Evaluation of adenosine deaminase seric activity in the diagnosis of bovine tuberculosis

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Determination of seric levels of adenosine deaminase (ADA), an enzyme produced by monocytes/macrophages and lymphocytes, has been used in the diagnosis of human tuberculosis (TB). In the present study, ADA seric activity was evaluated comparatively to the comparative tuberculin test in the diagnosis of bovine tuberculosis. Two hundred fifty-six cattle were classified by origin and by the comparative tuberculin test as TB-positive animals (n = 52, from herds where the Mycobacterium bovis had previously been isolated), and TB-negative animals (n = 204, TB-free herds). The mean ADA seric value from the TB-positive group (4.45 ± 2.33 U/L) was significantly lower (p = 0.008) than that observed in sera from the TB-negative group (6.12 ± 4.47 U/L). When animals from a herd with clinical cases of enzootic bovine leukosis of TB-negative group were withdrawn from analysis, the mean ADA seric values of TB-negative group (5.12 ± 3.75 U/L) was not significantly different anymore from that of the TB-positive group (p = 0.28). There was no agreement in the diagnosis of bovine TB between comparative tuberculin test and determination of ADA seric values, using two different cutoff points, being 6.12 U/L and 15.0 U/L, (kappa = –0.086 and kappa = –0.082, respectively). In conclusion, the determination of ADA seric activity was not a good auxiliary test for bovine TB, because it was not able to distinguish between TB-positive and TB-negative animals.

Key words: bovine tuberculosis - cattle - diagnosis - adenosine deaminase

Bovine tuberculosis (TB) is an important zoonotic disease in cattle caused by Mycobacterium bovis and is more frequent in developing than developed countries (Cosivi et al. 1998, Mota & Lobato 1998). Although the disease has been considered under control in many developed countries, its reemergence has been observed during the last couple of years (Collins et al. 1994). In Brazil, the mean prevalence of last couple of years (Collins et al. 1994). In Brazil, the mean prevalence of last couple of years (Collins et al. 1994). Reemergence has been observed during the last couple of years (Collins et al. 1994). The method recommended by the World Animal Health Organization (OIE) (Palmer 2004) and worldwide accepted for the diagnosis of bovine TB is the tuberculin test (Langenegger et al. 1981, Monaghan et al. 1994, O’Reilly 1995, Palmer 2004). However, because tuberculin testing is not very sensitive, the development of more sensitive and specific diagnostic tools could improve bovine TB control and eradication programs (Harboe 1990, Cousins et al. 1991, L’Herminez 1993, Wood & Rothel 1994, Collins 1995, Thoen & Bloom 1995).

The enzyme adenosine deaminase (adenosine amino hidrolase - E.C.3.5.4.4. - ADA) is an enzyme involved in the metabolism of purines (Koehler & Benz 1962, Ammann & Fudenberg 1980, Giusti & Galanti 1984) and is widely distributed in lymphoid tissues (Edwards et al. 1971). ADA acts in the differentiation of lymphoid cells (Barton & Goldshneider 1979, Bothamley et al. 1995), is secreted in biological fluids during the cellular immune response against intracellular pathogens (L’Herminez 1993), but can also be increased in other pathological processes (Chalhoub et al. 1996). The genetic deficiency of ADA production leads to a syndrome characterized by reduction of T cell population in humans (Clark 1995).


Rodrigues et al. (1997) evaluated ADA seric activity in 150 comparative tuberculin test (CTT)-negative animals and 10 CTT-positive animals from the state of Bahia, Brazil and ADA seric activity was significantly higher in diseased cattle.

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The aim of this study was to evaluate the ADA seric activity comparatively to the comparative tuberculin test in the diagnosis of bovine TB in cattle in the state of Minas Gerais.

MATERIALS AND METHODS

Animals - Two hundred fifty-six adult Holstein-Zebu crossbred cattle from six dairy farms in Minas Gerais, Brazil, were submitted to comparative tuberculin test. Based on their origin and results of the CTT, animals were classified in two groups: (i) a TB-positive group, including 52 CTT-positive animals from three farms where the \( M. bovis \) had previously been isolated from animals with clinical signs of tuberculosis (32 animals from Herd A, 2 from Herd B, and 18 from Herd C), (ii) TB-negative group, including 204 CTT-negative animals from three tuberculosis-free farms (5 animals from Herd D, 104 from Herd E, and 95 from Herd F).

Serum samples - Blood samples were collected through jugular venopunction and sera were prepared and stored at \(-20^\circ\text{C}\) until tested. The time between blood collection and serum storage was always shorter than 20 h. Only hemolysis-free sera were used because ADA activity from erythrocytes has been shown to increase ADA seric values (Giusti & Galanti 1984).

CTT - The CTT was performed according to Langenegger et al. (1981) using bovine and avian PPD (Laboratório Regional de Apoio Animal, Ministério da Agricultura, Pecuária e Abastecimento, Pedro Leopoldo, MG Brazil).

