Molecular mechanisms of *Trypanosoma cruzi* infection by oral route

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Frequent reports on outbreaks of acute Chagas’ disease by ingestion of food contaminated with parasites from triatomine insects illustrate the importance of this mode of transmission. Studies on oral Trypanosoma cruzi infection in mice have indicated that metacyclic trypomastigotes invade the gastric mucosal epithelium. A key molecule in this process is gp82, a stage-specific surface glycoprotein that binds to both gastric mucin and to target epithelial cells. By triggering Ca\(^{2+}\) signalling, gp82 promotes parasite internalisation. Gp82 is relatively resistant to peptic digestion at acidic pH, thus preserving the properties critical for oral infection. The infection process is also influenced by gp90, a metacyclic stage-specific molecule that negatively regulates the invasion process. *T. cruzi* strains expressing high gp90 levels invade cells poorly in vitro. However, their infectivity by oral route varies considerably due to varying susceptibilities of different gp90 isoforms to peptic digestion. Parasites expressing pepsin-susceptible gp90 become highly invasive against target cells upon contact with gastric juice. Such is the case of a *T. cruzi* isolate from an acute case of orally acquired Chagas’ disease; the gp90 from this strain is extensively degraded upon short period of parasite permanence in the gastric milieu. If such an exacerbation of infectivity occurs in humans, it may be responsible for the severity of Chagas’ disease reported in outbreaks of oral infection.

Key words: *Trypanosoma cruzi* - oral infection - metacyclic trypomastigotes - surface molecules - gp82 - gp90

In the past 20 years transmission of *Trypanosoma cruzi*, the causative agent of Chagas’ disease, has steeply declined in Southern Cone primarily due to control measures directed at the elimination of the domiciliary vector, *Triatoma infestans*. Notwithstanding the successful elimination of *T. cruzi* transmission by *T. infestans* in Brazil, Uruguay, Chile and substantial areas of Argentina, Bolivia and Paraguay (Schofield et al. 2006), many challenges remain. Increases in transmission have been reported in some areas of Argentina (Gürtler et al. 2005), Venezuela (Feliciangeli et al. 2003) and in the Brazilian Amazon (Coura 2006). Several countries, including Mexico, Peru, Colombia and Costa Rica, have not implemented a national programme for the control of *T. cruzi* transmission (Schofield et al. 2006). A matter of concern is also the report that wild populations of *T. infestans* are much more widespread throughout Bolivia than predicted, raising the possibility of a threat of reinfestations of treated areas and the eventual spread to neighbouring regions (Noireau et al. 2005).

*T. cruzi* acquisition via the oral route has recently been the focus of attention. In Brazil, after the control of *T. cruzi* transmission through *T. infestans* and by blood transfusion, the most important and frequent mode of transmission is by the oral route (Coura 2006). Outbreaks of acute Chagas’ disease through food contamination have occurred in distinct regions of the country in the last four years. The occurrences have been more prevalent in the Amazon Region, where triatomines other than *T. infestans* predominate. According to Coura et al. (2002), more than 50% of acute cases of Chagas’ disease registered in the Brazilian Amazon between 1968-2000 were attributable to microepidemics of orally transmitted infection.

Studies in mice have shown that insect-derived metacyclic trypomastigotes invade the gastric mucosal epithelium where the parasite replicates as amastigotes; no evidence of parasitic invasion anywhere within the oropharynx or oesophagus has been found (Hoft et al. 1996). Metacyclic forms express mucin-like surface glycoproteins that are highly resistant to proteolytic degradation (Mortara et al. 1992) and enable them to resist the harsh conditions in the stomach. This resistance to proteolysis of the mucins allows the metacyclic forms to proceed towards the target cells located in the gastric mucosa (Yoshida 2006), which constitute the unique portal of entry to systemic infection (Hoft 1996). Contact with gastric juice does not diminish the parasite’s infectivity. In fact, some *T. cruzi* strains may become more invasive upon transient residence in the gastric milieu (Cortez et al. 2006b, Covarrubias et al. 2007). This report summarises recent findings on the molecular mechanisms underlying the process of gastric mucosal infection by metacyclic trypomastigotes in a mouse model. Focused are also the questions of the modulation of *T. cruzi* infectivity by stomach components, which may either function as barriers to progression towards target cells or may act on parasites to render them more competent for invasion. This potential of stomach components to promote inva-
Mechanisms of oral infection, Chagas’ disease in orally acquired infection.

