

Trypanosoma cruzi: adaptation to its vectors and its hosts

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Abstract – American trypanosomiasis is a parasitic zoonosis that occurs throughout Latin America. The etiological agent, *Trypanosoma cruzi*, is able to infect almost all tissues of its mammalian hosts and spreads in the environment in multifarious transmission cycles that may or not be connected. This biological plasticity, which is probably the result of the considerable heterogeneity of the taxon, exemplifies a successful adaptation of a parasite resulting in distinct outcomes of infection and a complex epidemiological pattern. In the 1990s, most endemic countries strengthened national control programs to interrupt the transmission of this parasite to humans. However, many obstacles remain to the effective control of the disease. Current knowledge of the different components involved in elaborate system that is American trypanosomiasis (the protozoan parasite *T. cruzi*, vectors Triatominae and the many reservoirs of infection), as well as the interactions existing within the system, is still incomplete. The Triatominae probably evolve from predatory reduvids in response to the availability of vertebrate food source. However, the basic mechanisms of adaptation of some of them to artificial ecotopes remain poorly understood. Nevertheless, these adaptations seem to be associated with a behavioral plasticity, a reduction in the genetic repertoire and increasing developmental instability.

Trypanosoma cruzi / Triatominae / mammalian host / vectorial transmission / adaptative strategies

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1. INTRODUCTION

American trypanosomiasis is a zoonosis caused by the flagellate protozoa *Trypanosoma cruzi*, which is restricted to New World. The parasite is transmitted through the fecal droppings of infected blood-sucking insect vectors belonging to the subfamily Triatominae (Hemiptera and Reduviidae). Endemic foci of the human disease (also called Chagas disease) range from Mexico to the Northern half of Argentina and Chile, mostly in poor rural areas where houses are infested with domestic species of Triatominae (Figs. 1 and 2) [106]. As for animal trypanosomiasis, its distribution is widespread from the Southern United States to Patagonia (roughly 40° N to 45° S). More than 180 domestic, synanthropic and wild species of mammals, especially nest-building rodents and opossums (Marsupialia, Didelphidae), are likely to be infected by *T. cruzi* and to be involved in the transmission cycle of the disease [119].

After exclusively infecting animals for millions of years, *T. cruzi* began to infect humans relatively recently and, consequently, Chagas disease became an emerging infectious disease (EID). As with many EID, the key event in its emergence was a change in the vector-host-parasite ecology resulting from anthropogenic changes to wildlife habitats that allowed for vector-human contact and subsequent transmission of the zoonotic parasite to humans [118].

During the 20th century, Chagas disease was reported in all countries in Latin America, where it was the most important parasitic disease in terms of impact on national economies and public health systems [119]. The disease follows a chronic course, mainly affecting the heart (myocarditis) and digestive system (megaesophagus and megacolon). From the

beginning of the 1990s, regional control programs directed against domestic vectors were initiated in most Latin American countries. They led to a reduction of disease prevalence from 1990 estimates of 16–18 million people infected to current estimates of ~9 million [107]. However, many obstacles remain to the effective control of the human disease. Although Chagas disease is physically disabling and its lethality is unquestionable (at least one million Latin American people carrying the parasite will die unless a new scientific and/or political breakthrough occurs), the disease is no longer perceived as a major public health burden in many countries [76]. Part of the reason for this may be that Chagas disease most often afflicts vulnerable indigenous communities living below the poverty line. This situation has led to the recent classification of Chagas disease as a neglected disease by the Pan American Health Organization [6].

The epidemiology of Chagas disease is currently changing. The vector transmission of the parasite was until now mostly depended upon domestic vectors (especially *Triatoma infestans*, *Triatoma dimidiata* and *Rhodnius prolixus*) and displayed a hyperendemic pattern of the disease in many regions of Latin America. Gradually, and chiefly owing to the control programs employed against these domestic vectors, this pattern is being supplanted by emergent infection patterns and by previously secondary transmission patterns [107]. As such, *T. cruzi* congenital infection and transmission by blood transfusion is becoming important, not only in Latin America but also in European and Northern American countries where Latino immigration occurs [119]. Also of concern are the emergent vector transmission patterns that involve wild Triatominae displaying



Figure 1. Bug collectors searching for Triatominae in an adobe-walled house. (A color version of this figure is available at www.vetres.org.)



