Detection of Surra (trypanosomiasis) positivity in humans in an outbreak area of Indonesia

Dyah Haryuningtyas Sawitri,1 April Hari Wardhana,1 Mohamad Sadikin,2 Heri Wibowo3

ABSTRACT

BACKGROUND Surra is an infection caused by a blood protozoan parasite, Trypanosoma evansi, and transmitted by blood-sucking insects. The parasite generally infects only animals; however, it was reported to infect an Indian cattle farmer in 2004, followed by reports of other human cases. The most severe Surra outbreak in Indonesia occurred in Sumba Island during 2010–2012, resulting in the death of more than 2,000 livestock. This study was conducted to explore the serological status of farmers who have intensive contact with their livestock against T. evansi infection in Southwest Sumba district.

METHODS A total of 24 serum samples were collected from farmers living in the Surra outbreak area. All sera were tested using both card agglutination test for trypanosomiasis/T. evansi (CATT/T. evansi) and field enzyme-linked immunosorbent assay (FELISA).

RESULTS Of the 24 serum samples, 4 (16.7%) samples were seropositive for the antigen T. evansi using both tests. This is the first report of human trypanosomiasis (Surra) in Indonesia. Unfortunately, the clinical manifestations of farmers with positive Surra infection were not reported because all sera samples used in this study were obtained from the Public Health Service with no reports of clinical signs from the respondents.

CONCLUSIONS Farmers living in the Surra outbreak area have a high risk of being infected with T. evansi due to their potential frequent contact with Surra vectors. Therefore, T. evansi infection in humans requires attention as it might have the potential to develop as a new emerging zoonotic disease in Indonesia.

KEYWORDS Trypanosoma evansi, human serum, CATT/T. evansi, FELISA

Trypanosomes are blood protozoa that infect humans, livestock, and wild animals throughout the world and are most often transmitted by blood-sucking insects.1 Human trypanosomiasis is endemic in Africa and South America.2 African trypanosomiasis in humans is caused by Trypanosoma brucei gambiens (chronic form) or T. b. rhodei (acute form), which is the cause of sleeping sickness or human-animal trypanosomiasis, whereas South American trypanosomiasis (Chagas diseases) is caused by T. cruzi. Sleeping sickness and Chagas diseases are transmitted by Tsetse flies and reduviid bugs/Triatominae, respectively.3 In addition to these human trypanosomes, there are various other species that cause animal trypanosomiasis with a wide geographic distribution. T. evansi is a pathogenic trypanosome that causes Surra in livestock in Asia, Africa, South America, and parts of Europe and is transmitted by hematophagous flies (Tabanids and Stomoxys calcitrans).3 Another species, T. vivax, causes nagana disease in animals in Africa and is transmitted by Glossina morsitans. Furthermore, T. congolense and T. brucei brucei have been reported to cause trypanosomiasis in animals in Africa. In addition, T. equiperdum was reported to infect horses.
in Europe, America, North Africa, and India. Another non-pathogenic parasite, T. lewisi, has been reported to infect rodents and transmitted by fleas.

Human infection by animal species of Trypanosoma is considered to be impossible because of the presence of a trypanolytic factor in human serum. Nevertheless, the first case of human infection with T. evansi was reported in the central part of India (Seoni village, Taluka Sindevali, district of Chandrapur, Maharashtra State) in 2004. The disease was found in a cattle farmer who had intensive contact with his livestock. It was reported that the disease could have occurred as the farmer was deficient in apolipoprotein L1 (APOL1), a component of human serum with trypanocidal activity. In addition, three other cases have been reported worldwide (Table 1).

Human innate immunity against T. evansi is induced by the trypanolytic activity of a human-specific apolipoprotein bound to high-density lipoproteins, termed as APOL1. APOL1 is taken up by the parasite via endocytosis and triggers the formation of anion-selective pores in the lysosomal membrane, which induces uncontrolled osmotic swelling of this compartment and subsequent cell death. The lack of nonspecific immune protection of APOL1 in humans or the development of new capacity in the parasite to respond to nonspecific immunity may also lead to the evolution of most of the trypanosome pathogens. Similarly, the results of another study reported by Lai et al demonstrated tolerance toward T. evansi stock in response to normal human serum (NHS). This report was supported by Van Vinh Chau et al who described the first human case of T. evansi infection in Asia without any APOL1 deficiency (healthy individual).