**Metacyclic stage-specific surface molecule gp82 promotes *T. cruzi* infection by oral route**

Experiments on oral infection in mice using metacyclic trypomastigotes generated in culture have indicated that the stage-specific surface glycoprotein gp82 plays a key role in the invasion of the gastric mucosal epithelium (Neira et al. 2003). Gp82 of different *T. cruzi* strains resists degradation by pepsin at acidic pH, so that the parasites recovered from the stomach 1 h after oral inoculation into mice, preserve full infectivity (Cortez et al. 2006b). The process of gp82-dependent host cell invasion by metacyclic forms has been partially elucidated by in vitro studies with human epithelial HeLa cells. Parasite-host cell interaction mediated by gp82 triggers in both cells signal transduction pathways that lead to intracellular Ca\(^{2+}\) mobilisation (Ruiz et al. 1998), an event essential for *T. cruzi* internalisation (Docampo & Moreno 1996, Burleigh & Andrews 1998, Yoshiha 2006). In the parasites, the Ca\(^{2+}\) response is induced upon recognition of gp82 by an as yet undefined host cell receptor expressed in mammalian cells that are susceptible to *T. cruzi* infection, such as HeLa and Vero cells, but not in *T. cruzi*-resistant cells such as the human leukaemia K562 cells (Yoshida et al. 2000). The events upstream of Ca\(^{2+}\) mobilization appear to involve the activation of the protein tyrosine kinase and phosphorylation of p175, a protein that is undetectable in noninfected epimastigotes (Favoreto et al. 1998) and activation of phospholipase C. These pathways lead to the generation of inositol 1,4,5-triphosphate (IP\(_3\)), which promotes Ca\(^{2+}\) release from IP\(_3\)-sensitive stores (Yoshida et al. 2000). In host cells, phosphoinositide-3 kinase and protein kinase C appear to be part of the signalling cascades that culminate in Ca\(^{2+}\) mobilization from thapsigargin-susceptible compartments (Ferreira et al. 2006). Ca\(^{2+}\)-dependent disorganization of the target cell actin cytoskeleton results from gp82-mediated parasite entry (Cortez et al. 2006a). This may contribute to efficient infection, provided that microfilament rearrangement facilitates *T. cruzi* internalisation (Rodriguez et al. 1995). Several pieces of evidence indicate that bidirectional signalling triggered by gp82 ensures the most effective target cell invasion. Metacyclic forms of clone CL-14, for instance, which express gp82 at low levels but otherwise exhibit a surface profile indistinguishable from the highly invasive parental CL strain, invade cultured HeLa cells poorly (Atayde et al. 2004).

Gp82 is encoded by a multigene family that is part of the trans-sialidase/gp85 superfamily, which is expressed at the infective trypomastigote stage (Araya et al. 1994). It is a glycoprotein that contains N-linked oligosaccharides (Ramirez et al. 1993); however, the carbohydrate portion of the molecule is not involved in host cell binding. Therefore, the recombinant protein based on gp82 generated in bacteria has the same cell binding capacity as its endogenous counterpart (Ruiz et al. 1998). The gp82 molecule that has been identified by the monoclonal antibody (MAb) 3F6, which inhibits infection in vitro and in vivo when parasites are orally administered, is highly conserved between genetically divergent *T. cruzi* strains. Analysis of the amino acid sequences of gp82, as deduced from cDNA clones of G and CL strain metacyclic forms, has revealed an overall identity of 97.9% and of 100% within the central domain containing the target cell binding site (Ruiz 1998). At the carboxy-terminal end of the central domain, the cell binding site for gp82 is contiguous to, and partially overlapping, the epitope for MAb 3F6 (Manque et al. 2000). In addition to the cell invasion promoting activity, gp82 also has the ability to bind to gastric mucin.

