**FULL PAPER**

**Parasitology**

**Therapeutic Effect of Clindamycin and Tetracycline on *Babesia rodhaini* Infection in Mouse Model**

Agus WIJAYA, Retno WULANSARI, Hitoshi ANO, Hisashi INOKUMA and Susumu MAKIMURA

Laboratory of Veterinary Internal Medicine, Department of Veterinary Science, Faculty of Agriculture, Miyazaki University, Miyazaki 889–2192, and \(^1\)Faculty of Agriculture, Yamaguchi University, Yamaguchi 753–8515, Japan

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**ABSTRACT.** In order to identify the alternative effective chemotherapeutic agents for murine babesiosis, some selected drugs were examined for their efficacy against protozoan infection in the mouse-*Babesia rodhaini* (*B. rodhaini*) model. Clindamycin was not completely effective for elimination of parasites in a dose of 50 mg or 100 mg/kg BW/day b.i.d. but effective to prolong the life span of hosts, while it completely cured *B. rodhaini* infections in a dose of 200 mg. On the other hand, a double therapy consisting of 2 treatments with 100 mg clindamycin and 100 mg clindamycin and with 100 mg clindamycin and 100 mg tetracycline; respectively, and a single therapy with 100 mg tetracycline or 200 mg clindamycin, had a possibility to clear away *B. rodhaini* organisms from hosts. However, almost all the treatment groups, had a relapse of the infection within 10 days post treatment or re-treatment. Cured mice by treatment with clindamycin and clindamycin, or clindamycin and tetracycline showed complete resistance against challenge with *B. rodhaini*, while mice cured by administration of clindamycin at 200 mg or tetracycline at 100 mg showed incomplete resistance to challenge infection. The present data suggest that the two former chemotherapies can induce effective protective immunity (premunition), but the latter two chemotherapies induce incomplete premunization.

**KEY WORDS:** Babesia rodhaini, chemotherapy, clindamycin, effective protective immunity, premunization.

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Babesiosis has always been problematic both in human and animal medicine, and a better drug has been urgently needed in recent years due to the spreading geographic range of the infection. This condition is further complicated by the occurrence of pet animal babesiosis. Nowadays, many kinds of chemotherapy and vaccination have been applied in order to eradicate babesiosis without success. This condition, leaded some workers to use clindamycin plus quinine as a current treatment of choice, which have been reported to have antimalarial activity, but to have a little effect for parasitism or survival in the experimental babesiosis model. It is well known that almost no effective drug without any side effective problem can be found until now especially for treatment of babesiosis. In this study, we used clindamycin and tetracycline in experimental therapy of murine babesiosis. Tetracycline has a relatively low of toxicity, and shows slight side effects, and is generally safety, so that it would be strongly recommended to use if its efficacy was comparable to other classes of compounds. Since 1953, chlorotetracycline was reported effective in retarding the growth of *B. equi* in splenectomized donkeys [4]. In the study of *B. microti* in the hamster model, both clindamycin and the combination of clindamycin and oral quinine are safe, effective agents for fatal babesiosis [8]. However, some workers used a combination of quinine and clindamycin for the therapy of *B. microti* infection in the hamster model without constant results in parasitological cure [6].

In attempt to demonstrate the utility of the mouse model, three antiprotozoal drugs, diminazene diaceturate, clindamycin and oxytetracycline, were examined for their efficacy inhibiting the growth of *B. canis* in canine-red blood cell—SCID mice. The mouse model clearly showed that diminazene diaceturate and oxytetracycline were capable of eliminating *B. canis* from the canine-red blood cell—SCID mice, whereas clindamycin exhibited a static effect only as parasitemia relapsed upon cessation of drug administration [1]. Actually, the purpose and strategy of babesiosis treatment is not to eliminate parasite quickly and completely from hosts, but to suppress the proliferation of parasites and to decrease the symptom of babesiosis without any bad side effect, so that pretension is established. In the present study, we use clindamycin and tetracycline as the alternative effective drug for the treatment of babesiosis and compared the effectiveness of the two drugs and diminazene diaceturate against *B. gibboni* infection in mice, and more over in acquiring resistance to challenge infection in cured mice.