Figure 2. Crack offering refuge to triatomine bugs and fecal streaks in a mud wall. (A color version of this figure is available at www.vetres.org.)

synanthropic trends and coming into contact with humans. Finally, human cases arising from ingestion of contaminated food recently recorded in Brazil are puzzling to health authorities [122]. Although human-related ecological alterations seem to favor change in the patterns

of transmission, the basic mechanisms involved remain poorly understood.

This article reviews the different components that make up the elaborate system that is American trypanosomiasis and the interactions between components within the system.

2. THE DIFFERENT COMPONENTS OF AN ELABORATE SYSTEM

2.1. The parasite

Since the discovery of *T. cruzi* as the etiological agent of American trypanosomiasis (Chagas disease) by Carlos Chagas in 1909, our knowledge about the wide genetic variability and phylogenetic relationships among the different lineages of the parasite has grown exponentially [24]. This growth of knowledge has been supported by theoretical approaches based on population genetics analysis and the increasing development of molecular biology tools used to identify different genetic subtypes of the parasite [14, 15, 38, 39, 108, 110, 111]. Today, *T. cruzi*, a flagellate eukaryote of the order Kinetoplastida, is classified as a single species, with an intraspecific classification that recognizes different lineages and sublineages.

A general objective of most taxonomic classifications is to reflect, as accurately as possible, the evolutionary history of the taxonomic units considered. This is not a simple task for any organism and it becomes even harder if the studied organism has an atypical eucaryotic genetic biology, as is apparently the case for *T. cruzi* [46]. Moreover, when the organism under study is of medical importance, it is desirable that both the taxonomic classification and the data that lead to the identification of the taxonomic units be of epidemiological and/or clinical relevance. In this sense, and in spite of controversies about phylogenetic relationships among the different lineages within *T. cruzi*, molecular data allow us to draw six general conclusions based on current evidence: (i) there are major phylogenetic lineages of the parasite with great genetic distances between them [112]; (ii) there is a general agreement about intraspecific nomenclature: there are two major lineages, *T. cruzi* I (TCI) and *T. cruzi* II (TCII), with five sublineages within *T. cruzi* II (IIa, IIb, IIc, IIId and IIe) [3, 14]; (iii) the evolution of the different lineages has been basically clonal, with some ancient genetic exchange and recombination events that had an important impact on the generation of genetic diversity within this taxon [13, 14, 117]; (iv) there are lineages with

a hybrid origin arising from ancient hybridization events between parasites belonging to distantly related lineages [75, 117]; (v) at least some lineages have some capacity for genetic exchange and homologous recombination, as demonstrated experimentally [46]; (vi) there is evidence suggesting genetic exchange in natural cycles involving TCI strains, although there are no data about the frequency of these events or their impact on the genetic structure of natural populations of the parasite [22, 33]. These six statements constitute a useful framework that makes sense in both evolutionary and epidemiological terms. Moreover, the nomenclature used (*T. cruzi* I, IIa, IIb, IIc, IIId, and IIe) is a very good approximation of the likely “real” phylogenetic structure of the parasite. Indeed, the genetic markers available for lineage typing are very useful considering the complex and still unresolved epidemiology of the different lineages, as well as the possible relationship between the range of clinical manifestations and the infecting lineages [7, 14, 15].

The general pattern of the geographic distribution of *T. cruzi* lineages and sublineages is shown in Figure 3. Among all lineages and sublineages, TCI has the largest distribution, from the southern part of the USA to northern Argentina and Chile. TCIIa has been reported in the northern Amazon basin and in the USA, with some reports from south of the Amazon basin. TCIIc is present from the Amazon basin to southern endemic regions while TCIIId and TCIIe have only been reported in the Southern Cone countries. TCIIb occurs mainly in the Southern Cone countries; however it has also been reported in Colombia [123].