Gastric epithelia are protected by an extracellular mucus layer whose major components are the high molecular mass mucins. During oral *T. cruzi* infection, the metacyclic trypomastigotes, upon reaching the stomach, bind to gastric mucin through gp82 as a first step towards their destination, i.e., the mucosal epithelial cells. There have been reports that suggest that gp82 binding is selective to gastric mucin and not, for instance, to mucin from submaxillary glands. This specificity may explain why, in orally inoculated mice, parasites are not found in the oropharynx or oesophagus (Hoft et al. 1996). Such selectivity may determine the stomach cells as the portal of *T. cruzi* entry similar to the preferential adherence to colonic mucin of *Shigella dysenteriae*, whose pathogenic potential is correlated with its ability to invade and multiply within the cells of colonic epithelium (Sudha et al. 2001). The importance of the gastric mucin-binding property of gp82 in the establishment of *T. cruzi* infection by the oral route has been confirmed by studies that have used parasite strains that were deficient in the expression of gp82. Metacyclic forms of gp82-deficient strains express the surface glycoprotein gp30 which induces Ca\(^{2+}\) signalling in host cells and is also recognized by MAb 3F6 (Cortez et al. 2003). The lack of gp82 expression does not affect the parasite’s capacity to enter host cells in vitro, presumably because gp30 can fulfil the invasion-promoting function of gp82. Since the invasion of the target cell by gp82-deficient parasites is inhibited to the same extent by either purified native gp30 or by the recombinant gp82, it suggests that the same receptor may be recognizing these molecular species (Cortez et al. 2003).

While the in vitro infectivity of gp82-deficient *T. cruzi* strains is similar to that of the gp82-expressing CL strain, the course of infection upon oral administration in mice differs markedly. Gp82-deficient parasites produce very low parasitaemia levels, which may be attributed to the poor affinity of gp30 for gastric mucin (Cortez et al. 2003). Studies that have used systems that mimic the in vivo conditions have demonstrated that poor adhesion to gastric mucin can interfere with parasite entry into subjacent epithelial cells. In the presence of gastric mucin, penetration of gp82-deficient metacyclic trypomastigotes into HeLa cells was significantly reduced while the infectivity of gp82 expressing strains remained unaltered (Cortez et al. 2003).

Based on the findings from experiments in mice and in cultured epithelial cells, using CL and gp82-deficient strains, the following picture is envisaged (Fig. 1). Upon oral inoculation, metacyclic forms reach the stomach lu-
men, where the gp82-expressing parasites bind to gastric mucin as an initial step for translocation towards the target epithelial cells; this process appears to require parasite energy. For metacyclic forms of gp82-deficient \textit{T. cruzi} strains, which adhere, albeit poorly, to gastric mucin in gp30-mediated fashion, the mucus layer constitutes a barrier that prevents most parasites from reaching the mucosal epithelium. Following the initial binding, the process of cell invasion is initiated (Fig. 2). Metacyclic forms that express gp82 engage this molecule to accomplish the internalisation process by triggering the signalling cascades that lead to an increase in intracellular Ca\textsuperscript{2+} concentration. Gp82-deficient metacyclic forms that manage to traverse the mucus layer enter host cells in a gp30-mediated manner, thus inducing Ca\textsuperscript{2+} mobilization. After \~12 h post-invasion, most metacyclic forms have transformed into amastigotes, which go through replication cycles. By 96 h, nests of amastigotes are detectable at much higher numbers in the histological preparations of the stomach of mice infected with gp82-deficient parasites than in sections from animals that were inoculated with gp82-expressing strains (Fig. 3). There is no difference between strains with regards to the rate of intracellular multiplication so that the release of trypanomastigotes into the circulation should occur within a comparable time frame. However, reflecting the difference in the rate of invasion, parasitaemia levels in mice infected with gp82-deficient parasites are much lower than in mice inoculated with gp82-expressing parasites.

It should be noted that the CL strain and the strains deficient in gp82 expression referred in this report express gp90 molecule at low levels (Cortez et al. 2003). Therefore their infectivity is not influenced by this molecule. Gp90 is a metacyclic, stage-specific surface glycoprotein (Teixeira & Yoshida 1986) that is ubiquitously present in the \textit{T. cruzi} population (Yoshida 2006). This surface molecule binds to mammalian cells in a receptor-mediated manner without inducing Ca\textsuperscript{2+} response (Ruiz et al. 1998) and therefore acts as a negative regulator of target cell invasion (Málag & Yoshida 2001).