**MATERIALS AND METHODS**

**Animals and parasites:** An outbred strain of male ICR mice, 8–12 weeks old with around 30 g B.W. were purchased from Clea & Shizuoka Experimental Animal Supplier Co., Japan and housed in our laboratory. The pathogenic agent used in this study was *B. rodhaini* Australian strain kindly provided by Prof. K. Ono of Dept. of Clinical Pathology, Faculty of Agriculture, Tokyo University, and was routinely maintained in our laboratory by intra peritoneal passage with parasitized erythrocytes (pRBC) through ICR mice.

**Drugs:** The drugs used were clindamycin phosphate “Darunion” (Up John Co., Tokyo), oxytetracycline “Terramycin”
Long Acting (Pfizer Co., Tokyo), and 4-4' diazaminodibeza-
midine diaceturate (diminazene diaceturate), “Ganazeq” 
(Chiba-Geigy, Co., Japan). Clindamycin was given through a 
gastric tube into a group of mice, while oxytetracycline was 
subcutaneously injected into a group of mice, and Ganazeq 
was subcutaneously administered at a dose of 25 mg/kg/day in 
saline solution for 3 days as a comparison groups. The dosage 
of the antibiotics was calculated per kg body weight on the 
basis of human therapeutic dosages. The antibiotics, in saline 
or distilled water solution, were used in the following doses: 
50, 100, and 200 mg/kg BW b.i.d. of clindamycin and 100 
mg/kg BW s.i.d. of tetracycline. The dose used for tetracy-
cline was 100 mg/kg/day, considering that dogs tolerate intra 
muscular (IM) oxytetracycline in doses of 50–100 mg/kg/day. 
Oral doses of 75–465 mg/kg daily for as long as 8 weeks were 
attained by dogs without evident toxicity [2]. The mice in the 
time control groups were given distilled water only.

Experimental procedures: The experimental mice were 
divided into four groups; the first groups were treated with 
clindamycin for seven days, while the second groups were 
injected with tetracycline for seven days, and the third groups 
were administered with diminazene diaceturate for three days 
as a comparison groups. To the fourth groups was given dis-

Table 1. Effect of treatment with selected antiprotozoal drugs on B. rodhaini infection in mice

