Morphological Comparison of Axenic Amastigogenesis of Trypomastigotes and Metacyclic Forms of Trypanosoma cruzi

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Amastigogenesis occurs first when metacyclic trypomastigotes from triatomine urine differentiate into amastigotes inside mammalian host cells and a secondary process when tissue-derived trypomastigotes invade new cells and differentiate newly to amastigotes. Using scanning electron microscopy, we compared the morphological patterns manifested by trypomastigotes and metacyclic forms of Trypanosoma cruzi during their axenic-transformation to amastigotes in acidic medium at 37°C. We show here that in culture MEMTAU medium, secondary and primary axenic amastigogenesis display different morphologies. As already described, we also observed a high differentiation rate of trypomastigotes into amastigotes. Conversely, the transformation rate of in vitro-induced-metacyclic trypomastigotes to amastigotes was significantly slower and displayed distinct patterns of transformation that seem environment-dependent. Morphological comparisons of extracellular and intracellular amastigotes showed marked similarities, albeit some differences were also detected. SDS-PAGE analyses of protein and glycoprotein from primary and axenic extracellular amastigotes showed similarities in glycopeptide profiles, but variations between their proteins demonstrated differences in their respective macromolecular constitutions. The data indicate that primary and axenic secondary amastigogenesis of T. cruzi may be the result of different developmental processes and suggest that the respective intracellular mechanisms driving amastigogenesis may not be the same.

Key words: Trypanosoma cruzi - primary amastigogenesis - secondary amastigogenesis - differentiation processes

Trypanosoma cruzi, the etiological agent of Chagas disease is a Stercorarian trypanosome, transmitted as infective metacyclic trypanomastigotes to mammalian host via the feces or urine of the insect vector. Flagellated trypomastigotes are ingested by vertebrate cells through a process referred to as parasite-directed endocytosis (Burleigh & Andrews 1995) with formation of a parasite-containing endocytic vacuole which fuse with lysosomes from the host cell (Carvalho & De Sousa 1989, Tardieux et al. 1992). Within 2 h after infection, the trypomastigotes can leave the acidic environment as a phagosome before the transformation into amastigotes is completed (Ley et al. 1990) and enter the slightly alkaline environment of the cytoplasm, where they multiply as aflagellate amastigotes (Burleigh & Andrews 1995). These early events appear to occur at different rates depending on the type of vertebrate cells studied. It has been shown by optical microscopy that following vertebrate cell penetration, metacyclic trypomastigotes reorganize into amastigotes (in about 3 h) and remain morphologically quiescent in a lag period within the cell for about 35 h (at 35°C) prior to the onset of reproduction (Dvorak 1975). It has been proposed that the lag period observed, at least in part, must represent the time necessary for their transformation into amastigotes (Ley et al. 1988). The amastigotes then divide and transform into trypomastigotes. Upon leaving the cell, the tissue-derived trypomastigotes enter the bloodstream, and are capable of infecting surrounding cells or disseminate to other tissues via the bloodstream, repeating the intracellular events. The tissue-derived trypomastigotes can be eventually taken up by the insect host. In the intestine of the invertebrate host the blood trypomastigotes transform into epimastigotes. The epimastigotes then divide and give rise to infective metacyclic trypanomastigotes. Consequently, a primary amastigogenesis occurs when metacyclic trypomastigotes from triatomine urine differentiate into amastigotes inside mammalian host cells, and a secondary process is observed when tissue-derived trypomastigotes differentiate into amastigotes.

Most of the information concerning the in vitro and in vivo amastigogenesis process comes from the study of tissue-derived trypomastigote forms (Nogueira & Cohn 1976, Milder et al. 1977, Ley et al. 1988, Burleigh & Andrews 1995, Barros et al. 1996, 1997), but little is known about the developmental process driving the metacyclic forms to amastigotes. Although, metacyclic trypomastigotes can appear similar in morphology and share some biological properties with tissue-derived trypanomastigotes. However, the latter has the capacity to transform into epimastigotes at 27°C in blood or axenic medium, a property that is absent in metacyclic trypanomastigotes (Brener 1973). Both forms also have stage-specific antigens and they display different modes of interaction with host cells (Burleigh & Andrews 1995). For instance, metacyclic trypomastigotes express on their...
surface several glycoproteins that interact with mammalian cells, which have no counterpart in blood trypomastigotes (Ruiuz et al. 1993).