It is worth mentioning that different numbers of phylogenetic lineages within *T. cruzi* have been proposed: 3 or 4 major lineages, based on mitochondrial or nuclear DNA sequences, respectively [75]; 3 lineages, using microsatellite analysis [87]; and also 3 lineages based on sequence analysis of an 1 100 bp DNA fragment [96]. Today, it seems clear that most of these inconsistencies among different phylogenetic analyses are related to the hybrid origin of some lineages, which generates different phylogenetic relationships according to the

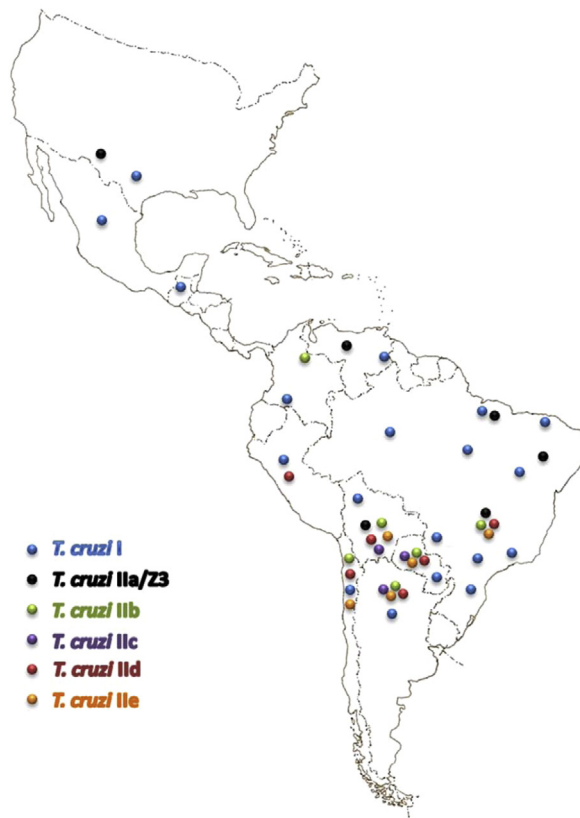


Figure 3. General pattern of distribution of *T. cruzi* lineages and sublineages; the isolates typed as *T. cruzi* II using the mini-exon gene (uncharacterized sublineages) are not represented on the map.

number and type of examined markers, as well as the number and genetic diversity of the studied isolates.

Based on molecular data, a very plausible model for the evolution of the different *T. cruzi* lineages suggests an ancestral fusion of strains belonging to lineages TCI and TCIIb, producing a heterozygous hybrid which through homogenization of its genome became the homozygous progenitor of the TCIa and TCIIc lineages [117]. A second hybridization event between TCIIb and TCIIc strains could have generated the TCIId and TCIIe lineages. Under this hypothetical model, every hybridization event was separated by long periods of clonal evolution. In light of such a complex evolutionary framework and the existing evidence, the current

consensus of six lineages or DTUs (discrete typing units), namely *T. cruzi* I, IIa, IIb, IIc, IId, and IIe, seems to be the most parsimonious and useful in epidemiological terms, and is probably also the most accurate phylogenetically [113].

Accepting the intraspecific nomenclature proposed before, and assuming a general genetic stability in space and time of the lineages as suggested by the strong linkage disequilibrium systematically observed in *T. cruzi*, different authors have hypothesized about the evolutionary history of the different lineages. These evolutionary models are based on present data regarding associations among parasite lineages, insect vectors and mammalian hosts; present and inferred ancient geographical distribution of vectors and

mammalian hosts; and reconstructions of the molecular evolutionary history of parasite lineages [12, 45, 75]. All these analyses, beyond generating basic knowledge about the evolutionary history of *T. cruzi*, have the goal of allowing us to obtain a picture of the eco-epidemiological characteristics involved in the transmission dynamic of the different *T. cruzi* lineages. Certainly, the understanding of the parasite evolutionary history and the time of divergence of the different lineages could explain some present associations among different hosts and vectors. The occurrence of common evolutionary histories among parasite lineages, hosts and vectors, could help investigators identify phenomena that led to the enhanced fitness of some lineages under certain epidemiological situations, and provide examples of co-evolutionary processes. An examination of such processes could also lead to the identification of adaptative mechanisms that would allow different lineages to infect different mammalian species (including humans), and may also explain the diversity of clinical manifestations of Chagas disease.

