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Leishmania braziliensis panamensis: Increased Infectivity Resulting from Heat Shock

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SMEJKAL, R. M., WOLFF, R., AND OLENICK, J. G. 1988. Leishmania braziliensis panamensis: Increased infectivity resulting from heat shock. Experimental Parasitology 65, 1–9. Promastigotes of Leishmania braziliensis panamensis were subjected to a heat shock transformation yielding an amastigote-like stage. During the process of conversion, the heat-induced differentiating form displayed an increase in infectivity (as determined by lesion size) accompanied by a total protein composition unlike that of the promastigote and a morphology resembling that of the amastigote. These biological/functional changes may be related to an involvement of a heat shock response in the differentiation of leishmania, thus having important implications in the development of prevention and treatment strategies.

INDEX DESCRIPTORS AND ABBREVIATIONS: Leishmania braziliensis panamensis; Protozoa, parasitic; Hemoflagellate; Heat shock; Infectivity; Heat-induced differentiating form; Developmental change; Morphology; Dulbecco’s phosphate-buffered saline (PBS); Sodium dodecyl sulfate (SDS); Polyacrylamide gel electrophoresis (PAGE); Heat shock protein (Hsp).

INTRODUCTION

Leishmania spp., the causative agents of a wide spectrum of diseases, are protozoan parasites whose life cycle requires them to live and multiply under extremes of environmental conditions. The promastigote form is a slender, motile flagellate found in nature in the alimentary tract of the sandfly vector at a temperature approaching that of ambient air (Killick-Kendrick 1979; Lainson 1982). During a blood meal, promastigotes from an infected sandfly are transmitted to the warmer environment of the bloodstream of the mammalian host. Once in the host, the parasites are engulfed by macrophages, and the round, amotile intracellular amastigote form is found inside the phagolysosomes of these macrophages (Mauel 1984; Pearson et al. 1983; Sadick and Raff 1985).

A great deal of recent effort in leishmania research has centered on determining how and where the promastigote is transformed into the amastigote form: in the bloodstream or inside the macrophage. The amastigote is more resistant to destruction by the macrophage than is the promastigote (Mauel 1984). Therefore, determining the trigger of the promastigote to amastigote transformation is potentially of great importance in developing protective strategies.

One possible trigger is the temperature shift that the parasite undergoes when transferred from the sandfly vector to the mammalian host. In response to such temperature shifts or to other environmental stresses, bacteria (Yamamori et al. 1978), yeast (McAlister et al. 1979), mammalian cells (Corces et al. 1981; Mirault et al. 1982; Pelham and Bienz 1982), and other cell types (Ashburner and Bonner 1979; Craig et al. 1982; Kelley and Schlesinger 1978; Pelham and Bienz 1982; Voellmy and Rungger 1982) produce a set of proteins
which seem to be highly conserved through evolution (Craig et al. 1982) and which function to protect the cells from the toxic effects of such stresses (Ashburner 1982). The onset of the synthesis of these heat shock proteins is rapid and, in some cases, accompanied by a concomitant decrease in the synthesis of proteins normally produced (Ashburner 1982; Ashburner and Bonner 1979; Craig et al. 1982; Duncan and Hershey 1984; McKenzie et al. 1975).

Differences in protein composition, especially in protein surface components, may be expected in forms of an organism found in very different environments in nature. This is true for the promastigote and amastigote forms of Leishmania. Immunological analyses of promastigotes and amastigotes of several Leishmania species have resulted in the identification of stage-specific and common protein antigens (Handman and Curtis 1982; Handman et al. 1984; Smejkal et al. 1984; Sadick and Raff 1985; Pan 1986). In general, a comparison by these investigators of the total or surface complement of protein antigens has revealed mostly shared, and only a few (3–7) stage-specific, antigens; some antigens, although shared, were found to be predominantly expressed in either promastigotes or amastigotes. During the intracellular transformation of L. mexicana from promastigotes to amastigotes in macrophages, changes have been observed mostly in protein bands ranging from Mₚ, 24,000 to 68,000 (Chang and Fong 1982). Several proteins specific to the amastigote have been postulated to provide protection against the hostile environment of the phagolysosome (Handman and Greenblatt 1977; Gottlieb and Dwyer 1981) and to play a role in the increased survival rate of amastigotes in macrophages (Mauel 1984).