Using axenic conditions, several investigators have reported the morphological differentiation of bloodstream trypomastigotes into amastigotes and the morphological and ultrastructural changes throughout transformation have been described (Villalta & Kierzenbaum 1982, Andrews et al. 1987, Kambara et al. 1990, Tomlinson et al. 1995). In a carefully performed work, Andrews et al. (1987) studied the morphological transformation by transmission electron microscopy (SEM), of tissue-culture derived trypomastigotes into amastigotes and described a complex pattern of morphological rearrangements which occurred during 24 to 48 h of incubation and found that the morphological changes were associated with the expression of specific membrane antigens. Tomlinson et al. (1995) confirmed by SEM and antigenic analysis that the incubation at 37°C of tissue-culture derived trypomastigotes in acidic media accelerated greatly the morphological transformation, which was also accompanied by the acquisition and loss of specific membrane antigens. Other authors have obtained extracellular round amastigote-like forms from metacyclic trypomastigotes using highly enriched media (Kimura et al. 1978, Pan 1978, Rondinelli et al. 1988). Taking these observations together along with the little attention given to comparative morphology between the transformation of trypomastigotes and metacyclic forms into amastigotes raised the question of whether both transformations occurred following similar or different developmental patterns.

To address this question, we have studied by SEM the morphological events that occur during the primary and secondary amastigogenesis of T. cruzi. Furthermore, we have compared intracellular amastigotas with their corresponding extracellular amastigotas.

**MATERIALS AND METHODS**

**Parasites and stages** - Throughout this study, cloned EPm6 isolate of T. cruzi was used. The parasite was cloned and maintained by alternate triatomine/mouse passages as previously described (Contreras et al. 1994). The procedures to obtain in vitro-induced metacyclic and extracellular-derived metacyclic amastigotes were described in details previously (Contreras et al. 2002).

**In vitro primary amastigogenesis** - The transformation kinetic to produce extracellular-amastigotes from in vitro-induced metacyclic trypomastigotes (EMA) was described (Contreras et al. 2002). In brief, DEAE-52-purified metacyclic trypomastigotes were transferred to sterile plastic tissue culture flasks (175 cm² Falcon Labware, Oxnar, CA) containing 14 ml of MEMTAU medium pH 5.8, which consists of a 1:1 mixture of TAU3AAG medium and MEM 10% FBS medium, supplemented with 70 mM Sucrose, 20 µg/ml bovine or human Hemoglobin, 200 U/ml Penicillin, 200 µg/ml Streptomycin, and 20 mM MES [2 (N-morpholinoethanesulfonic) acid hydrate], followed by incubation at 37°C, without agitation in a 5% CO₂ atmosphere for one, two or three days, namely as pre-incubation phase. After three days of pre-incubation, the parasites were centrifuged, resuspended, transferred to sterile culture flasks and incubated for one, two or three days under the same conditions described above, namely as the re-incubation phase.

**In vitro secondary amastigogenesis** - Extracellular-derived trypomastigote amastigotes (ETA) were obtained essentially as described by Tomlinson et al. (1995) except by incubating in MEMTAU medium. Tissue-culture trypomastigotes from infected Vero cells were concentrated by centrifugation and resuspended in MEMTAU medium at 37°C, without agitation in a 5% CO₂ atmosphere for 3, 6, 9, 12, 18 and 24 h.