Widely different times for the divergence between the TCI and TCII lineages have been proposed [12, 59, 75]. Whatever is the divergence time between TCI and TCII, certain data support the association of TCI with the triatomine tribe Rhodniini and the marsupial genus *Didelphis* (opossum), while TCII has been proposed to have evolved in association with the triatomine tribe Triatomini and edentate placental mammals such as armadillos (Xenarthra, Dasypodidae) [45, 121]. This hypothesis suggests that both, *T. cruzi* I and *T. cruzi* II, would have evolved contemporarily in South America up to 65 million years [45]. A relatively recent report presented the finding of *T. cruzi* IIc, IIb and II d lineages in armadillos in the wild in Paraguay [121]. This study included a very comprehensive analysis of historically published host records of *T. cruzi* lineages that clearly suggests the proposed association of the genus *Didelphis* with TCI, and armadillos with TCII [121]. Whether these data reflect an ancient association that could imply adaptation to marsupials (*T. cruzi* I) and placentals (*T. cruzi* II), remains to be confirmed.

A recent and comprehensive phylogenetic study of the genus *Trypanosoma* and co-evolution with its hosts suggests that “ecological host fitting”, a process by which organisms are able to colonize new hosts as a result of traits they already possess, may have played a principal role in trypanosome evolution, even more than co-evolution [50, 58]. If this general pattern of evolution proposed for the genus *Trypanosoma* is also the rule for the evolution of the different lineages within the *T. cruzi* taxon, adaptations of different lineages to specific hosts could have not occurred.

2.2. Hosts and reservoirs

The concept of what constitutes a reservoir host has changed over time. Carlos Chagas, who described the etiological agent of the human disease that later received his name, also described one of its wild hosts, an armadillo species that he identified as *Tatusia novencincta*. In his description, Carlos Chagas named this infected armadillo as the “depository” of *T. cruzi*. The use of this term reflects the once generally accepted idea that a wild mammal species harboring a parasite capable of infecting humans is an almost inert repository of this parasite that acts as the source of human infection. The notion of a host as a complex living system, able to suffer and apply selective pressure on parasites according to their environmental conditions, thereby playing active and distinct roles in the transmission cycle is much more recent. The complexity of variables that influence a given parasite transmission cycle, in time and space scales, is especially important when considering the study of multi-host parasites such as *T. cruzi*. This protozoan fully embodies the characteristics of a successful parasite since it is maintained in nature by numerous vector species and mammal host species distributed across almost all biomes and habitats in the Americas. *T. cruzi* is also able to colonize almost all tissues in its many mammalian hosts, even unconventional sites, such as the scent glands of Didelphid marsupials [29].

Despite the identification of a large list of wild mammal species naturally infected by *T. cruzi*, there are many unanswered questions



Figure 4. *Didelphis albiventris*, which may simultaneously act as a reservoir and a vector for *T. cruzi*. (A color version of this figure is available at www.vetres.org.)

concerning the role played by these species in the maintenance and/or transmission of *T. cruzi* in the wild [8, 119]. Most studies are geographically restricted and do not include data on the broader environmental conditions, such as local fauna diversity, the relative abundance of infected animals or the population structure. Here, we reappraised the concept of reservoir by including, when possible, the peculiarities of the interaction of *T. cruzi* with its known wild and domestic hosts. We define a reservoir as one species or an assemblage of species responsible for the long-term maintenance of a given parasite in a given environment [4].

2.2.1. Marsupials

These ancient mammals are considered, together with Xenarthra (armadillos and anteaters), to be the earliest hosts of *T. cruzi* as South American mammals were restricted to these taxa after separation of this continent. Marsupial species of the genus *Didelphis* (Fig. 4)

display a unique interaction with *T. cruzi*: they are able to concomitantly maintain amastigotes multiplying in the tissues and epimastigotes multiplying and differentiating in their scent glands (Fig. 5) [29]. These findings showed that *Didelphis* sp. may act simultaneously as reservoir and a vector of *T. cruzi*, and suggest a possible ancient transmission mechanism of this parasite before the acquisition of hematophagous habits by triatomines. Metacyclogenesis of *T. cruzi* in the lumen of the scent gland occurs without adhesion of the parasite and may be significant (involving up to 50% of the *T. cruzi* present), however the “vectorial” capacity of Didelphidae in natural conditions is still unknown (Fig. 4) [23]. The infectivity of opossum scent gland metacyclic forms was proven by successful experimental infection of Swiss mice. Monogenetic trypanosomatids (*Leptomonas*, *Crithidia* and *Herpetomonas*) may also efficiently colonize the scent glands of *Didelphis aurita* under experimental conditions when the trypanosomatids are directly