Several groups (Hunter et al. 1982, 1984; Hansen et al. 1984; Smejkal et al. 1984; Van der Ploeg et al. 1985) have described an in vitro temperature-induced conversion of the promastigote of various Leishmania spp. to a round, aflagellated, extracellular form resembling the amastigote. Known variously as an axenic amastigote (Hansen et al. 1984; Smejkal et al. 1984), a culture-derived amastigote (Hunter et al. 1982), a differentiating form (Hunter et al. 1984), and a high-temperature promastigote (Van der Ploeg et al. 1985), this form is obtained from promastigotes in culture by raising the temperature from 26 to 34°C. During temperature-induced differentiation, seven actively synthesized proteins (underexpressed or absent from the promastigote) have been identified with Mₚ, 83,000, 70,000, 68,000, 27,000, 23,000, and 22,000 (Hunter et al. 1984). These proteins correspond in molecular weight to a small set of proteins, termed Hsps; the synthesis of these Hsps has been found to be induced as a response to a heat shock or stress situation in every eukaryotic and prokaryotic organism thus far examined (Bienz 1985). Indeed, sequences related to cloned heat shock genes, Hsp70 and Hsp83, from Drosophila have been detected in the nuclear DNA of L. major with hybridization analyses producing similar patterns for the lesion amastigotes and the differentiating high-temperature promastigote (Van der Ploeg et al. 1985).

However, to our knowledge, there has been no evidence that these responses and changes reflect development of promastigotes into an infective stage. We now report that, concomitant with changes in morphology and protein composition, the infectivity of heat-shocked leishmania promastigotes greatly exceeds that of promastigotes.

**Materials and Methods**

*Leishmania braziliensis panamensis*, stock WR470, was originally obtained from a patient presenting at the Walter Reed Army Medical Center with cutaneous leishmaniasis. Promastigotes were grown at 26°C in Schneider’s Drosophila Medium (Biotluids) containing 25% fetal bovine serum (GIBCO) and 50 µg gentamycin/milliter of medium unless otherwise stated. Fresh stablates from the same freeze-down were defrosted for each experiment.
To convert promastigotes to heat-induced differentiating forms, log-phase promastigote cultures were washed extensively in PBS, counted, and aliquotted into separate flasks in fresh Schneider’s medium at 5–10 \times 10^5 cells/ml. Those cultures to be converted were then transferred to 34°C, while control cultures were maintained at 26°C. In our hands, the parasites remained just as viable and converted just as well in Schneider’s medium as in Medium 199 as described by Hansen et al. (1984). Carrying out the conversion in Schneider’s medium allowed for a more direct comparison between cells grown at 26 and 34°C, controlling for any possible artifacts generated by growing the cells in two different media.

At time 0 and at each time point thereafter, an aliquot of each culture was counted and checked for viability and morphological type by the method of Jackson et al. (1985) using a Zeiss microscope equipped with uv optics. This test for viability is a double dye technique, using ethidium bromide and fluorescein diacetate. The diacetate is metabolized in viable cells to yield the green fluorescing dye, while dead cells emit the red color characteristic of ethidium bromide. Samples of culture were harvested by centrifugation, washed, and then suspended in the same cell density (5–10 \times 10^5/ml). An aliquot for animal infections was removed from each sample and the remainder of the cell suspension was centrifuged and stored as a dry pellet at −70°C until all samples were ready for analysis by PAGE.

The infectivity of each sample was tested in adult (age 5 months) male Syrian hamsters (Charles River). Between 24 and 48 hr prior to infection, the hamsters were shaved at the base of the tail. Parasites from the aliquots reserved for infecting were centrifuged and suspended to approximately 5 \times 10^6 cells/ml in Medium 199. One animal was mock infected by the injection of Medium 199 alone. Three animals per experimental condition were injected intradermally at the base of the tail with 30 μl Medium 199 containing 2–5 \times 10^5 parasites. The 2.5-fold difference in parasite number is within acceptable limits to ensure reproducibility of results (Pan and Honigberg 1985, unpublished observations). The animals were checked periodically for appearance of a papule or lesion, and the affected area was measured.

The protein composition of cells at each time point was analyzed by SDS-PAGE according to the method of Laemmli (1970). Pellets, each containing 10^6 organisms, were solubilized in SDS sample buffer (2% SDS, 5% mercaptoethanol, 2% Nonidet-P40, 10% glycerol, and 0.001% bromphenol blue). Extracts were heated at 95°C for 3 min, loaded onto a 0.75-mm-thick slab gel containing a 6% stacking gel and a 12.5% separating gel, and then electrophoresed at constant voltage. Gels were silver stained using the method of Morrissey (1981).