The number of atypical human infections caused by T. evansi has been increasing, due to which some scientists have created a new network on atypical human infection by animal trypanosomes (NAHIAT). This network is intended to coordinate information and research activities regarding atypical human trypanosomiasis. Indonesia is an endemic country for Surra that infects livestock, and there have been few outbreaks in 2010–2016. However, there is a lack of a study on atypical human trypanosomiasis in Indonesia. Therefore, the primary aim of the present study was to assess the details regarding the disease, particularly when there were Surra outbreaks. One of the methods used for diagnosing Surra is the serological test. Various serological tests such as indirect immunofluorescent antibody test, enzyme-linked immunosorbent assay (ELISA), and card agglutination test for trypanosomiasis (CATT) are used to identify specific antibody responses. CATT/T. evansi is highly specific in detecting circulating antibodies as it targets the variant surface glycoprotein RoTat 1.2. The CATT/T. evansi technique for detecting antibody was originally described by Bajyana Songa and Hamers and converted into a test kit by the Institute of Tropical Medicine, Belgium. This kit has a high measure of validity, and it is economical and applicable in large-scale screening. Furthermore, CATT/T. evansi is a rapid and easy test that can be performed under both laboratory and field conditions. High test accuracy has been reported for CATT/T. evansi in livestock raised in various regions such as Punjab, India, Mesir, Indonesia, and Kenya.

Field ELISA (FELISA) is an immunostick assay technique based on ELISA and developed as an alternative serological method with high accuracy. This diagnostic kit was constructed by the Indonesian Research Center for Veterinary Science, Bogor, and designed for rapid and practical application in the field to detect antibodies in the serum of infected animals. Sumba Island was the only region to be free of T. evansi infection until 2009. However, the Surra outbreak that occurred in 2010–2012 resulted in the

<table>
<thead>
<tr>
<th>Country</th>
<th>Occupation (number of farmers)</th>
<th>Year</th>
<th>Parasite identification methods</th>
<th>Clinical symptom</th>
<th>Treatment</th>
<th>Result</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>India, Seoni</td>
<td>Farmer (1)</td>
<td>2004</td>
<td>Morphology, serology, PCR</td>
<td>Fever</td>
<td>Suramin</td>
<td>Recovered</td>
<td>5</td>
</tr>
<tr>
<td>India, Kolkata</td>
<td>Farmer (1)</td>
<td>2005</td>
<td>Morphology</td>
<td>Fever</td>
<td>-</td>
<td>Death</td>
<td>8</td>
</tr>
<tr>
<td>Mesir</td>
<td>Farmer (1)</td>
<td>2010</td>
<td>Morphology</td>
<td>Fever</td>
<td>-</td>
<td>Recovered</td>
<td>9</td>
</tr>
<tr>
<td>Vietnam</td>
<td>Factory worker (1)</td>
<td>2015</td>
<td>Morphology, serology, PCR</td>
<td>Fever</td>
<td>Suramin</td>
<td>Recovered</td>
<td>10</td>
</tr>
</tbody>
</table>
death of more than 2,000 livestock (buffaloes, cattle, and mostly horses) due to *T. evansi* infection.\(^{21}\) It was reported to be the most severe Surra outbreak in Indonesia, and the parasite was believed to be introduced from Sumbawa Island (Surra endemic area) when there was traditional horse racing. Considering Surra cases in humans and to support the NAHIAT agenda, this study was performed to conduct serological tests using both CATT/*T. evansi* and FELISA kits on farmers who have intensive contact with their livestock in some Sumba outbreak regions.

### METHODS

#### Serum samples

This study was approved by the Ethics Committee of the Faculty of Medicine, Universitas Indonesia, number 809/UN2.F1/ETHICS/2015. A total of 24 sera samples (22 males and 2 females) were collected from farmers living in three subdistricts (West Wewewa, Kodi Bangedo, and Loura) of Southwest Sumba district in 2012 during an outbreak of Surra. Furthermore, 8 serum samples were obtained from Jakarta (Surra non-endemic areas) that served as seronegative controls (Table 2). Jakarta was selected for collecting seronegative samples because there has been no report of Surra cases in Jakarta infecting either livestock or humans. The seronegative control samples were selected on the basis of few parameters, i.e., the individuals were healthy, they never had contact with livestock, they lived in areas with no farms (town), and there were no vectors of Surra found in the area where they lived. Based on these parameters, all the available samples were found to be collected from female individuals. There was no difference in response to the serological test for Surra infection between males and females. All sera were maintained at -20°C for further analysis.