**In vitro intracellular amastigotes** - Intracellular-derived metacyclic trypomastigotes (IMA) and intracellular-derived trypomastigote amastigotes (ITA) were obtained by disruption of infected Vero cells prior to trypanostagogenosis, i.e., approximately three days after the infection of the monolayers with in vitro induced-metacyclic trypomastigotes or tissue-culture trypomastigotes, respectively. Infected monolayers were gently trypsinized, and the detached cells were recovered by centrifugation and resuspension in PBS (0.15M sodium chloride, 0.02M sodium phosphate, pH 7.2) supplemented with 1% BSA (Bovine Serum Albumin) (PBS-BSA). A 10 ml aliquot of the resuspension was disrupted by passage through a 25-gauge needle (Piras et al. 1982). The intracellular amastigotes were separated from the cell debris by centrifugation in a 15-21% discontinuous Metrizamide gradient (Carvalho & De Souza 1983).

**SEM** - Purified metacyclic trypomastigotes (day 0), pre-incubated (days 1, 2 and 3), re-incubated (days 1, 2 and 3), IMA, ITA, trypomastigotes (0 h) and from different times of incubation (3, 6, 9, 12, 18 and 24 h) were analyzed by SEM. Sample processing was carried out essentially as described by Andrews et al. (1987) using glass coverslips precoated with 10 µg/ml poly-L-lysine. Glutaraldehyde-fixed parasites were post-fixed with 1% OsO₄ in sodium cacodylate buffer, dehydrated in graded ethanol and critically point dried from liquid CO₂. Specimens were coated with gold-palladium in a BAL-TEC SCD050 evaporator before being examined in a Philips XL-20 scanning electron microscopy.

**Analysis of polypeptides and glycopeptides by SDS-PAGE** - EMA and ETA proteins were analyzed by SDS-PAGE (Laemmli 1970) on 10% uniform polyacrylamide slab minigels. Wells were loaded with 7 µg of protein. The resulting protein patterns were stained with a combined Coomassie Blue-Silver staining procedure (De Moreno et al. 1985) for proteins and combined Periodic Acid-Alician Blue-Glutaraldehyde-Silver (PAABGS) stain procedure for glycoproteins (Dubray & Bezard 1982, Moller et al. 1993, Moller & Poulsen 1995). For glycoproteins, two types of markers were run in the same gel: the broad molecular weight range was used to estimate molecular weight and control the reversal of silver staining (unstaining) of proteins and a mixture of proteoglycans and glycoproteins (6 µg/well). The following glycoconjugates were used: Chondroitin 6-Sulfate from shark cartilage (type C, Sigma C-4384), Mucin from bovine sub maxillary glands (type I-S, Sigma M-3895), Fetuin from fetal calf serum (Sigma F-2379), and Bovine α-Acid Glycoprotein (orosomucoid, Sigma G-9014). The gels were then scanned.
RESULTS

In vitro transformation of tissue-culture derived trypomastigotes into amastigotes - A detailed morphological analysis of the extracellular differentiation kinetic of tissue culture-derived trypomastigotes into amastigotes was carried out using SEM. Parasites released at the third day after infection from vero cells in culture showed a typical trypomastigote bloodstream morphology (Fig. 1a), characterized by C- or S-shaped, thick and elongated bodies (total length 12.88 ± 1.98 µm; main width 1.19 ± 0.20), prominent kinetoplast terminal or sub-terminal (1.47 ± 0.48 µm from posterior end, arrow Fig. 1a), thick moderate to long flagella and a conspicuous undulating membrane. When these parasites are incubated in MEMTAU medium at 37°C, an accelerated appearance of forms resembling amastigotes occurs within 24 h (Fig. 1b-j). Within 3 h of incubation, broad trypomastigotes (Fig. 1b) and parasites with twisted body and sharp posterior cell end (Fig. 1c) were detected. At 6 h, it was possible to observe a progressive sinking of the flagellum into the cytoplasm of the parasites (Fig. 1d-e) maintaining sharp the posterior cell end. At 9 h of incubation, the parasites present a nut-shaped morphology with longitudinal cracks and short free flagellum (Fig. 1f). In the next 12 h the parasite body fold around itself in helical fashion and an increasing degree of internalization of the flagellum can be seen (Fig. 1g). Between 18 h (Fig. 1h) and 24 h of incubation, typical amastigotes became predominant and the transformation process accomplished (Fig. 1i-j). During the transformation process, different types of trails were released by the parasites, as described in the legend to Fig. 1.