<table>
<thead>
<tr>
<th>Treatment groups</th>
<th>1&lt;sup&gt;a&lt;/sup&gt;</th>
<th>2&lt;sup&gt;a&lt;/sup&gt;</th>
<th>3&lt;sup&gt;a&lt;/sup&gt;</th>
<th>4&lt;sup&gt;a&lt;/sup&gt;</th>
<th>5&lt;sup&gt;a&lt;/sup&gt;</th>
<th>6&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>%Parasitemia&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>day 0&lt;sup&gt;c&lt;/sup&gt;</td>
<td>1.5 ± 0.72</td>
<td>1.3 ± 0.24</td>
<td>1.4 ± 0.38</td>
<td>1.4 ± 0.38</td>
<td>0.6 ± 0.51</td>
<td>0.9 ± 0.17</td>
</tr>
<tr>
<td>day 3</td>
<td>27.4 ± 8.13</td>
<td>29.0 ± 3.24</td>
<td>29.2 ± 2.92</td>
<td>9.4 ± 1.19</td>
<td>6.7 ± 6.60</td>
<td>32.2 ± 6.62</td>
</tr>
<tr>
<td>day 4</td>
<td>18.5 ± 19.08</td>
<td>46.0 ± 12.20</td>
<td>55.0 ± 13.18</td>
<td>0.2 ± 0.19</td>
<td>0.8 ± 0.95</td>
<td>80.6 ± 13.62</td>
</tr>
<tr>
<td>day 5</td>
<td>1.0 ± 0.64</td>
<td>100.0 ± 5.00</td>
<td>NT</td>
<td>0.0 ± 0.00</td>
<td>0.1 ± 0.13</td>
<td>NT</td>
</tr>
<tr>
<td>day 7</td>
<td>0.5 ± 1.03</td>
<td>NT</td>
<td>NT</td>
<td>0.0 ± 0.00</td>
<td>NT</td>
<td>NT</td>
</tr>
<tr>
<td>day 8</td>
<td>0.0 ± 0.00</td>
<td>NT</td>
<td>NT</td>
<td>NT</td>
<td>NT</td>
<td>NT</td>
</tr>
<tr>
<td>Mortality&lt;sup&gt;b&lt;/sup&gt;</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>day 4</td>
<td>0/6</td>
<td>2/6</td>
<td>4/6</td>
<td>0/5</td>
<td>1/8</td>
<td>4/5</td>
</tr>
<tr>
<td>day 5</td>
<td>0/6</td>
<td>6/6</td>
<td>6/6</td>
<td>0/5</td>
<td>1/8</td>
<td>5/5</td>
</tr>
<tr>
<td>day 6</td>
<td>1/6</td>
<td>NT</td>
<td>NT</td>
<td>0/5</td>
<td>1/8</td>
<td>NT</td>
</tr>
<tr>
<td>day 7</td>
<td>1/6</td>
<td>NT</td>
<td>NT</td>
<td>0/5</td>
<td>1/8</td>
<td>NT</td>
</tr>
<tr>
<td>day 8</td>
<td>2/6</td>
<td>NT</td>
<td>NT</td>
<td>0/5</td>
<td>1/8</td>
<td>NT</td>
</tr>
</tbody>
</table>

a) Normal ICR mice were infected intra-peritoneal with 1 × 10<sup>6</sup> parasitized RBC. b) Group 1. Clindamycin in a dose of 200 mg/ 
kg b.i.d. × 7 days, via oral route. Group 2. Clindamycin in a dose of 100 mg/kg b.i.d. × 7 days, via oral route. Group 3. 
Clindamycin at a dose of 50 mg/kg b.i.d. × 7 days, via oral route. Group 4. Ganazeq in a dose of 25 mg/kg s.i.d. × 3 days, by S.C. 
route. Group 5. Tetracycline in a dose of 100 mg/kg s.i.d. × 7 days, by S.C. route. Group 6. Control (no treatment), supplied with 
distilled water only. Each treatment group was composed of 5, 6 or 8 mice. c) All the treated animal groups had a relapse within 
10 days post treatment. d) Results are expressed as mean % parasitemia suppressed by drugs in treated and non-treated animals.

RESULTS

Effect of treatment with selected antiprotozoal drugs against B. rodhaini infection in mice: Three selected antipro-
protozoal drugs, clindamycin, tetracycline, and diminazene 
diaceturate were used for treatment of B. rodhaini infection 
in ICR mice at various doses. The effect of those drugs is seen in 
Table 1. Parasitemia was cleared in group 1 (clindamycin in a 
dose of 200 mg/kg/day), group 4 (diminazene diaceturate), 
and group 5 (tetracycline). Parasites were cleared from the red 
blood cells in group 1 (clindamycin at a dose of 200 mg/kg/ 
day) at day 8 post treatment, while in group 5 (tetracycline), 
they disappeared earlier than in group 1, namely at day 7 post 
treatment. On the other hand, the eliminating time of B. 
rodhaini from the murine-red blood cells was earlier in group 
4 (Ganazeq) than in group 1 (clindamycin in a dose of 200 mg/ 
kg/day) and group 5 (tetracycline), in which parasites disap-
peared at day 5 post treatment. However, the mean percent suppression of parasitemia post treatment was rapidly increased with time in group 2 (clindamycin with dose of 100 mg/kg/day), group 3 (clindamycin with dose of 50 mg/kg/day) and group 6 (control), whereas in groups 1, 4 and 5 in the first time similar conditions were also seen in the initial course after treatment but finally parasitemia level peaked until day 3 and then was followed by the decrease in parasitemia or parasitic clearance in the next day. Nevertheless, the parasitemia relapsed in all of the treated animal groups within ten days post-parasitic clearance. No or little mortality was observed especially in group 4 (diminazene diaceturate), group 1 (clindamycin in a dose of 100 mg/kg/day) and group 5 (tetracycline). In group 2 (clindamycin with dose of 100 mg/kg/day), group 3 (clindamycin with dose of 50 mg/kg/day) and group 6 (control), all the animals died at day 5.