### Table 2. Sera samples of farmers in Southwest Sumba district used in the present study

<table>
<thead>
<tr>
<th>Status area</th>
<th>Name of community health center (Puskesmas)</th>
<th>Location (Subdistrict)</th>
<th>Sex, n</th>
<th>Range of age (years old)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Outbreak</td>
<td>Watu Kuwala</td>
<td>West Wewena</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>Outbreak</td>
<td>Wallandimu</td>
<td>Kodi Bangedo</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>Outbreak</td>
<td>Rada Mata</td>
<td>Loura</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>Outbreak</td>
<td>Waimangura</td>
<td>West Wewena</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Non-endemic</td>
<td></td>
<td>Jakarta (City)</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td>22</td>
<td>10</td>
</tr>
</tbody>
</table>

#### FELISA

Each sample was diluted in phosphate-buffered saline (PBS, pH 7.2) containing 0.05% Tween 20 (1:10) and shaken homogeneously. Subsequently, 800 µl diluted serum sample was poured into an immunostick tube, incubated at room temperature for 10 min, and then removed and washed with tap water. Each immunostick was introduced and immersed in tubes containing conjugated (1:6000) anti-human Ig G-HRP (Sigma Chem, USA), incubated at room temperature for 10 min, and then washed with tap water. Finally, the immunostick was submerged in a tube containing the substrate ODN (o-Dianisidine; Sigma Chem, USA) for 3 min and then washed with tap water. A positive response was marked by a color change on the immunostick surface coated with antigen.\(^{20}\)

#### CATT/*T. evansi*

Approximately 45 µl of the antigen was poured into the circular reaction zones of the supplied plastic card and mixed with 25 µl of the test sera according to the manufacturer’s instructions. The dropper was held vertically, and drops were allowed to fall freely without touching the card to maintain a constant volume. The antigen and serum mixture was mixed and spread properly using a clean plastic stirring rod to approximately 1 mm from the rim of the test area; the stirring rod was wiped with a sterile filter paper after each use. Each plastic card contained two reaction zones for the positive and negative controls. The card was agitated for 5 min at 70g after positioning on a flat bed of an electric orbital rotator. After 5 min, the results were read before removing the card from the rotator. A reaction was considered to be positive when the agglutination was visible to the naked eye.\(^{22}\) Human sera that were positive for anti-trypanosome antibodies by CATT demonstrated different degrees
of agglutination graded as +++ (very highly positive), ++ (highly positive), + (moderately positive), ± (mildly positive), and − (negative).

**RESULTS**

All samples from Southwest Sumba district were collected from farmers who had intensive contact with livestock to identify whether they had an opportunity to have contact with any vectors of Surra, such as the tabanid fly, the stable fly (*S. calcitrans*), or other hematophagous flies. When the vectors bite the farmers, they might show a positive antibody to *T. evansi* in their serum sample. Conversely, all samples from Jakarta were collected from those people who never had any contact with livestock and vectors, and hence they were used as the seronegative control group. Unfortunately, there were no reports of any clinical manifestations of farmers who were positive for Surra because all serum samples used in this study were obtained from the Public Health Service without the details of any clinical signs from the respondents. Based on the results of both FELISA and CATT/*T. evansi*, 4 of 24 farmer serum samples (16.67%) collected from Southwest Sumba district were confirmed to be seropositive (moderately positive), whereas all human serum samples collected from the non-endemic area (Jakarta) showed a seronegative response (Table 3). A positive CATT/*T. evansi* response was shown by the sand formation in sample numbers 1 and 5 (Figure 1), indicating that there was agglutination between the antigen CATT/*T. evansi* and anti-*T. evansi* antibodies from the farmers’ sera. Meanwhile, a positive FELISA response was shown by the color change from white to an orange dot on the immunostick. Nevertheless, these experiments were not followed by examination of blood samples to confirm the presence of parasites in the farmers’ peripheral blood samples based on either microhematocrit centrifuge technique or polymerase chain reaction (PCR). Furthermore, no clinical symptoms were reported from all respondents who showed either seropositive or seronegative responses to *T. evansi*.

**DISCUSSION**

This study was conducted to assess whether farmers were exposed to *T. evansi* infection during the outbreak in 2010–2012. The results demonstrated that a few farmers (16.7%) living in Southwest Sumba district were serologically positive for Surra tests. Another Surra surveillance study was also conducted by Wardhana et al. who collected serum samples from livestock from the same area in the same year (Southwest Sumba). They reported that 38% (57/150) of livestock were seropositive on the basis of CATT/*T. evansi*.