ADA seric activity - ADA seric activity was determined by ADA-Kit (Biokits, Contagem, MG, Brazil), according to the manufacturer, whose methodology is based on Giusti and Galanti (1984), modified by Rodrigues et al. (1993).

Statistical analysis - Mean values of ADA seric activity from TB-positive and TB-negative cattle were compared by Student’s t-test after logarithmic transformation \([\log_{10}(\text{ADA}+1)]\) (Snedecor & Cochran 1980). Pearson correlation coefficient was computed for data on ADA seric activity and comparative tuberculin test (Snedecor & Cochran 1980). The frequency and dispersion of ADA seric activity values between TB-positive and TB-negative groups were evaluated graphically. The agreement between ADA seric values and CTT results were evaluated by the kappa statistic (Smith 1995) and Fisher exact test (Snedecor & Cochran 1980). The statistical analysis were performed by Epi Info (Dean et al. 1995), Minitab (Minitab for Windows, release 9.2, Minitab Inc., US) and WinEpiscope (Frankena et al. 1990) softwares.

RESULTS

The minimum and maximum ADA seric values (U/L), the mean values and standard deviation from TB-positive and TB-negative animals, as well as the global values for the two groups, are shown in Table I.

The mean ADA seric values for TB-positive animals was significantly lower \((P = 0.008)\) than that observed for TB-negative animals. Also, the highest ADA seric values were found in TB-negative animals from two herds. When animals from Herd E, which had the largest percentage (16%) of animals with ADA seric values over 11.22 U/L (the highest ADA seric values found among TB-positive animals) were withdrawn from analysis, the mean ADA seric values between TB-negative and TB-positive cattle did not differ significantly anymore \((P = 0.28)\).

The dispersion and frequency of ADA seric values between TB-positive and TB-negative groups are shown in Figs 1 and 2, respectively.

To study the agreement between the determination of ADA seric values and the CTT in the diagnosis of bovine TB, two cutoff points were established. One was based

![Fig. 1: adenosine deaminase (ADA) seric values (U/L) in sera from animals found positive (52) and negative (204) by the comparative tuberculin test.](image)

**TABLE I**

Values of adenosine deaminase (ADA) seric values (U/L) in sera from tuberculosis (TB)-positive and TB-negative cattle by the comparative tuberculin test

<table>
<thead>
<tr>
<th>Comparative tuberculin test</th>
<th>ADA seric values (U/L)</th>
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<tr>
<td></td>
<td>Mean ± SD</td>
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<tr>
<td>Positive ((n = 52))</td>
<td>4.45 (^{c/d}) ± 2.33</td>
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<tr>
<td>Negative (^a) ((n = 204))</td>
<td>6.12 (^c) ± 4.47</td>
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<tr>
<td>Negative (^b) ((n = 100))</td>
<td>5.12 (^d) ± 3.75</td>
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<td>Total ((n = 256))</td>
<td>5.78 ± 4.24</td>
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\(^a\): including animals from Herd E; \(^b\): excluding animals from Herd E; \(^c\): \(P = 0.008\); \(^d\): \(P = 0.28\)
on the mean ADA seric value of 6.12 U/L from animals of the TB-negative group (Table II) and the other on the mean ADA seric value of 15.00 U/L observed in ten TB-free animals from the state of Bahia, Brazil (Rodrigues et al. 1997). The kappa values between ADA seric value determination and the CTT were 0.086 and –0.082 using the 6.12 U/L and the 15.00 U/L cutoff points, respectively (Tables II, III). The Fisher exact test did not demonstrate any association between ADA seric activity and the CTT (P > 0.05).

The Pearson correlation coefficient between the ADA seric activity and the CTT was low (r = 0.129).

**DISCUSSION**

The increase of ADA seric activity in biological fluids has been used, with good sensitivity and specificity, as an auxiliary diagnostic tool in human TB, mainly for extrapulmonary TB as in patients with pleural effusions, pericardial effusions, meningitis, and ascitis (L’Herminez 1993, Pérez-Rodriguez & Castro 2000, Golden & Vikram 2005), in areas where the prevalence of TB effusions exceeds other effusion causes (Burgess et al. 1995, Pérez-Rodriguez & Castro 2000, Golden & Vikram 2005).