**Effect of single and double therapies using tetracycline and clindamycin drugs against B. rodhaini in mice:** Double or single therapy was applied to choose the drugs efficacious against murine babesiosis. The results are shown in Table 2. The peak level of parasitemia suppressed in the first treatment of double therapy with clindamycin & clindamycin and clindamycin & tetracycline was 19.8% and 17.5%, respectively and lower than that of the single therapy with tetracycline of 11.2%. In contrast, at day 6 post single therapy, the level of parasitemia drastically decreased until parasitemia was cleared, while in the first treatment of double therapy, parasitic clearance was reached at day 7 post treatment. At day 10 post the first treatment of double or single therapy, all of the treated groups relapsed. In the second treatment of double therapy with clindamycin and clindamycin two out of five mice relapsed and the peak level of parasitemia suppression was 78.4% at day 3 post treatment, while one out of five mice in the second treatment of double therapy with clindamycin and tetracycline relapsed and the peak level of parasitemia suppression was 52.3% at day 0. On the other hand, three out of five mice in single therapy with tetracycline, the peak level of parasitemia suppression was 33.1% at day 3 post the second treatment or at day 23 post parasites infection. Mice which had a relapse of parasitemia at a level of more than 3% finally died, even though treated. Parasitemia was cleared at the same day (day 5 and 6 post the second treatment) in all the treatment groups. No mortality was found in the treated animal groups either in the first treatment of double therapy or single therapy. On the other hand, at day 4 post treatment mortality was 100% for the untreated group although no data are shown.

**Effect of challenge with B. rodhaini on mice which were cured with selected antiprotozoal drugs:** Mice cured by the
Generally in Japan, diminazene diaceturate is used against pathic polyneuritis, Landry-Guillain-Barre Syndrome [9]. This drug has some bad side effects, one of which is that diminazene diaceturate has been used as a choice drug against babesiosis. Nowadays, diminazene diaceturate is used against babesiosis. A variety of drugs were used in order to choose the efficacious drugs against babesiosis. Nowadays, diminazene diaceturate has been used as a choice drug against babesiosis. This drug has some bad side effects, one of which is that human patient with B. microti infection develop an acute idiopathic polyneuritis, Landry-Guillain-Barre Syndrome [9]. Generally in Japan, diminazene diaceturate is used against B. gibsoni infected dogs. Some small animal clinicians, experienced some toxic effects in dogs, such as vomit and central nervous system damage. Valinomycin and particularly, gramicidin D are significantly effective against B. rodhaini and T. parva infections. However, in the present study, clindamycin and tetracycline were effective for parasitic clearance on murine babesiosis at doses of 200 and 100 mg/kg/day, respectively. For identifying drugs effective for the treatment of human B. microti infections, 20 selected antiprotozoal agents or combinations of agents were tested for activity against B. microti in mongolian jirds (Meriones unguiculatus) [10]. In the present study 3 selected antiprotozoal agents, tetracycline, clindamycin and diminazene diaceturate, were used to examine the activity against B. rodhaini in ICR mice. Several members of a series of 1-(chlorophenoxyalkyloxy)-4,6-diamino-1,2-dihydro-2,2-dimethyl-1,3,5-triazinobromides (I) have shown marked suppressive activity against B. rodhaini as well as P. berghei infection in mice [5]. Clindamycin plus quinine was also effective but less so effective than atovaquone. When treatment was not started until parasitemia became established, Atovaquone in doses of 300, 150, and 80 mg/kg/day was effective in the recovery of all animals compared with 50% of those receiving 10 mg/kg/day of the drug and 10% of untreated control [3]. We found that, clindamycin alone was less effective than tetracycline. When treatment was not started until parasitemia became established, tetracycline in a dose of 100 mg/kg/day or clindamycin in 200 mg/kg/day.