\textbf{Susceptibility of gp90 molecule to proteolysis influences \textit{T. cruzi} infectivity by oral route}

Expression of gp90, the metacyclic stage-specific surface glycoprotein that down modulates parasite infectivity (Málag & Yoshida 2001), varies in different \textit{T. cruzi} strains and its expression in a strain is inversely correlated with the ability of that strain to invade cul-

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**Fig. 1:** interaction of \textit{T. cruzi} with gastric mucin in oral infection. Upon inoculation into mice (A), gp82-expressing and gp82-deficient metacyclic trypomastigotes reach the stomach lumen (B). Parasites expressing gp82 bind to gastric mucin in the mucous layer by engaging this surface molecule, as a first step towards underlying epithelial cells. By contrast, a few gp82-deficient parasites bind to gastric mucin, possibly using gp30.

**Fig. 2:** penetration of \textit{Trypanosoma cruzi} into gastric epithelial cells in oral infection. Upon reaching the target cells, gp82-expressing metacyclic forms accomplish the invasion process by triggering the gp82-mediated signaling cascades that lead to an increase in intracellular Ca\textsuperscript{2+} concentration. Metacyclic forms of gp82-deficient \textit{T. cruzi} strains that managed to traverse the mucus layer enter host cells in gp30-mediated manner, inducing Ca\textsuperscript{2+} mobilization. The upper part of the figure is a composition of a histological section of mouse stomach stained by hematoxylin-eosin and Giemsa-stained metacyclic trypanomastigotes that are not in the same scale.
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Two variant forms of gp90 have been identified based on the reactivity with MAbs 1G7 and/or 5E7 (Teixeira & Yoshida 1986). The complete peptide sequence of either of these gp90 molecules remains to be determined. Attempts to clone the gene encoding gp90 have, so far, been unsuccessful. What is known is that the sequence is present in multiple copies in the genome, encodes a gp90 carboxy-terminal portion containing the epitope for MAb 5E7 (Franco et al. 1993), an antibody that reacts with metacyclic forms of all *T. cruzi* strains examined to date (Yoshida 2006). The recombinant protein corresponding to the carboxy-terminal domain of gp90 binds to target cells in a receptor-mediated manner that is indistinguishable from that of native gp90 (Ruiz et al. 1998). However, the receptors for the recombinant and the native molecules may be distinct. MAb 1G7, which does not react with the recombinant protein, binds to live metacyclic trypomastigotes and inhibits parasite-induced intracellular Ca\(^{2+}\) mobilization (Ruiz et al. 1998). Thus, the gp90-reactive MAb 1G7 has an opposing effect to that of gp82-reactive MAb 3F6, which triggers an increase in cytosolic Ca\(^{2+}\) concentration upon binding to metacyclic forms (Ruiz et al. 1998). Of interest is that while MAb 3F6 induces the activation of the protein tyrosine kinase and phosphorylation of 175, (Favoreto et al. 1998), MAb 1G7 promotes p175 dephosphorylation (Manque et al. 2003). These findings are compatible with the idea that activation of protein tyrosine kinases and intracellular Ca\(^{2+}\) mobilization are associated events (Yoshida et al. 2000).

Metacyclic forms of *T. cruzi* strains expressing gp90 at high levels invade cultured cells poorly. However, it is not possible to predict from this in vitro behaviour the course of infection after oral administration into mice. Recent studies have indicated that in vivo infectivity of these parasites is determined by gp90 susceptibility to peptic digestion. This notion is illustrated in the case of two *T. cruzi* strains that were derived from patients during the acute phase of the disease (Fig. 4). Metacyclic forms of these strains, which exhibited a similar surface profile (with high expression of gp82, gp30 and gp90), entered cultured epithelial cells at low numbers (Cortez et al. 2006b). This provides evidence that gp90-mediated parasite-host cell interactions predominate over those interactions that are mediated by the gp82 or gp30 molecules. When orally inoculated into mice, the two strains exhibited distinct infective properties. One of these strains entered gastric mucosal epithelial

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**Fig. 3:** intracellular replication of *Trypanosoma cruzi* in gastric mucosal epithelium. Following oral inoculation of metacyclic trypomastigotes into mice, the parasites invade target cells in the stomach and replicate as amastigotes. Shown are the histological sections of the mouse stomach stained by hematoxylin-eosin four days post-infection. Nests of replicating amastigotes (white arrows) can be seen in higher numbers in mice infected with gp82-expressing parasites (A) than in animal inoculated with gp82-deficient strains (B).