**Table 3.** Results of serological tests of farmers’ sera based on CATT/*T. evansi* and FELISA analyses

<table>
<thead>
<tr>
<th>Location (village, subdistrict)</th>
<th>CATT/<em>T. evansi</em></th>
<th>FELISA</th>
<th>Sex</th>
<th>Age (years old)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>+</td>
<td>−</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>Watu Kawula, West Wewewa</td>
<td>1</td>
<td>5</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Walla-Ndimu, Kodi Bangedo</td>
<td>−</td>
<td>7</td>
<td>−</td>
<td>7</td>
</tr>
<tr>
<td>Lete Konda, Loura</td>
<td>2</td>
<td>4</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Waimangura, West Wewewa</td>
<td>1</td>
<td>5</td>
<td>1</td>
<td>5</td>
</tr>
<tr>
<td>Jakarta</td>
<td>0</td>
<td>8</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td>Total</td>
<td>4</td>
<td>28</td>
<td>4</td>
<td>28</td>
</tr>
</tbody>
</table>

CATT=card agglutination test for trypanosomiasis; FELISA=field enzyme-linked immunosorbent assay
evansi results. This indicates the presence of active infection of T. evansi in the area, where the parasite might attack both humans and livestock.

Shegokar et al. mentioned that an epidemiologic surveillance study is required to follow up the seropositive findings of people living in a Surra outbreak area. They conducted whole blood analysis and demonstrated that 410 of 1,806 (22.7%) people living in a human Surra outbreak area were seropositive to T. evansi. However, no parasites were detected in the blood of 60 persons who were positive at a high serum dilution (≥1:4). The authors suggested investigation of trypanosomiasis in any human Surra outbreak region where T. evansi is endemic in livestock, such as in Indonesia. Another study also reported that 1 of 30 (3.3%) workers raising camel in Egypt was positive on the basis of ELISA antibodies and blood smear analyses. Although some islands in Indonesia are known to be Surra endemic areas, there are no data reporting a serological study of the disease in humans who live and have contacts with livestock. Therefore, a serological survey of Surra in humans was conducted in Sumba Island when the outbreak occurred in 2010–2012.

The present study focused on farmers exposed to T. evansi who were in contact with livestock. Serological surveillance of anti-T. evansi antibodies was conducted in farmers living in three subdistricts of Southwest Sumba. The majority of the people living in the villages consists of traditional farmers and raise livestock. They have intensive contact with their livestock every day, which puts them at a high risk, particularly if the livestock is infected by infectious agents such as T. evansi.

Previous studies have also used CATT/T. evansi and FELISA to detect antibodies in the serum or plasma of farmers. The CATT/T. evansi technique has been applied in several countries for serological surveys in humans and livestock, whereas the FELISA technique was developed at our laboratory based on ELISA applied to livestock only. The antigen coated on the FELISA immunostick was obtained from an Indonesian strain of T. evansi and can detect various strains of T. evansi in Indonesia. Meanwhile, the antigen used in CATT/T. evansi is commercially available. The latter method has been generally applied to detect seropositivity of T. evansi in humans (India, Egypt, and Vietnam). In terms of the results of the serological tests, there were no differences in responses between positive samples analyzed using either CATT/T. evansi or FELISA. However, based on sera dilution of the samples, the FELISA method was found to be more sensitive than CATT/T. evansi.

At least two hypotheses can be used to explain why the CATT/T. evansi result had less sensitivity compared to that of FELISA. First, the antigen coated on the CATT/T. evansi kit might not have been isolated from the Indonesian strain of T. evansi. Second, the parasite could have been lysed in the body or the parasite did not exist in the body for a long time. The second hypothesis was consistent with that reported of Joshi et al. who evaluated the level of specific antibodies in a patient at a high plasma dilution (1:64) when he was infected with high parasitemia (approximately 10^4 trypanosomes/ml). After 24 hours or 3 months after being treated with suramin, the CATT/T. evansi result was positive at plasma dilutions of 1:16 and 1:4, respectively. The sensitivity of plasma gradually decreased after 6 months of treatment. Positivity was detected at only 1:2 sera dilution. These results indicated that the decrease in seropositivity was probably due to lysis of the parasite in the body. In the present study, the farmers’ sera were collected in 2012 when the number of Surra cases in Sumba had declined due to intensive drug treatment, surveillance, and vector elimination. This might have contributed to the low level of seropositivity result.