**RESULTS**

Promastigotes of *Leishmania braziliensis panamensis*, strain WR470, were converted to heat-induced differentiating forms by transferring culture to 34°C. The cultures were monitored at various time intervals by sampling an aliquot immediately before harvesting. A cell count was obtained, and the cells were observed for viability, morphology, and motility. As shown in Fig. 1 and Table I, promastigotes cultured at 26°C remained elongated and motile throughout the course of the experiment. The number of viable cells dropped slightly by the end of 120 hr, probably due to a depletion of nutrients, since cultures were not supplemented during the experiment. The cells shifted to 34°C began to change their morphology between 2 and 5 hr, and by 9 hr, most had become round and amoeboid, resembling amastigotes. Although the percentage of viable cells at 48 hr was slightly less in the culture grown at 34°C than in the one grown at 26°C, the viability of both cultures was essentially the same by 120 hr.

The infectivity of the parasites at various stages in the conversion of promastigote to heat-induced differentiating form was determined by injecting a suspension of parasites into a susceptible animal at various time points and observing the appearance and size of resulting papules or lesions. The results are shown in Table II. The inoculum obtained at time 0 produced a papule or lesion by 40 days after infection. The size of the affected area grew quickly to 9.5–11 mm and was evident for 60 days. Essentially, the same infectivity was manifested by cells grown for 2 hr at 26°C. However, after 2 hr growth at 34°C, the inoculum caused a papule or lesion which appeared 4 days earlier, grew to almost twice the size of that produced by the cells from time 0 or 2 hr at 26°C, and lasted 84 days. The size of the affected area and the length of time it was observed continued to in-
FIG. 1. Fluorescence micrographs of *Leishmania braziliensis panamensis* promastigotes and organisms undergoing temperature-induced conversion. (a) Promastigotes after 48 hr at 26 C. The cells retain the morphology they had at time 0; (b) 2 hr at 34 C; (c) 4 hr at 34 C; (d) 24 hr at 34 C; (e) 48 hr at 34 C.
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TABLE I
Conversion of Promastigotes to Heat-Induced Differentiating Forms

<table>
<thead>
<tr>
<th>Time (hr)</th>
<th>Temperature (C)</th>
<th>% Viable cells</th>
<th>Morphology</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>26</td>
<td>&gt;99</td>
<td>Slender, motile</td>
</tr>
<tr>
<td>2</td>
<td>26</td>
<td>&gt;99</td>
<td>Slender, motile</td>
</tr>
<tr>
<td>2</td>
<td>34</td>
<td>&gt;99</td>
<td>Slender, motile</td>
</tr>
<tr>
<td>5</td>
<td>34</td>
<td>94.1</td>
<td>50% Round and amotile</td>
</tr>
<tr>
<td>9</td>
<td>26</td>
<td>&gt;99</td>
<td>Slender, motile, and clumped</td>
</tr>
<tr>
<td>9</td>
<td>34</td>
<td>96.7</td>
<td>&gt;75% Round and amotile</td>
</tr>
<tr>
<td>24</td>
<td>26</td>
<td>&gt;99</td>
<td>Slender, motile, and clumped</td>
</tr>
<tr>
<td>24</td>
<td>34</td>
<td>95.7</td>
<td>Round, amotile</td>
</tr>
<tr>
<td>48</td>
<td>26</td>
<td>93.5</td>
<td>Slender, motile, and clumped</td>
</tr>
<tr>
<td>48</td>
<td>34</td>
<td>87.2</td>
<td>Round, amotile, and clumped</td>
</tr>
<tr>
<td>120</td>
<td>26</td>
<td>82.2</td>
<td>Slender, motile, and clumped</td>
</tr>
<tr>
<td>120</td>
<td>34</td>
<td>81.3</td>
<td>Round, amotile, and clumped</td>
</tr>
</tbody>
</table>

Note. Promastigotes in log phase culture were aliquotted into flasks, which were maintained at 26 C or transferred to 34 C. At the times indicated, a flask from each temperature was sampled and observed for viability and morphology.

crease throughout the experiment, and after 120 hr, the infectivity of both cultures was about equal.