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**Fig. 4:** increased infectivity of *Trypanosoma cruzi* strains expressing gp90 molecules susceptible to peptic digestion. Upon oral infection, metacyclic trypomastigotes of parasite strains bearing pepsin-susceptible gp90\(^*\) molecules express much less of these cell invasion down modulators, so that they can enter target cells more effectively. By contrast, parasites bearing pepsin-resistant gp90 molecules bind to target cells and relay inhibitory signal to both cells, thus hampering parasite entry.
cells in high numbers and, as a consequence, resulted in a high number of replicating parasites in amastigote nests by day 4 as determined by haematoxylin-eosin stained histological sections of the stomach. At later times, the infected mice developed high parasitaemias and about 30% of them died within 30 days of infection. By analysing the parasites recovered from the mouse stomach 1 h after oral infection, the gp90 molecule was found to be extensively degraded, in a manner similar to in vitro treatment of parasites with pepsin at acidic pH. Metacyclic forms with digested gp90, either upon transient residence in the gastric milieu or upon pepsin treatment in vitro, displayed an increased capacity to enter target cells (Cortez et al. 2006b). In contrast with these observations, mice inoculated with the second T. cruzi strain developed very low parasitaemias. Consistent with this, at day 4 post-inoculation the number of amastigote nests found in the stomach was very low and mortality was null. Analysis of parasites recovered from the mouse stomach 1 h after oral infection showed that the gp90 molecule remained intact, thus explaining why metacyclic forms of this strain are as poorly infective in oral infection as they are towards cultured epithelial cells. It should be noted that, apart from the differential susceptibility of gp90 to peptic digestion, the two T. cruzi strains did not exhibit marked differences, with both strains preserving fully functional gp82 and gp30 molecules after being recovered from the gastric milieu (Cortez et al. 2006b).

Overall, these findings suggest that the severity of acute Chagas’ disease that has been reported in patients infected by the oral route may be associated with up-regulation of parasite infectivity, due to digestion of the cell invasion inhibitor, gp90. Support for this hypothesis has been provided by the first study of a T. cruzi isolate derived from an acute case of Chagas’ disease acquired by oral route. This isolate, from an outbreak of oral T. cruzi infection that occurred in the southern state of Santa Catarina in 2005 and resulted in three deaths, was designated SC. Metacyclic trypomastigotes of SC, characterized by high levels of expression of gp82, gp30 and gp90, entered host cells in culture at low rates (Covarrubias et al. 2007). When inoculated orally into mice, however, the parasites produced high parasitaemias and high mortality with more than 50% of infected animals being dead by day 20 post-inoculation. Analysis of SC parasites recovered from the mouse stomach 1 h after oral administration failed to detect the presence of gp90 molecule, indicating that it was completely digested upon contact with the gastric juice. However, while the amino-terminal portion of gp82 was digested, the domain implicated in gastric mucin-binding and host cell invasion was fully preserved. With a fully functional gp82, in addition to a pepsin-resistant gp30, the internalisation of the parasite increased.

Do T. cruzi gp35/50 molecules play a role in oral infection?