The mean seric value of ADA observed among TNegative animals in the present study was lower than that reported by Rodrigues et al. (1997), but similar to that observed by Yasuda et al. (1996). These differences can be related to the presence of other pathogens or uncontrolled pathologies, as enzootic bovine leukosis, as has been described by Yasuda et al. (1996). Surprisingly, the mean ADA seric value observed here was higher in the TB-negative than in the TB-positive group, and this was mainly due to the high frequency (16.34%) of animals from Herd E that showed ADA seric values over 11.22 U/L, similar to the highest value found among TB-positive animals. When cattle from Herd E were removed from the statistical analysis, no significant difference was found between the two groups. The high ADA seric values

<table>
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<th>Table II</th>
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<tr>
<td>Agreement of adenosine deaminase (sADA) values, with a cutoff of 6.12 U/L, and comparative tuberculin test in the diagnosis of bovine tuberculosis</td>
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<tr>
<td>sADA</td>
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<td>------</td>
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<tr>
<td>Positive</td>
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<tr>
<td>Negative</td>
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<tr>
<td>Total</td>
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<tr>
<td>a: Kappa = –0.086; b: animals positive to the comparative tuberculin test; c: animals negative to the comparative tuberculin test; d: animals positive to ADA (≥ 6.12 U/L); e: animals negative to ADA (&lt; 6.12 U/L); P &gt; 0.05.</td>
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<th>Table III</th>
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<td>Agreement of adenosine deaminase (sADA) values, with a cutoff of 15.00 U/L, and comparative tuberculin test in the diagnosis of bovine tuberculosis</td>
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<td>sADA</td>
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<td>Positive</td>
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<tr>
<td>a: Kappa = –0.082; b: animals positive to the comparative tuberculin test; c: animals negative to the comparative tuberculin test; d: animals positive to ADA (≥ 15.0 U/L); e: animals negative to ADA (&lt; 15.0 U/L); P &gt; 0.05.</td>
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found in animals from Herd E could be due to infection by agents other *M. bovis*, such as the bovine leukemia virus (BLV). Indeed, clinical signs of enzootic bovine leukosis, confirmed by histopathology, were detected in some animals from Herd E. According to Yasuda et al. (1996), animals with clinical signs of enzootic bovine leukosis have high levels of ADA seric activity.

Although elevated ADA seric levels were reported from human TB patients (Lakshimi et al. 1992, Rodrigues et al. 1993, Orphanidou et al. 1996), measurement of this enzymatic activity had medium to low sensitivity compared to ADA determination in pleural fluid (Lakshimi et al. 1992, Orphanidou et al. 1996). Furthermore, Orphanidou et al. (1996) could not find any correlation between ADA activity in serum and in pleura. In the present study, the lack of a significant difference in ADA seric values between TB-positive and TB-negative cattle could be due to the difference between the production of ADA isoenzymes during immune responses in unlike species to diverse stimuli and to the clinical status of TB-positive animals.

Cattle infected by BLV showed higher ADA seric values when clinical signs of disease were present (Yasuda et al. 1996). In the case of humans infected with *M. tuberculosis*, ADA seric levels were also higher in patients with positive acid-fast bacilli smear and strong skin test reaction (Lakshimi et al. 1992). Therefore, for human TB, ADA activity is generally employed as an auxiliary diagnostic tool for confirmation of clinical suspicion and correct therapeutic approach (Chalhoub et al. 1996).

The majority of the CTT-positive animals studied here had no clinical signs of TB and this could also partly explain the ADA seric concentration found in this group. Rodrigues et al. (1997) reported ADA seric values higher than 15 U/L in ten animals with TB but did not describe the clinical status of the animals; more advanced disease in these animals could explain the higher ADA seric values in that study when compared to the present. Even if the determination of ADA seric activity could distinguish animals with signs of TB from the rest of the herd, this would have a restricted use, because just the culling of animals with clinical signs would not eradicate the infection from the herd (Mota & Lobato 1998).

The comparative tuberculin test, a type IV hypersensitivity reaction, is a cell-mediated response to mycobacterial antigens (Monaghan et al. 1994). Unger et al. (1992, 1994) showed that the ADA activity, especially the ADA2 isoenzyme, can also be a useful marker of cell-mediated immunity against *M. tuberculosis* in human TB. Upon regression analysis of ADA seric values and values of skin thickness increasing from CTT, we could not demonstrate an association between these two variables. This suggests that ADA seric activity in cattle does not correspond to a specific cell-mediated response against *M. bovis*, probably because the ADA2 isoenzyme, the major responsible for ADA activity in human pleural effusions (Unger et al. 1992; 1994), seems to be absent from cattle sera (Tanabe 1993).

The dispersion of ADA seric values between TB-positive and TB-negative cattle in the present study showed a unimodal distribution, i.e., one population curve is inside the other (Figs 1, 2), making it almost impossible to establish a cutoff to separate the two populations and develop an ADA based test with any diagnostic or epidemiologic meaning. In parallel, analysis of ADA seric values and CTT, measured by the kappa statistics, using the two cutoff points showed a complete lack of agreement between the two tests.

The use of ADA seric activity as a diagnostic tool for bovine TB becomes even more unlikely because, similar to what happens in man (Chalhoub et al. 1996), other pathologies such as the clinical presentation of bovine enzootic leukosis could also increase ADA seric activity (Yasuda et al. 1996); the latter disease is widespread in Brazil (Leite et al. 2001).

In conclusion, due to the impossibility of establishing a meaningful cutoff of ADA seric values for separation of TB-positive and TB-negative animals, as compared to the CTT, this test should not be used as an auxiliary test in the diagnosis of bovine TB.

REFERENCES


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