Comparing ETA from 24 h of incubation (Fig. 1i-j) to their corresponding ITA isolated by Metrizamide gradient (Fig. 1k-l), a remarkable, gross morphological similarity can be seen. The bodies of both types of amastigotes are rounded or oval-shaped, have longitudinal cracks and a short protruding flagellum. However, their surface membranes display differences: ITA presents a smooth surface, while ETA exhibits a finely grained surface.

In vitro transformation of metacyclic trypomastigotes into amastigotes - With the aim of looking at the overall aspect of parasites during the transformation process, we examined the differentiating cells by SEM (Figs 2, 3). The TAU3AAG-purified parasites of the pre-incubation phase show a typical morphology of metacyclic trypomastigotes (Fig. 2a-b), characterized by a long and slender body (total length 20.02 ± 0.51 µm; main width 1.28 ± 0.25) slightly curved or S-shaped, sub-terminal kinetoplast (3.83 ± 0.76 µm from posterior end, arrows Fig. 2a-b), a thick flagellum edging the body of the parasite, with or without very short free flagellum. When these parasites are transferred from 27 to 37°C and maintained in MEMTAU medium for 1, 2 or 3 days, they do not transform to amastigotes, but morphological alteration were observed in the trypomastigote morphology (Fig. 2c-h). During the first day of pre-incubation, almost all of the metacyclic trypomastigotes display a slight lengthening and thinning (total length 21.41 ± 4.99 µm; main width 1.09 ± 0.26), showing a small-sized kinetoplast usually very far from its pointed posterior end (4.95 ± 1.06 µm from posterior end, arrows Fig. 2c-d), and a poorly-developed undulating membrane. A similar morphological pattern was observed on the second day of pre-incubation, though the total length and width of the parasites was reduced (16.51 ± 3.44 and 0.97 ± 0.17 µm, respectively) having a variable kinetoplast sub-terminal (5.13 ± 1.29 µm from posterior end, large arrows Fig. 2e-f). This morphology is notably kept in parasites pre-incubated by three day, except by a progressive shortening of the entire organism (total length 12.03 ± 3.03 µm; main width 0.65 ± 0.13), with an inconspicuous undulating membrane (Fig. 2g-h).

Although notable cellular rearrangements were not observed during the pre-incubation phase, the metacyclic trypomastigotes exhibited a lengthening followed by a shortening of the body as the pre-incubation time progressed (Fig. 2c-h). This enlargement can be seen to initiate by a sharpening at either the posterior or anterior end of the parasites (Fig. 2c-d, short arrows), while the shortening process seems to occur preferentially by a sloughing off of the surface membrane from the posterior end of the parasite (Fig. 2e, g, h, arrowheads). This process is accompanied by a continuous loss of surface membrane, the lengthening at the posterior end and the slenderness of the metacyclic trypomastigote along with the release of a barely-visible material (Fig. 2c, f, short arrows). Moreover, small flagellar swellings, protruding from the flagellum at different levels and membrane blobbing were seen in some instances along the entire surface of the parasites (Fig. 2e, short arrows).

When MEMTAU-purified metacyclic trypomastigotes were re-incubated in fresh MEMTAU medium at 37°C, an accelerated appearance of forms resembling amastigotes occurs in the subsequent 72 h (Fig. 3). This phase of the transformation process was not synchronous, but the decrease in the proportion of metacyclic trypomastigotes was compensated by an increase of amastigotes and a decrease in the proportion of amastigotes (5.13 ± 1.29 µm from posterior end, arrows Fig. 2e-f). This morphology is notably kept in parasites pre-incubated by three day, except by a progressive shortening of the entire organism (total length 12.03 ± 3.03 µm; main width 0.65 ± 0.13), with an inconspicuous undulating membrane (Fig. 2g-h).

