Leishmania major: Parasite Interactions Suggesting Sexuality

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In five experiments, Leishmania (Leishmania) major (MRHO/SU/59/P-strain) grew poorly when seeded in FYTS medium supplemented with 15% fetal calf serum, but presented several peculiar pairs of promastigotes diametrically opposed and attached at their posterior ends (5.8-13.5%). As seen in Giemsa-stained smears, a ring-like line and/or an enlargement, generally occurred at the parasite junction. A close proximity of nuclei, which sometimes were difficult to distinguish from each other, was also observed at this junction. Several of these pairs appeared to be composed of fused cells in which the nuclei could be apparently fused, as shown by fluorescence microscopy to detect β-tubulin and DNA, and by scanning electron microscopy. Under other culture conditions these pairs were absent or occurred at very low rates (0.2-2.2%). Such pairs differ markedly from longitudinally dividing cells and resemble those described in two other Leishmania species, as well as in Herpetomonas megaseliae and Phytomonas davidii, suggesting steps of a putative sexual process.

Key words: Trypanosomatidae - Leishmania major - sexuality - morphology - immunocytochemistry - scanning electron microscopy

The occurrence of sexuality in Leishmania has been a subject of interest since the beginning of this century, but several older cytological observations ascribed to sexual events were unconvincing (Rogers 1904, Adie 1921-22, Christophers et al. 1926, Wenyon 1926, Wenrich 1954). This subject was discredited, until Maazoun et al. (1981) reported heterozygous patterns in enzyme electrophoretic variants of three Leishmania species, and their paper was followed by several others describing naturally occurring putative hybrids using biochemical and/or molecular methods (Le Blancq et al. 1983, Le Blancq & Peters 1986, Evans et al. 1987, Pagès et al. 1989, Kelly et al. 1991, Cupolillo et al. 1992, Belli et al. 1994, Piarroux et al. 1994, Dujardin et al. 1995). However, genetic recombination between species or strains of Leishmania has not yet been demonstrated under laboratory conditions (Gradoni et al. 1986, Evans et al. 1989, Panton et al. 1991, Shehata et al. 1991), although Lanotte and Rioux (1990) had recorded by videocinematography fusion of pairs of asposed promastigotes in cultures of two species, and also evidenced the possibility of nuclear fusion by examining these cultures in Giemsa-stained smears.

In this paper we describe pairs of asposed promastigotes occurring at relatively high percent in L. (L.) major (P strain) when seeded in a medium described for insect trypanosomatids (Roitman et al. 1972), which we had supplemented with fetal calf serum. Such pairs were studied on Giemsa-stained smears, by fluorescence microscopy to examine the β-tubulin distribution and nuclear DNA features, as well as by scanning electron microscopy. The frequency of such pairs under other culture conditions was also examined.

MATERIALS AND METHODS

Promastigotes of a L. (L.) major strain (MRHO/SU/59/P) grown in BHI+LIT medium overlaying blood-agar (Jaffe et al. 1984) were seeded in FYTS medium (Roitman et al. 1972) supplemented with 15% heat-inactivated fetal calf serum (FYTS+15%FCS), pH 7.0, which was distributed in 4-ml-volumes in 16x150 mm screwcap tubes, and kept at about 27°C. Following the finding in fresh preparations of unusual pairs of apposed cells, Giemsa-stained smears were prepared as previously described (Sousa 1994) for morphological studies. Subsequently, two experiments were performed to compare the frequency of such pairs in FYTS+15%FCS with those in LIT medium (Chiari & Camargo 1984), LIT/blood-agar and FYTS+15%FCS/blood-agar. The inoculum was always promastigotes from BHI+LIT/blood-agar cultures, which in the first experiment had no pair of asposed cells, while in the second one presented 0.5%. The cultures were maintained at about 27°C and the percent of pairs of asposed cells was determined daily, from the 2nd to the 7th days, by

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examining at least 500 cells in fresh preparations. The maximum parasite growth was compared in two culture conditions, FYTS+15%FCS and LIT/blood-agar, in both cases beginning with about 8.5x10⁵ cells/ml. The occurrence of pairs of apposed cells in FYTS+15%FCS was also verified by using promastigotes from LIT/blood-agar as inoculum (3 assays).