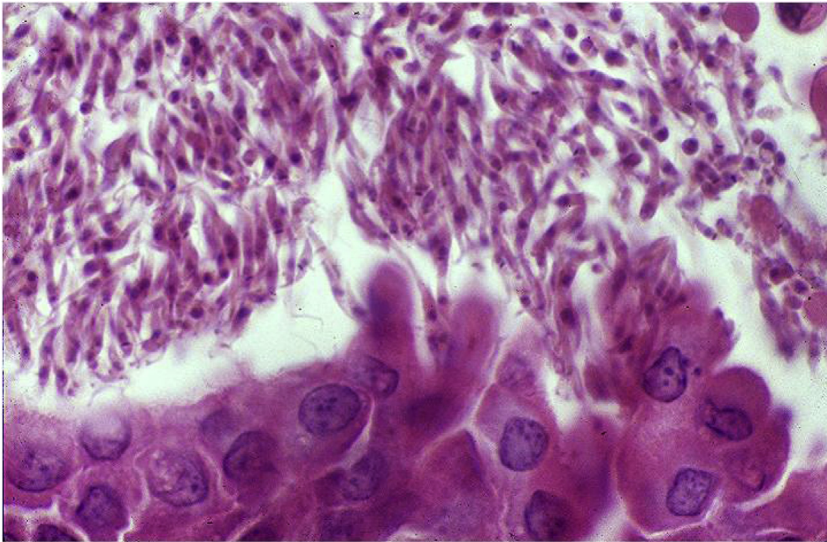


Figure 5. *T. cruzi* epimastigotes multiplying and differentiating in the scent gland of *Didelphis* sp. (A color version of this figure is available at www.vetres.org.)

inoculated into the glands [57]. Marsupials of the genus *Didelphis* are non-specialized mammals and may be found on the ground and in the canopy. They are excellent climbers but move clumsily on the ground. *Didelphis* may provide excellent reservoirs of *T. cruzi* since these animals may maintain long-lasting infections and high parasitemias. Notably, their competence as reservoir varies in space and time. This is exemplified by the differences of *T. cruzi* infection prevalence that ranged between 11–90% in *D. aurita* collected from various sites in Rio de Janeiro State, Brazil [40]. Furthermore, due to its synanthropic behavior, the genus *Didelphis* should always be considered an excellent reservoir of *T. cruzi* in peridomestic, domestic and disturbed habitats. In a pristine wild environment, other taxa assume the role of reservoir. Opossums are known for their adaptability to human environments and are considered to be indicators of environmental disturbance [86]. In experimental conditions, *D. aurita* has been described as being able to maintain stable infection by TCI isolates while having the capacity to control and even eliminate TCII isolates. This observation was reinforced by the regular

isolation of TCI and not TCII isolates in naturally infected animals [40]. However, these experimental studies were undertaken with only two TCII isolates, which is not representative of the diversity of the heterogeneous TCII lineage. In fact, naturally TCII-infected *D. aurita* and *Didelphis albiventris* animals have been detected in several biomes showing that this marsupial may act as an effective biological filter for some subpopulations of *T. cruzi* but not necessarily for the entire TCII lineage. Animals from Didelphidae family display unique interaction patterns with *T. cruzi*. For instance, *Philander frenatus*, in contrast to *D. aurita*, maintains a stable infection with a high and long lasting parasitemia in experimental infections with *T. cruzi* Y strain [67]. It is likely that these two marsupials apply distinct selective pressures on *T. cruzi* subpopulations and consequently play dissimilar roles in the maintenance and transmission of this parasite in nature.