The comparison of IMA isolated by Metrizamide gradient (Fig. 3i-j) with their corresponding EMA shows a notable gross morphological similarity, yet display differences on their membrane (Fig. 3g-h). While, the EMA exhibit a
rough surface membrane and longitudinal cracks, the IMA present a smooth, granulose and slanting crack on their surfaces. However, both type of amastigotes are rounded or oval-shaped and have a short protruding flagellum.

Protein and glycoprotein's patterns from EMA and ETA - Total protein patterns of the two amastigote types were visualized by Coomassie Blue-Silver as shown in Fig. 4A. As can be seen, ETA and EMA show polypeptide patterns of different molecular weight and intensity, even though both profiles appear to share many of the detected bands (Fig. 4A). In contrast, the glycopeptide patterns obtained by PAABGS of EMA and ETA are al-

Fig. 1: scanning electron micrograph of Trypanosoma cruzi during the extracellular transformation of trypomastigotes to amastigote in MEMTAU medium at 37°C. Tissue culture trypomastigotes from the supernatant of Vero monolayer (a) before of the incubation in acidic medium present variable morphologies. After 3 (b, c), 6 (d, e), 9 (f), 12 (g), 18 (h) and 24 h (i, j) of incubation an accelerated cellular rearrangement can be observed. Arrow in a indicate a terminal kinetoplast (k). Morphological changes were accompanied by release of trails like-bead associated with the parasite (short arrows in d, f, g, j, k, l). Short arrows in f point to other type of trails being released at anterior end and the flagellum tip of the parasite. Short arrow in k and l indicate different type of trails being released by intracellular trypomastigote-derived amastigotes obtained by disruption of infected Vero cells.
most indistinguishable from each other (Fig. 4B), with ETA displaying higher glycopeptides complexity (lane ETA vs lane EMA, Fig. 4B). These results indicate that ETA and EMA are very similar in their glycoprotein content but differ substantially in their protein profile.

DISCUSSION

Trypanosomes are organisms in which shape is dictated, at least in part, by the subpellicular array of microtubules that are cross-linked to each other and to the plasma membrane through MAPs (Gull 1999). It has been
emphasized that the extreme morphological and motility differences of the trypomastigote and amastigotes stages of \textit{T. cruzi} offer an excellent chance to study microtubules-based mechanism of morphological control which would be difficult to discern in systems where such change is more subtle (Gull 1999, Tyler & Engman 2001). We have shown by SEM that in axenic conditions the morphological transformation of the tissue-cultured trypomastigotes and the in vitro-induced amastigotes from metacyclic trypomastigotes are different. The comparison of extracellular and intracellular amastigotes morphology showed that they share many similarities but are not mor-

Fig. 3: scanning electron micrograph of \textit{Trypanosoma cruzi} during the extracellular transformation of metacyclic to amastigote. When metacyclic trypomastigotes are re-incubated in fresh MEMTAU medium at 37°C for 8 (a, b), 24 (c, d), 48 (e, f) or 72 h (g, h) a notable cellular rearrangement can be observed. Large arrow in a indicate a prominent subterminal kinetoplast (k). The morphological changes are accompanied by release of different type of trails associated with the parasite (short arrows). Short arrows in c, e and f point to different type of trails being released at either the posterior or anterior end of the parasites. Short arrows in i and j point to different type of trails being released by intracellular metacyclic-derived amastigotes obtained by disruption of infected Vero cells.
phologically identical. The comparison of protein and glycoprotein profiles by SDS-PAGE analyses also uncovered differences between the two types of amastigotes, especially among their respective set of proteins. Therefore, our work suggests that the morphological transformations of *T. cruzi* may serve as a robust model system to study the molecular mechanisms of morphological control.