Pairs of apposed cells from a 5-day old culture in FYTS+15%FCS were studied under fluorescence microscopy to examine concomitantly the β-tubulin distribution and the nuclear DNA. Then, the cells were washed in phosphate buffered saline (PBS), pH 7.2, allowed to adhere to 0.1% poly-L-lysine-coated coverslips, fixed for 5 min at room temperature with 4% paraformaldehyde in PBS and rinsed three times in PBS containing 0.5% Triton X-100. Subsequently, the cells were incubated for 1 hr at 37°C with 1:50 anti-β tubulin monoclonal antibody (Sigma), rinsed and incubated for 1 hr with 1:25 fluorescein isothiocyanate (FITC)-coupled rabbit anti-mouse IgG; following a 10 min incubation with 2 mg/ml 4,6-diamidino-2-phenylindole (DAPI; Sigma), the coverslips were rinsed in 0.85% NaCl and mounted in 1.4-diazabicyclo-(2.2.2)-octane (DABCO; Sigma). The preparations were observed and photographed with a Zeiss microscope equipped for epifluorescence using selective FITC or DAPI filters.

For scanning electron microscopy, cells from a 5-day old culture in FYTS+15%FCS were adhered to poly-L-lysine coated coverslips as above-described, fixed for 1 hr at 4°C in 2.5% glutaraldehyde in 0.1 M Na-cacodylate buffer, pH 7.2, plus 3.5% sucrose, and subsequently washed in this same buffer. After post-fixation in 1% OsO₄ for 30 min, the material was dehydrated in graded series of acetone, dried with CO₂ by the critical-point method, mounted on aluminum stubs and coated with a 20 nm-thick gold layer. The samples were examined with a Carl Zeiss DSM 940 scanning electron microscope.

RESULTS
FYTS+15%FCS medium was unsuitable for growing *L. major* promastigotes, whose population increased somewhat more than twice within 48-72 hr, but thereafter gradually declined. However, such cultures presented a relatively high percent (5.8-13.5%) of peculiar pairs of promastigotes, at times very slender, diametrically opposed to each other and attached at their posterior ends, there generally occurring a ring-like line and/or an enlargement (Figs 1-9) sometimes resembling a hat brim (Fig. 5).
5), as seen in Giemsa-stained smears. In such pairs both nuclei were usually in close proximity and in some instances it was difficult to distinguish one from the other, either in Giemsa-stained slides or under DAPI fluorescence (Figs 3-6, 9, 12B-14B, 15, 16). Several pairs seemed composed of fused cells, as suggested either by indirect immunofluorescence of β-tubulin (Figs 12A, 13A, arrows) or by scanning electron microscopy, which also evidenced the peculiar enlargement and meeting of nuclei at the level of the parasite junction (Figs 17-21). Typically dividing cells could also be seen in FYTS+15%FCS (Figs 10, 11).

In fresh preparations, the pairs under study were actively moving and presented an uncommon brightness at the level of nucleus. The phenomenon of formation of these pairs in FYTS+15%FCS medium was reproducible in 5/5 assays, the inoculum being promastigotes either from BHI+LIT/blood-agar or LIT/blood-agar. The percent of these pairs in different culture conditions are given in Table. L. major grew luxuriantly in LIT/blood-agar and to a lesser extent in other media (FYTS+15%FCS/blood-agar>LIT>FYTS+15%FCS). Beginning with about 8.5x10^5 promastigotes/ml, the maximum growth in LIT/blood-agar was nearly 6x10^7 cells/ml (5th day), whereas in FYTS+15%FCS it was around 2x10^6 cells/ml (2nd day).