2.2.2. Primates

Primates were incorporated much later into the transmission cycle of *T. cruzi* than

marsupials, as they are supposed to have colonized the American continent 35 million years ago [42]. However, they also display an ancient co-evolutive history with *T. cruzi*. In the golden lion tamarin (*Leontopithecus rosalia*), an endangered callithrichid species maintained in a conservation program in a biological reserve in the state of Rio de Janeiro, Brazil, it was shown that a same animal host species may display different competences as a reservoir, even in the same forest unit. In this reserve, *T. cruzi* infection with the genotype TCII lineage was demonstrated in 46% of the golden lion tamarins [70]. However, infectivity, demonstrated by high parasitemia levels (positive hemocultures), varied between colonies. The subsequent observation of a positive association between the frequency of tamarins with *T. cruzi*-positive blood cultures and the presence of Trichostrongylidae showed that concomitant parasitic infections may increase the transmissibility of *T. cruzi* and, consequently, the competence of this host species as a reservoir [81]. Infection by genotype TCI was detected only in a small number of the tamarin specimens. In contrast, in the Amazon basin region of Brazil, only the TCI lineage was detected in the primate species examined [71].

2.2.3. Bats

In a recent survey of 93 specimens belonging to four different bat families (Molossidae, Noctilionidae, Phyllostomidae and Vespertilionidae), 15% had *T. cruzi* positive hemocultures [72]. The majority (80%) of these positive hemocultures were derived from the generalistic species *Phyllostomus hastatus*, suggesting that this bat species may play an important role in the maintenance and dispersal of *T. cruzi*. An electrophoretic analysis of multiple enzyme loci (MLEE) of the isolates did not show any correlation between zymodeme, bat genera, species and geographic origin. The analysis of minixon genes showed that 10 out of 14 *T. cruzi* bat isolates corresponded to the TCII lineage. The existence of mixed infections with TCI, TCII (unspecified sublineage) and Z3 (TCIIa) in bats indicates that these mammals may be involved in the dispersal of various *T. cruzi* genotypes [72].

2.2.4. Free ranging carnivores

The oral transmission of *T. cruzi* is probably especially important among carnivores. A study conducted in the Brazilian Pantanal region focused on *T. cruzi* infection in free ranging coatis (*Nasua nasua*), a carnivore species commonly observed in this biome, reported a high antibody prevalence. The high number of positive hemocultures obtained showed that coatis may be highly infective. Single infections by TCII (unspecified sublineage) (32.1%) and TCI (28%) as well as Z3 (TCIIa) (7.1%) were most commonly observed, however, mixed infections TCI/TCII (10.7%) and TCI/Z3 (TCIIa) (3.6%) were also detected. The follow up of recaptured coatis showed that infection with high parasitemia levels is commonly long lasting in this species. Due to their high biomass, coatis certainly play a role in the amplification and dispersal of the main *T. cruzi* subpopulations, demonstrating that predator-prey links may be excellent mechanisms for *T. cruzi* transmission and perpetuation in the wild [54]. Similarly, other carnivores, such as raccoons (*Procyon lotor*) and gray foxes (*Urocyon cinereoargenteus*) are frequently exposed to *T. cruzi* infections [51, 99].

2.2.5. Armadillos

In spite of being the very first wild mammals in which *T. cruzi* infection was observed, the interaction of these Xenarthrans (Fig. 6) with *T. cruzi* remains largely uncharacterized. *T. cruzi*-infected armadillos have been reported from the USA to Uruguay, and as with other wild mammal species, infection rates do vary from region to region. In Louisiana, USA, infection by *T. cruzi* was detected in 26% of 80 armadillos [120]. In the same state, but in another region, only 3.9% of 415 wild nine-banded armadillos (*Dasypus novemcinctus*) were infected [88]. A retrospective study conducted in French Guiana described this species as the main *T. cruzi* reservoir behind the marsupial *Didelphis marsupialis* [93]. More recently, high infection with TCII was detected in nine-banded armadillos and six-banded armadillos (*Euphractus sexcinctus*) in Paraguay [121].



Figure 6. *Dasypus novemcinctus* (nine-banded armadillo). (A color version of this figure is available at www.vetres.org.)