In general, T. evansi cannot infect humans because of the presence of the trypanolytic factor APOL1, which eliminates the parasite from the body. The case of Surra that infected the farmer in India was due to the unavailability of APOL1 in his body. During that time, all scientists believed that a defect in APOL1 was the primary factor responsible for atypical human trypanosomiasis infection. However, this theory became controversial after a case of Surra infection reported in an individual in Vietnam. In that case, the Vietnamese lady had normal serum with APOL1 and did not suffer from immunosuppression, but she was infected with T. evansi and exhibited long intermittent fever as a clinical symptom. Unfortunately, this study was not followed by APOL1 analysis, due to which we are not aware whether the farmers who were seropositive were deficient in APOL1. Surra infections in farmers have been reported to be caused due to their frequent contact with vectors of Surra. Ekawasti et al. reported the presence of a highly abundant population of hematophagous flies in Sumba, which increased the
probability of transmission of *T. evansi* to the farmers.

As far as the authors know, the present study using serological tests in humans living in Sumba Island is the first report of seropositivity to *T. evansi* in Indonesia. This finding is extremely important to demonstrate that Surra might be a new potentially emerging disease that exposes people in an endemic area to a high risk of infection. In addition, this finding has implication as an early warning to people who have contact with livestock. It is highly recommended to pay attention when seropositive Surra infection is found in humans, particularly in an endemic area.

According to Brun, there are at least two scenarios through which humans can be infected with *T. evansi*. First, the parasite is mutated to a form that can resist the lytic factor in the human plasma. Due to mutation, the parasite could escape from the trypanolytic activity. Lai et al reported that *T. evansi* stock isolated from China could survive in NHS. The isolate was found to be resistant after being challenged with NHS and inoculated into mice for 25 times. In the 20th passage, the *T. evansi* stock isolated from China became resistant to NHS, including the serum containing APOL1 (the trypanolytic factor). Lai et al also demonstrated that *T. evansi* stocks isolated from different geographic regions developed a high level of natural resistance to NHS. They examined 15 isolates of *T. evansi* from different stocks, i.e., China, Vietnam, Kazakhstan, and the Philippines, against NHS both in vitro and in vivo and found that 5 of 15 isolates were resistant to NHS.

Second, the human host could be deficient in the lytic factor (APOL1) in the plasma. Vanhollebeke et al demonstrated that *T. evansi* infection was caused due to frameshift mutations in both APOL1 alleles in their study patient. Two mutations within the gene encoding APOL1, in codons 142 and 266, were identified and associated with *T. evansi* susceptibility in the patient. This generated an unexpected termination of protein translation by internal stop codons, which resulted in the total absence of APOL1. The defective APOL1 in the serum of the person with *T. evansi* infection did not demonstrate lytic activity against *T. evansi*.

According to Desquesnes et al, the abnormalities found in the production mechanism of APOL1 are not the only factor that lead to the development of *T. evansi* infection in humans, but people suffering from immunosuppression are also at high risk of developing *T. evansi* infection, particularly among those living in regions where the parasite is endemic. Wabale et al reported a case of trypanosomiasis caused by *T. evansi* in a patient infected with retrovirus and lymphadenitis from Mumbai, India. The patient was immunocompromised, which may have facilitated the infection of *T. evansi*.

A recent case of human trypanosomiasis caused by *T. evansi* infection was reported from Vietnam. The parasite infected a 38-year-old woman breastfeeding a healthy infant. Parasitological, serological, and PCR analyses confirmed that the woman had Surra infection. Interestingly, the woman was a healthy person who had not previously described immunological risk factors for Trypanosoma infection and did not have the genetic mutations associated with APOL1 deficiency and serum concentrations. She became infected with the parasite through a wound while butchering raw beef. This indicates that a normal person is also at a high risk of being infected with *T. evansi*.

Although the present study had successfully demonstrated seropositivity to *T. evansi* in farmers, there are still certain limitations. Because the farmers’ sera were obtained from sera bank, there were no data about the clinical symptoms. Moreover, the presence of *T. evansi* in the farmers could not be confirmed by molecular analysis such as PCR, even for parasitological examination, because of the unavailability of blood samples. The study was also not followed by APOL1 analysis.

In conclusion, farmers who have intensive contact with livestock in Surra outbreak areas can potentially develop antibodies to *T. evansi*. This indicates that there is frequent contact between the positive vectors such as hematophagous flies carrying *T. evansi* and the farmers. Additional large-scale studies are warranted to assess the zoonotic potential of this disease by conducting joint research with relevant public health authorities and the Agricultural Ministry on *T. evansi* infection (Surra).

Conflict of Interest
The authors affirm no conflict of interest in this study.

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