**DISCUSSION**

A variety of drugs were used in order to choose the efficacious drugs against babesiosis. Nowadays, diminazene diaceturate has been used as a choice drug against babesiosis. This drug has some bad side effects, one of which is that human patient with B. microti infection develop an acute idiopathic polyneuritis, Landry-Guillain-Barre Syndrome [9]. Generally in Japan, diminazene diaceturate is used against B. gibsoni infected dogs. Some small animal clinicians, experienced some toxic effects in dogs, such as vomit and central nervous system damage. Valinomycin and particularly, gramicidin D are significantly effective against B. rodhaini and T. parva infections. However, in the present study, clindamycin and tetracycline were effective for parasitic clearance on murine babesiosis at doses of 200 and 100 mg/kg/day, respectively. For identifying drugs effective for the treatment of human B. microti infections, 20 selected antiprotozoal agents or combinations of agents were tested for activity against B. microti in mongolian jirds (Meriones unguiculatus) [10]. In the present study 3 selected antiprotozoal agents, tetracycline, clindamycin and diminazene diaceturate, were used to examine the activity against B. rodhaini in ICR mice. Several members of a series of 1-(chlorophenoxyalkyloxy)-4,6-diamino-1,2-dihydro-2,2-dimethyl-1,3,5-triazinobromides (I) have shown marked suppressive activity against B. rodhaini as well as P. berghei infection in mice [5]. Clindamycin plus quinine was also effective but less so effective than atovaquone. When treatment was not started until parasitemia became established, Atovaquone in doses of 300, 150, and 80 mg/kg/day was effective in the recovery of all animals compared with 50% of those receiving 10 mg/kg/day of the drug and 10% of untreated control [3]. We found that, clindamycin alone was less effective than tetracycline. When treatment was not started until parasitemia became established, tetracycline in a dose of 100 mg/kg/day or clindamycin in 200 mg/kg/day.

**Table 3. Effect of B. rodhaini challenge infection on mice**

<table>
<thead>
<tr>
<th>Treatment groups*</th>
<th>1 Clindamycin &amp; Clindamycin</th>
<th>2 Clindamycin &amp; Tetracycline</th>
<th>3 Tetracycline</th>
<th>4 Clindamycin</th>
<th>5 Control (Non-treated)</th>
</tr>
</thead>
<tbody>
<tr>
<td>%Parasitemia**</td>
<td>day 0 0.8 ± 1.30 0.5 ± 0.41</td>
<td>day 0 0.4 ± 0.25 5.7 ± 9.64</td>
<td>day 4 1.7 ± 2.94 3.5 ± 4.31</td>
<td>day 5 0.1 ± 0.23 1.6 ± 2.13</td>
<td>day 7 0.0 ± 0.00 0.0 ± 0.00</td>
</tr>
<tr>
<td>Mortality†</td>
<td>day 4 0/3 0/4 0/7 1/4 0/5</td>
<td>day 5 0/3 0/4 4/7 1/4 5/5</td>
<td>day 6 0/3 0/4 7/7 0/4 NT</td>
<td>day 7 0/3 0/4 0/7 0/4 NT</td>
<td>day 9 0/3 0/4 6/7 2/4 NT</td>
</tr>
</tbody>
</table>