The protein composition of the cells at each sampling time was analyzed by PAGE (Fig. 2). The protein composition of the cells in culture at 26 C showed few qualitative and only some quantitative changes until 120 hr, at which point some notable changes were observed. In contrast, the cultures transferred to 34 C showed a dramatic decrease in the number of proteins present as the time at the elevated temperature increased, a phenomenon observed in other systems, as well (Craig et al. 1982; Ashburner and Bonner 1979; Ashburner 1982; McKenzie et al. 1975; Duncan and Hershey 1984). Bands at Mr 84,000 and Mr 81,000 (c, d) had decreased by 48 hr and were gone by 120 hr at 34 C. Also notable was the disappearance of a number of bands below Mr 40,000. For example, after 9 hr at 34 C, bands at Mr 38,000 and Mr 36,000 (o, p) began disappearing. Another band at Mr 29,000 (r) decreased in intensity after 24 hr at 34 C. A band at Mr 25,000 (t)

TABLE II
Infectivity of Promastigotes and Heat-Induced Differentiating Forms

<table>
<thead>
<tr>
<th>Sample (hr°C)</th>
<th>Time of appearance (days after infection)</th>
<th>Time of maximum size (days after infection)</th>
<th>Maximum size (mm)</th>
<th>Duration (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>40</td>
<td>48-70</td>
<td>9.5-11</td>
<td>60</td>
</tr>
<tr>
<td>2/26</td>
<td>40</td>
<td>48-66</td>
<td>10-12</td>
<td>50</td>
</tr>
<tr>
<td>2/34</td>
<td>36</td>
<td>50-73</td>
<td>17-18.5</td>
<td>84</td>
</tr>
<tr>
<td>9/26</td>
<td>36</td>
<td>48-70</td>
<td>12-15</td>
<td>64</td>
</tr>
<tr>
<td>9/34</td>
<td>34</td>
<td>51-71</td>
<td>20-21</td>
<td>86</td>
</tr>
<tr>
<td>48/26</td>
<td>35</td>
<td>47-65</td>
<td>17.5-20</td>
<td>90</td>
</tr>
<tr>
<td>48/34</td>
<td>33</td>
<td>48-70</td>
<td>25-32</td>
<td>92</td>
</tr>
<tr>
<td>120/26</td>
<td>32</td>
<td>41-67</td>
<td>22-28</td>
<td>93</td>
</tr>
<tr>
<td>120/34</td>
<td>32</td>
<td>42-61</td>
<td>22-28</td>
<td>93</td>
</tr>
</tbody>
</table>

Note. Cells at each timepoint were injected intradermally into Syrian hamsters at the base of the tail. The animals were checked periodically for appearance of a papule or lesion, and the affected area was measured.
was one of the few prominent bands in this region after 120 hr at 34 C. This band, however, was present in both culture systems after 9 hr at their respective temperatures. The band was more prominent in the 26 C samples and persisted with equal intensity throughout the time frame of incubation. A weaker, but similarly persistent, band was noted for the 34 C samples.

Protein bands at Mr's 61,000, 59,000, 55,000, 52,000, 49,000, 46,000, 43,000, and 39,000 (g-n, respectively) were present in all samples, but some were more abundant in the later 120-hr, 34 C samples. The Mr 55,000 and Mr 52,000 bands most likely represent the α and β subunits of tubulin. These bands were more prominent in the 26 C samples but markedly decreased in intensity at 120 hr; 34 C samples began to show a progressive decrease beginning at 9 hr. During the course of the experiment, a number of bands at Mr's 93,000, 87,000, 72,000, 68,000, and 28,000 (a, b, c, f, s) were observed earlier or were more prevalent in the 34 C samples. A doublet band at Mr 36,000 and Mr 35,000 (p, q), appearing in early 26 and 34 C samples, maintained its Mr 35,000 intensity but gradually lost the discrete Mr 36,000 component at 26 C; at 34 C both bands weakened considerably with successive sampling, disappearing at 48 hr with only the Mr 35,000 component strongly reappearing at 120 hr.

**DISCUSSION**

*Leishmania* spp. promastigotes transform into intracellular amastigotes after being transmitted by the sandfly to the warmer environment of the mammalian host. Amastigotes have a greater infectivity for experimental animals than do promastigotes, possibly due to their increased ability to survive the defense mechanisms of the host (Mauel 1984). Like true amastigotes, the infectivity of *L. braziliensis panamensis* heat-induced differentiating forms also appeared to be greater than that of promastigotes. Upon culture at 34 C, promastigotes of *L. b. panamensis* changed in morphology to that resembling intracellular amastigotes. However, by 2 hr after transfer to 34 C and even before the change in morphology began, the infectivity of the cells had increased (Tables I and II).