T. cruzi strains referred to in preceding sections, derived mostly from chagasic patients, rely to a certain extent on gp82 or gp30 to establish infection in mice by oral route. For these strains, mucin-like surface glycoproteins primarily serve to protect the parasites against the action of gastric juice. However, gp35/50 mucins have an additional role in T. cruzi strains that preferentially engage these molecules to invade target cells in vitro. Among T. cruzi strains examined to date, two groups have emerged based on the expression of mucin-like gp35/50 glycoproteins: those reacting solely with MAb 2B10 and those reacting with both MAb 2B10 and MAb 10D8. Strains from human origin that have been studied so far express gp35/50 molecules reactive with MAb 2B10 but not with MAb 10D8, whereas most strains derived from wild animals or wild triatomine insects express MAb 10D8-reactive gp35/50 (Mortara et al. 1992). The gp35/50 mucins are encoded by a large multigene family (Di Noia et al. 1998) and are expressed in metacyclic forms as well as in epimastigotes (Yoshida et al. 1989). These mucins are GPI-anchored glycoproteins rich in threonine residues. In the protein core, glycans are O-linked to the threonine residues through N-acetylglucosamine rather than N-acetylgalactosamines as usually found in vertebrate mucins. (Schenkman et al. 1993). Depending on the T. cruzi strain, they may contain galactofuranose residues in addition to galactopyranose (Prevato et al. 1994, Acosta-Serrano et al. 1995, Salto et al. 2000). MAb 10D8 reacts with gp35/50 epitopes containing galactofuranose, whereas MAb 2B10 appears to recognize galactopyranose-containing epitopes present in all strains. Several lines of evidence have indicated that MAb 10D8-reactive gp35/50 is implicated in host cell invasion as both the antibody and the purified mucin are capable of inhibiting parasite internalisation (Yoshida et al. 1989, Ruiz et al. 1993). Gp35/50 glycoprotein binds to target cells in a receptor-mediated manner and induces a bidirectional Ca\(^{2+}\) response, but to a lesser degree than gp82 (Ruiz et al. 1998). It is possible that most parasite strains that rely preferentially on gp35/50 to enter host cells are poorly invasive because of the relatively weak signalling activity of these molecules. However, the low invasive capacity may also be due to the expression of gp90, which in general is present at high levels in metacyclic forms of T. cruzi strains that are recognized by MAb 10D8 (Yoshida 2006). As these strains also exhibit high expression of gp82, the results support the hypothesis that the gp90-mediated interaction with the host cell is dominant acting. Gp35/50 mucins contain sialic acid but sialyl residues are not required for invasion and, depending on the parasite strain, they may even impair recognition by target cells (Yoshida et al. 1997). For instance, treatment of metacyclic forms of G strain with bacterial neuraminidase removes gp35/50 sialic acid and increases the parasite infectivity (Yoshida et al. 1997). This can be reverted by resialylation of gp35/50 upon incubating the parasites with T. cruzi trans-sialidase and sialyl lactose, which results in parasite internalisation to similar levels as that seen before desialylation. Consistent with this result, the ability of desialylated gp35/50 to bind to target cells and to trigger Ca\(^{2+}\) response was found to be higher than that of its sialylated counterpart (Yoshida et al. 1997).

G strain metacyclic forms that engage MAb 10D8-reactive gp35/550 mucins to enter host cells in vitro are
poorly infective when given orally into mice and fail to produce detectable parasitaemia upon microscopic examination of blood samples (Yoshida 2006). Analysis of histological preparations of stomach from mice infected with G strain has revealed very scarce parasites. These observations are consistent with the fact that this strain expresses high levels of a gp90 isoform resistant to pepsin digestion (Covarrubias et al. 2007). If metacyclic forms expressing 10D8-reactive gp35/50, but not gp90, could be generated, the role of T. cruzi mucins in infection in vivo may be clarified. Finding proteases and/or conditions that preserve the gp35/50 molecules and degrade gp90 would facilitate researchers to gain insight into the role that mucins play in infection.

Studies on experimental oral infection by T. cruzi metacyclic trypomastigotes, in parallel with in vitro epithelial cell invasion assays, have revealed several features of parasite-host interactions, which would not have been predicted solely based on in vitro experiments. The gastric mucosal epithelium as the preferential portal of entry for metacyclic forms inoculated in mice by the oral route may be due to the selective binding of parasites to the gastric mucin. In the gastric milieu, metacyclic forms resist digestion due to the presence of surface mucin-like gp35/50 glycoproteins. Metacyclic stage-specific surface molecule gp82, implicated in gastric mucin-binding and in promoting host cell invasion, is relatively resistant to pepsin digestion and remains fully functional upon transient contact with gastric juice. Gp90, a metacyclic stage-specific surface molecule that functions as a negative regulator of target cell entry and is the main determinant of parasite infectivity, has strain-dependent susceptibility to proteolysis. Metacyclic forms expressing gp82 and gp90 at low levels traverse the mucus layer protecting the gastric mucosa through gp82-mediated binding to its major components, the mucin molecules, and efficiently invade the target epithelial cells engaging gp82. For metacyclic forms deficient in gp82, the mucous layer represents a barrier that hampers the journey towards host cells. For T. cruzi strains that express the cell invasion-promoting molecules gp82 and gp30, in addition to gp90 at high levels, their infectivity will be determined by the susceptibility of gp90 isoform to pepsin digestion. If they express pepsin-resistant gp90, they will be poorly infective. However, if they express pepsin-susceptible gp90 molecules, they can become highly infective upon exposure to gastric juice. The latter situation appears to be the case in a T. cruzi isolate from an acute case of Chagas’ disease acquired by oral infection, which had its infectivity enhanced in the mouse gastric milieu.

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