**DISCUSSION**

To our knowledge, in *Leishmania* species, pairs of promastigotes such as these described herein have not been reported previously. They also differ from those found in another L. major strain during experimental infection in *Lutzomyia longipalpis* (Walters et al. 1993), which were apposed paramastigotes. These pairs also markedly differ from cells joined laterally during longitudinal binary fission (compare Figs 1-9 with 10 and 11). It is worthy mentioning that although dividing trypanosomatids can occasionally be attached by their posterior ends before complete separation, they neither present any enlargement nor the proximity of nuclei at the level of their junction, as commonly seen here. However, the pairs of promastigotes described by us are somewhat similar to those reported in *Herpetomonas megaseliae* and *Phytomonas davidi* (Sousa 1991, 1994), and resemble those formed by fusion previously described in *L. infantum* and *L. tropica* (Lanotte & Rioux 1990), since the parasite attachment occurs by the posterior ends, the cells remaining apposed to each other, and on several occasions nuclear fusion appears to take place. Our studies to detect both β-tubulin and DNA in such pairs, as well as our findings by scanning electron microscopy, strongly suggest cellular fusion in the majority of cases, as well as close contact or even fusion of the nuclei (Figs 12A-21). Taking together our findings and those from the literature, we have considered the possibility that a fusion process to pro-
Leishmania major in Roitman’s FYTS medium supplemented with 15% heat-inactivated fetal calf serum. Scanning electron microscopy. Fig. 17: pair of very slender cells, diametrically opposed to each other, and presenting the nuclei at the level of their junction (X 3,100). Fig. 18: this pair illustrates the possibility of cellular fusion (X 7,000). Fig. 19: pair of very slender cells fused at their posterior ends, there occurring close contact of nuclei, arrow (X 3,340; X 6,500). Fig. 20: note the peculiar hat brim-like enlargement (arrow) at the parasite junction (X 3,600). Fig. 21: nuclear meeting at the cellular junction (X 6,750).

Mote nuclear interactions could have generated the pairs here described, this strongly suggesting a sexual process. Our data confirm that in trypanosomatids phenomena can occur in which the cytological bases for genetic recombination are clearly present.

The finding of putative hybrids of other L. major isolates (Le Blancq et al. 1983) and of L. major/L. arabica (Evans et al. 1987, Kelly et al. 1991) support the hypothesis of occurrence of genetic exchange in these species, although it has been considered infrequent (Le Blancq et al. 1983, Kelly et al. 1991). In the present study, we verified that the pairs of apposed promastigotes suggesting a sexual process were not common in media other than FYTS+15%FCS, they usually not being found or occurring in low numbers. It is accepted that genetic recombination mediated by sex is generally a major mechanism promoting diversity within a species, consequently enhancing its chances of survival in a fluctuating environment. As our paper shows close association between two cells and their nuclei, the question arises whether the unsuitable conditions for L. major growth in supplemented FYTS medium could have triggered the formation of such pairs to propitiate some type of
Percent of pairs of apposed promastigotes in Leishmania major under different culture conditions; parasites from BHI+LIT/blood-agar were used as inoculum

<table>
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<th>Days</th>
<th>FYTS+15%FCS</th>
<th>LIT</th>
<th>FYTS+15%FCS</th>
<th>LIT/blood-agar</th>
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<tr>
<td>2</td>
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<td>7.3</td>
<td>0.3</td>
<td>0.7</td>
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<tr>
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<td>13.5</td>
<td>2.0</td>
<td>1.2</td>
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<tr>
<td>4</td>
<td>7.7</td>
<td>5.8</td>
<td>0.3</td>
<td>1.1</td>
</tr>
<tr>
<td>5</td>
<td>8.5</td>
<td>5.7</td>
<td>2.2</td>
<td>0.9</td>
</tr>
<tr>
<td>6</td>
<td>7.8</td>
<td>5.8</td>
<td>0.8</td>
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</tr>
<tr>
<td>7</td>
<td>5.8</td>
<td>6.3</td>
<td>0</td>
<td>1.4</td>
</tr>
</tbody>
</table>

(1) and (2), respectively, the 1st and 2nd experiments.

BHI: brain heart infusion medium; LIT: liver infusion-tryptose medium; FYTS+15%FCS: folic acid-yeast extract-tryptose-sucrose medium supplemented with 15% fetal calf serum.

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REFERENCES


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