its VNTR type with spoligotype 22, and one had the unique VNTR type 7555*32.1.

BrockTB Stat-Pak test and/or lesions as diagnostic criteria. A lateral flow immunoassay based on the detection of serum antibodies to M. bovis antigens (BrockTB Stat-Pak test; Chembio Diagnostic Systems, Inc.) (14) was evaluated for the first time using unseparated thoracic blood samples obtained from animals found dead. The test was originally designed for use with serum obtained from live animals, where it was demonstrated to have a specificity of 51% (95% CI, 46 to 55%) and a sensitivity of 93% (95% CI, 91 to 95%) following validation on 1,532 badgers against culture postmortem (20). Recently, the test has been shown to also work with unclotted (freshly obtained) whole blood or plasma or even diaphragm extract from white-tailed deer experimentally inoculated with M. bovis (20). In this study, 379 badgers of the total 459 were examined by using the BrockTB Stat-Pak (37 samples from infected badgers, 337 samples from noninfected badgers, and 5 of the 6 badgers for which only histological evidence of infection was obtained). In some cases it was not possible to obtain blood from the carcass, and testing by BrockTB Stat-Pak was not carried out at the end of the study due to test kits being temporarily unavailable.

The ratio of males to females submitted to postmortem examination was 3.2:1 and was no different for the subset tested with the Stat-Pak (P = 0.57, Fisher’s exact test). For all badgers examined, the median weight was 9.6 kg, the median body length was 68 cm, and the median tooth wear was 0.5. The subset of animals tested with the Stat-Pak was representative of the whole in each respect (P > 0.5, Mann-Whitney Test). Based on the weight and tooth wear data to approximate the age of the badgers, there were 164 cubs (36%) and 295 adults (64%) examined. These proportions were the same for the subset tested by Stat-Pak (P = 0.79, Fisher’s exact test).

Blood could be obtained from 226 animals (60%) through the chest wall before the carcass was opened. In a further 152 cases (40%) this was not possible. Reasons for this were either that the needle became blocked by a blood clot or thoracic viscera that had become macerated through trauma or that there was only a small amount of blood in the heart/thorax. In these cases the sample was taken during the necropsy. Details of the procedure used were not recorded in one case. Blood quality was significantly poorer (P < 0.0001, Mann-Whitney test) if it had to be obtained during necropsy. The median
The median score for the condition of the blood from all samples was 2 (range, 1 to 4, indicating best to worst conditions). As we have previously found the hemolysis of badger serum to significantly reduce the sensitivity of the BrockTB Stat-Pak test result with bTB being detected in more carcasses (7). However, although culture may miss *M. bovis* infection in some cases, it would be difficult to replace this gold standard method with either the BrockTB Stat-Pak test or postmortem examination alone.

Despite the relatively low sensitivity, the close association of the BrockTB Stat-Pak test result with bTB status was highly significant (*P* < 0.0001, Fisher’s exact test), and the proportion of seropositive animals that were infected divided by the proportion of test-negative animals that were infected (relative risk) was 11, compared with 3 for the application of visible lesions (Table 1). Calculation of the odds ratio for the BrockTB Stat-Pak test was 45 (95% CI, 14 to 149), which means bTB positivity was 45 times more likely to be confirmed in antibody-reactive animals than in the seronegative population. This is in contrast to an odds ratio of 3 (95% CI, 2 to 6) for the presence of visible lesions indicating bTB infection.

The PPV of the BrockTB Stat-Pak test was 76% (95% CI, 50 to 93%) and represented the percentage of badgers with a positive test that were confirmed by culture to have bTB at the prevalence of bTB represented by the 374 test samples (37/374; 10%). The bTB prevalence calculated from all animals submitted to culture was 12% (55/459), so the subset used for BrockTB Stat-Pak testing was representative of the whole. The broad confidence intervals for the estimated test sensitivity and PPV reflect the fact that only 17 seropositive results were obtained from the analytical sample used from the analysis as having ambiguous disease status.

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TABLE 1. Association between infection in badgers and result of blood test or presence of lesions

<table>
<thead>
<tr>
<th>Test (no. tested)</th>
<th>Infected* badgers (55 examined)</th>
<th>Uninfected badgers (398 examined)</th>
<th>Relative risk</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. negative</td>
<td>No. positive</td>
<td>Sensitivity (%)</td>
<td>No. positive</td>
</tr>
<tr>
<td>Brock TB Stat-Pak (374)</td>
<td>24</td>
<td>13</td>
<td>35</td>
<td>4</td>
</tr>
<tr>
<td>Presence of any lesion (453)</td>
<td>34</td>
<td>21</td>
<td>38</td>
<td>65</td>
</tr>
</tbody>
</table>

* As determined by positive culture confirmed by spoligotyping/VNTR (*n* = 55) only. Six badgers with AFB present histologically but from which no positive culture was obtained were discounted from the analysis as having ambiguous disease status.

b Proportion of test-positive animals that were infected divided by the proportion of test-negative animals that were infected. (mean, 15 sites). In some badgers it was not possible to examine all sites due to traumatic damage to viscera or the inability to locate all lymph nodes. Using carcass lesions as a marker of infection was significantly less accurate (78%; 95% CI, 74 to 82%) than the BrockTB Stat-Pak test (93%; 95% CI, 89 to 95%; *P* < 0.0001, Fisher’s exact test), reflecting the fact that other infections can produce visible lesions reminiscent of bTB (12, 13). An example of this was evident in the current study. One badger had an enlarged/lesioned popliteal lymph node from which *M. avium* was subsequently isolated. This badger was negative in the Stat-Pak test. Taking the sum of the lesion scores in each tissue as a relatively crude indicator of the severity of infection among *M. bovis* culture-positive badgers, there was evidence that Stat-Pak positivity was associated with more severe bTB, as reported previously (2). The median score for the 13 Stat-Pak-positive badgers was 2 (range, 0 to 9), in contrast to a score of 0 (range, 0 to 11) for the 24 Stat-Pak-negative badgers. This difference was significant (*P* < 0.05, Mann-Whitney test).

The estimates of sensitivity and specificity were relative, having been measured against culture on a limited range of tissues. Gross examination of more tissues and obtaining of more cultures from each badger would result in bTB being detected in more carcasses (7). However, although culture may miss *M. bovis* infection in some cases, it would be difficult to replace this gold standard method with either the BrockTB Stat-Pak test or postmortem examination alone.

FIG. 1. The influence of blood condition, scored from 1 (fresh with little hemolysis) to 4 (severely hemolysed), on the Brock TB Stat-Pak result for badgers confirmed as infected by culture only.
obtained from a total of 379 badgers tested. Based on these results, over three-quarters of BrockTB Stat-Pak test-reactive badgers didn’t have been submitted to detailed postmortem examination and culture in order to confirm their bTB status. However, the PPV is dependent on both the properties of the test and the population studied, because the lower the disease prevalence, the lower the predictive value of the diagnostic test. Also, failure to attempt culture from a badger would by definition mean a loss of molecular typing data, which may be undesirable.

The ability of the BrockTB Stat-Pak test to detect bTB in badgers and wild deer (20) using samples that we were able to obtain from dead animals suggests that this may be a useful approach generally applicable to other wildlife species. Given its high specificity, low cost, ease of operation, and rapidity, the BrockTB Stat-Pak test could be used to reduce the number of badger carcasses submitted for examination, especially in situations where resources are limited, a prompt management decision is needed, and/or investigation by the gold standard method of culture is not possible. Accordingly, a positive BrockTB Stat-Pak test result would be accepted as a true positive, while only animals negative by the test would be submitted to more detailed postmortem examination and culture.

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