* a) Cured mice were challenged with 1 × 10³ parasitized RBC around 2 weeks post parasitic clearance. b) Group 1. Clindamycin (the 1st treatment) and clindamycin (the 2nd treatment) in a dose of 100 mg/kg b.i.d. × 7 days, via oral route. Group 2. Clindamycin (the 1st treatment) and tetracycline (the 2nd treatment) in a dose of 100 mg/kg b.i.d. or s.i.d. × 7 days, via oral or S.C. route. Group 3. Tetracycline in a dose of 100 mg/kg s.i.d. × 7 days, by S.C. route. Group 4. Clindamycin in a dose of 200 mg/kg b.i.d. × 7 days, via oral route. Group 5. No treatment (control) with distilled water *ad libitum*. Each treatment group was composed of 3, 4, 5 or 7 mice. Challenge was performed within two weeks post parasitic clearance. c) All the treated animal groups had a relapse within 10 days post treatment, except group 4. d) Results are expressed as mean % parasitemia suppressed by drugs in treated and non-treated animals. Data were recorded on the day following challenge infection. e) The first day of treatment was defined as day 0. f) Not tested, because all the mice died. g) Ratio of dead and live mice in treated and non-treated groups.
was effective in the recovery of all the animals compared with clindamycin in doses of 100 mg/kg/day and 50 mg/kg/day and non-treatment. Although within 10 days post treatment with tetracycline or clindamycin the mouse group had a relapse, the peak level of parasitemia suppression in the first treatment of clindamycin-treated mouse group was higher than that in tetracycline-treated mouse group. On the other hand, parasitemia was cleared earlier in tetracycline-treated mouse group than in the first treatment of clindamycin-treated mouse group.

Based on our study, challenge with a dose of 10^7/head pRBC to cured of treated or re-treated mouse group within two weeks post parasitic clearance provoked a little parasitemia, however, finally resulted in clearance of parasitemia. This means that an immune state was present or established in the mice.

Our findings, showed that tetracycline was effective against B. rodhaini infection. Parasitemia could be eliminated from the murine-red blood cells by subcutaneous administration of tetracycline in a dose of 100 mg/kg. On the other hand, clindamycin or tetracycline was not completely effective against B. rodhaini infection of mice became they had a relapse of parasitemia within 10 days post treatment. This finding, however, does not necessarily indicate no effectiveness of clindamycin for murine babesiosis, because in immunocompetent animals static drugs may be able to work synergistically with mycin for murine babesiosis, because in immunocompetent ever, does not necessarily indicate no effectiveness of clindamycin or tetracycline was not completely effective against B. rodhaini infection of mice became they had a relapse of parasitemia within 10 days post treatment. This finding, however, does not necessarily indicate no effectiveness of clindamycin or tetracycline. Because in immunocompetent animals static drugs may be able to work synergistically with host protective immune responses. The experiments with mouse model clearly showed that diminazene diaceturate was capable of eliminating B. canis from the canine-red blood cell—SCID mice [1].

Based on our study, the administration of tetracycline or clindamycin did not provoke any kind of clinical symptoms such as diarrhea effect to the treated mice. We can propose a hypothesis that the first treatment with clindamycin for 7 days brought about such static effect as parasitemia relapsed upon cessation of drug administration [1], while the second treatment for 7 days with tetracycline was useful in eliminating of blood parasites. Various antiprotozoal drugs can not completely eliminate the parasites from the host. Relapse often occurs post administration of anti protozoal drugs, therefore, repeated treatments are necessary. Nowadays, almost all anti protozoal drugs have several side effects, so it is very difficult to find antiprotozoal drugs without side effects. Basically, the purpose of chemotherapy against protozoal infection is not to eliminate the parasite completely, but to induce persistent infection and finally to establish premunization. This condition actually clears hosts of clinical symptoms. Our experimental findings, suggest that clindamycin or tetracycline is possibly used as a drug of choice for antiprotozoal chemotherapy because of their fewer side effects. However, clindamycin more induces premunization against babesiosis in mice than tetracycline.

ACKNOWLEDGMENTS. The authors thank Prof. K. Ono of University of Tokyo for providing the materials of B. rodhaini, an Australian strain.

REFERENCES