Our results show that the pH-induced transformation of tissue-derived trypomastigotes previously reported by Tomlinson et al. (1995) was also observed using a different acidic medium. The parasites transformed into amastigotes at 18 to 24 h after incubation in MEMTAU medium at 37°C indicating a faster differentiation rate than previously reported (Andrews et al. 1987, Tomlinson et al. 1995). However, the transformation process of trypanosomastigotes adhered to the morphological patterns described by Andrews et al. (1987). It is very likely that the accelerated morphological changes observed must be accompanied by antigenic changes, since we could observe by Western blots with stage-specific antisera changes in antigen profiles (unpublished results). An advantage of our system is that the elapsed time is shorter and the transformation process is highly synchronized.

Recently, we have also shown by light microscopy, SDS-PAGE and Western blots that pre-incubation at 37°C of metacyclic trypomastigotes in MEMTAU medium triggered, 24 h after incubation, early transformation events such as an accelerated macromolecular differentiation to amastigotes without noticeable morphological changes. Late transformation events which occurred when the culture medium is changed, however, were characterized by profound morphological rearrangements with few molecular changes (Contreras et al. 2002). The data presented here provide a more detailed morphological analysis of the parasites during the pre-incubation and re-incubation phases. The images indicate that within 24 h after incubation, the metacyclic trypomastigotes exhibit a lengthening followed by a shortening of the body as the pre-incubation time progresses. We postulate that the apparent enlargement of the parasite may be the result of a progressive shedding of its posterior surface as a compaction process probably driven by changes in the rate of shortening of microtubules and which is faster than the compaction of the cellular membrane occurs (Karp 1999). Such mechanism may explain the loss of massive surface membrane observed right before and during the start of the rounding process as breaking during this transition was often observed. Concomitantly, a release of fine trails along the entire surface of the parasite was observed which could be associated with the sequential expression and loss of specific membrane antigens as previously reported by other authors (Andrews et al. 1987, Zeledon et al. 1988, Barros et al. 1996).

The observation that the first morphological events induced by pre-incubation are a transient lengthening of the posterior end of the metacyclic form, an apparent shift of the kinetoplast, and a thinning of the parasite body resulting from a slight unfolding of the undulating membrane, seem to indicate that an extensive cytoskeleton remodeling associated with high turnover of surface membrane may be taking place. It is conceivable that MAPs which connect the subpellicular microtubules to the plasma membrane may participate in this process (Gull 1999). Whether the early morphological changes and the “escaping” of the metacyclic trypomastigote from endocytic

![Fig. 4: SDS-10% PAGE analysis of extracellular amastigotes of *Trypanosoma cruzi*. The gels (6 µg/well) were stained with a combined Coomassie Blue-Silver stain procedure for proteins (A) or with a combined Periodic Acid-Acian Blue-Gluteraldehyde-Silver (PAABG) stain procedure for glycoproteins (B) as described into Materials and Methods. EMA corresponds to extracellular metacyclic-derived amastigotes and ETA to extracellular trypomastigote-derived amastigotes. Lanes m shows molecular weight markers and s a mixture of glycoproteins (6 µg) for controlling staining procedure. The numbers on the left side correspond to relative mobility (Mr) of the markers in kilodaltons (kDa).]
vacuoles are causally related is unknown. It is noteworthy, however, that the transformation process can be continued and completed only after renewed the incubation medium. Assays to examine whether the product release from the parasite during pre-incubation phase, are involved in the inhibition of the morphological transformation are in progress and might be useful to explain why the metacyclic trypomastigote can also escape from endocytic vacuole into the host cell cytoplasm (Dvorak 1975).