The heat shock phenomenon has been implicated in the transformation of promastigotes to heat-induced differentiating forms by Hunter *et al.* (1982, 1984) and confirmed by Van der Ploeg *et al.* (1985), who demonstrated sequence homology be-
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tween the nuclear DNA of L. major and the
Drosophila heat shock genes. These genes
were found to be similar in the mammalian
and high-temperature culture forms of the
parasites. In many organisms, the tran-
scriptional activation of heat shock genes
in response to an increase in temperature
causes the rapid appearance of heat shock
proteins. In some case, as these genes are
activated, others coding for proteins nor-
rmally produced are turned off, causing a
decrease in synthesis of the latter proteins
(McKenzie et al. 1975; Kelley and Schles-
singer 1978; Ashburner and Bonner 1979;
Ashburner 1982; Craig et al. 1982; Voellmy
and Rungger 1982; Duncan and Hershey
1984). The decrease in intensity of protein
bands progressing to eventual disappear-
ance and the predominance of a small set of
proteins (MW 28,000 to 87,000), coupled
with the rapid change in infectivity when
promastigotes in culture were transferred
to 34°C, supports the involvement of a heat
shock response that may promote differen-
tiation into a more infective form or stage
similar to the amastigote. It is reasonable to
assume that the shift in temperature not
only activates heat shock genes to produce
traditional Hsps but may also trigger the
production of stage-specific proteins. such
as those described by us (Smejkal et al.
1984) and by Sadick and Raff (1985). Cer-
tain of these proteins, particularly if sur-
face membrane-associated, may play an
important role in parasite infectivity or viru-
ulence. In this regard, the increased infect-
vity and morphologic change likely reflect
the alterations in the total protein profiles
with one or more newly or predominantly
synthesized proteins becoming responsible
for or involved in the developmental differ-
etiation leading to an increase in viru-
ulence. The identity of such proteins and
their presumed role in developmental
changes have yet to be demonstrated. It is
likely that the stimulus resulting in in-
creased infectivity is concurrently the
trigger for synthesis of those proteins
which appear as changes in the electropho-
retic pattern (Fig. 2) beginning at 9 hr after
transfer and which may be responsible for
the change in morphology coinciding with
their appearance. One or more of these
proteins may also be responsible for the in-
crease in infectivity. The differences in the
electrophoretic patterns produced by cells
cultured at 34°C and those at 26°C are par-
ticularly significant in view of the observed
changes in infectivity and the evidence that
infectivity may be dependent on the ability
of surface components to protect the para-
site from host defense mechanisms (Mauel
1984).

Recently, heat shock has been shown to
trigger the induction of sexual reproduction
in Volvox, the females of which otherwise
reproduce asexually (Kirk and Kirk 1986).
The rapid conversion by heat shock of one
cell type to another in other organisms is,
thus, probable. Developmental processes
occurring during oogenesis may also acti-
vate heat shock genes (Bienz 1985). In con-
trast to the rapid response of heat shock
genes to environmental stress, the develop-
mental activation process appears to be
slow. Increased attention is currently being
given to both types of heat shock responses
in organisms whose life cycles take them
from an environment that is temperature
regulated to one that is not. Both mecha-
nisms of activation may occur during the
life cycle of leishmania.

Developmental activation of some heat
shock genes may occur during the “matu-
ration” process of promastigotes in the in-
sect vector. Promastigotes of Leishmania
tropica were found to be more infective
when taken from stationary phase cultures
or from sandflies 7 to 10 days after a blood
meal than from cultures growing logarith-
mically or from sandflies 3 days after a
blood meal (Sacks and Perkins 1984). We
have shown that L. b. panamensis promas-
tigotes, while maintaining a similar mor-
phology over the course of 120 hr, in-
creased in infectivity with time in culture at
26°C (Tables I and II). Injection of promas-
tigotes into a warm-blooded host by the
sandfly may trigger a heat-induced conversion similar to that produced in culture, where, in response to an increase in temperature, leishmania promastigotes undergo a conversion to heat-induced differentiating forms. Macrophages may then engulf the parasites already in the process of converting. Whether the changes that occur in the promastigote in culture as it transforms into the high-temperature form indeed mimic those that occur in a natural infection is unknown since an in vivo stage intermediate to the promastigote and amastigote has not yet been identified. However, the high-temperature differentiating forms will be of use in studying the promastigote to amastigote transformation and concomitant changes in infectivity and protein composition.

ACKNOWLEDGMENTS

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