Tyler and Engman (2000) have shown that the depletion of glucose from the medium triggers elongation of the epimastigotes flagellum. Later, the authors proposed that the intracellular glucose concentration is the key factor for both elongation of the flagellum and sustaining the differentiation of the metacyclic trypomastigote (Tyler & Engman 2001). In agreement with these data, we observed that the thickening or shortening process exhibited during the pre-incubation phase is not accompanied by elongation of the metacyclic trypomastigote flagellum. On the contrary, a thick and very short free flagellum edging the body of the parasite was noted, as expected for incubation of parasites in a glucose-rich medium.

The most striking cellular rearrangement of the parasites occurred when the culture medium was changed. The sudden shortening and the blunting of the posterior end observed in the metacyclic trypomastigote within the first 8 h after re-incubation provides clues as to the dynamics of posterior region release during amastigogenesis. The subsequent morphological changes involving thickening of the posterior end and progressive shortening of the parasite around itself in a helical fashion, along with a gradual internalization of the flagellum resemble the changes for tissue-culture-derived trypomastigotes as described above. The fact that early and late morphological changes are associated with the different environmental conditions surrounding extracellular primary amastigogenesis may provide an opportunity to study the role of ubiquitin (UB)-proteosome system, which was recently shown to be involved in the remodeling process of trypomastigote transformation (De Diego et al. 2001). It is worth noting that the early events described during primary amastigogenesis were not observed when tissue-culture-derived trypomastigotes transformed into amastigotes. These morphological differences may indicate that the metacyclic form possesses a specific way of responding and adapting to environmental changes that is distinct from that shown by bloodstream trypomastigotes.

It has been reported that while the transformation of trypomastigotes is triggered by Phosphatidylinositol-Specific Phospholipase C (PI-PLC), the metacyclic forms are not affected (Mortara et al. 2001). In addition, the pH-induced transformation of tissue-derived trypomastigotes is not observed when metacyclic trypomastigotes are incubated under the same conditions (Tyler & Engman 2001). Interestingly, it is known that the first intracellular cycle of metacyclic trypomastigotes is significantly shorter than the cycle of tissue-culture-derived trypomastigotes of the same strain that has been growing as an obligate intracellular parasite for several cycles (Dvorak 1975). These differences may be explained by the fact that even though ETA and EMA may appear to be undistinguishable at the gross morphological level, they nevertheless display noticeable differences at the ultrastructural and molecular levels. For instance, under the scanning electron microscope, ETA exhibit a finely granular surface while EMA present a rough membrane surface. Furthermore, when their polypeptide and glycopeptides profiles were compared, we detected significantly different protein profiles between the two forms of amastigotes, suggesting that the ultrastructural differences found in ETA and EMA are underscored by differences in macromolecular constitution (Ruiz et al. 1993, Burleigh & Andrews 1995).

It is noteworthy that the intracellular amastigotes showed similar gross morphology, yet their crack patterns were different. Whether or not such difference is related to distinct biological properties is unknown, but may reflect differences in their development. Analogy can be made with works on human plasmodia (Aikawa & Atkinson 1990, Rey 1991, Bannister et al. 2000), where it has been described that the processes of shizogony occurring in the vertebrate host yield exoerythrocytic merozoites and erythrocyte merozoites with similar biological capacities, but with partly different morphologies and antigenicities. Thus, it is tempting to speculate that the IMA may be the parasitological equivalent to exoerythrocytic merozoites derived from infective sporozoites transmitted by the vector insect, while the ITA may be equivalent to the erythrocyte merozoites. If so, one would expect that primary and secondary amastigogenesis in the vertebrate host will be different as well.

In conclusion, this paper reports on the first morphological comparison of the developmental processes giving rise to in vitro amastigotes from trypomastigote and metacyclic forms. Although it is clear that axenic amastigogenesis cannot mimic intracellular amastigogenesis in absolute terms, the data presented here demonstrates the utility of our experimental system as a model in which to begin the molecular study of parasite development in vivo. The ultrastructural and molecular differences observed between seemingly identical amastigotes (ETA and EMA) intimate that the mechanisms controlling primary and secondary intracellular amastigogenesis in T. cruzi may be different and should be investigated further.

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