2.2.6. Rodents

Rodents are ubiquitous mammals that are found, in most areas, in high relative abundance and have a broad range of habitats. Furthermore, both wild and synanthropic species have been reported as being highly involved in *T. cruzi* transmission in several regions. Caviomorph rodents (i.e. guinea pigs and allies) are especially interesting as they have an ancient co-evolutionary history with *T. cruzi* since, like primates, they arrived in the Americas about 35 million years [42]. Naturally infected *Thrichomys laurentius* and *T. pachiurus*, two sibling caviomorph rodent species, have been reported in two very dissimilar biomes in Brazil: in the Caatinga, the semi arid region of the northeastern Brazil (white shrub) and in the Pantanal, a flood plain region in the central part of the country, respectively [55]. TCI, TCII and Z3 (TCIIa), as well as one mixed TCI/Z3 (TCIIa) infection, were detected in these rodents [53]. The abundant rodent species *Oecomys mamorae* (arboreal rice rat) was also demonstrated to be involved in the maintenance and transmission of *T. cruzi* [53]. These results do not support an association between the *T. cruzi* main genotypes and the mammalian host taxa. In addition, the coexis-

tence of multiple strains circulating in a same biome and even in a same animal is probably a very common feature in wild mammals [101]. The distinct subpopulations and clones of *T. cruzi* are continuously submitted to distinct selective pressures in their vertebrate and invertebrate hosts. These selective pressures are the outcome of traits inherent to host species but are also, and perhaps much more strongly, related to populational and even individual conditions. Nutritional status, age, stress conditions, previous and current parasitic infections of the hosts besides inter- and intraspecific competition and/or cooperation among parasites should also be considered as components of the habitat that a parasite encounters. The coexistence of *T. cruzi* lineages and sublineages has been observed in wild and peridomestic mammals in Chile [101]. In the same study, no association was found between the genotype of the parasite and host species, suggesting that the profile of animal infection by *T. cruzi* depends on the predominant genotype in a given locality that is defined by the interplay of all biotic factors present at the time of transmission. As such, contradictory data from different studies may reflect different epidemiological scenarios and methods, limited or restricted data and/or biased or incomplete sampling of habitats.

2.2.7. Domestic animals as reservoirs

As observed for wild mammals, domestic species may also play various roles in the *T. cruzi* transmission. They may act as a link between wild and peridomestic or domestic habitats or they may merely be dead end hosts.

2.2.7.1. Dogs and cats

Dogs and cats are very often found with a high *T. cruzi* infection prevalence. As for all other mammal species, their role in the transmission cycle of *T. cruzi* varies as shown by the patterns of infection in distinct study areas. In the Argentinian Chaco, both cats and dogs have been reported to be epidemiologically important and highly infective for triatomines [49]. Similar findings were described in western Venezuela where both TCI and TCII genotypes infect dogs [26]. An active canine *T. cruzi* transmission cycle with severe symptoms affecting a broad range of dog breeds and age groups was observed in several counties in Texas, USA [61]. In contrast, the absence of positive dogs by IFAT testing led to conclude that dogs are not important in the dispersion of *T. cruzi* in several urban areas of Brazil and Colombia [99]. In our experience, in spite of the high antibody prevalence, we have not observed parasitemias upon direct examination of fresh blood smears, nor have we observed positive hemoculture in various areas of Brazil including the Amazon basin and the Northeast and Southwest regions. A direct correlation of seropositivity between humans and dogs was observed in Mexico, emphasizing the importance of sentinel dogs as a surveillance measure in that country [37].

2.2.7.2. Pigs

An interesting situation is represented by feral pigs in Pantanal, the seasonally flooded basin in the central part of Brazil. These animals, derived from domestic swine (*Sus scrofa*) that went back to the wild, have been implicated in the maintenance of *T. cruzi* in nature [52]. In addition, pigs may be attractive hosts for triatomines such as *Panstrongylus*

geniculatus, as observed in new human settlements in the Amazon basin [116].

2.2.7.3. Goats

The limited number of studies on *T. cruzi* in goats suggests that the role played by these peridomestic mammals in the transmission cycle of *T. cruzi* is mainly secondary in spite of their high exposure to infection. There are no reports of high parasitemias in naturally-infected animals, and the presence of the parasite was detected only by PCR testing or indirectly by the presence of specific antibodies [55, 101].

2.2.8. Birds and others non-reservoir feeding hosts

Birds are refractory to *T. cruzi* infection owing to a complement-mediated lytic effect of their blood on the parasite [60]. Nevertheless, they may play an indirect but significant role in the maintenance of *T. cruzi* parasites in both wild and peridomestic environments since they are important feeding sources for Triatominae. In fact, the schedule of blood intake by infected triatomine bugs influences the *T. cruzi* population density and the proportion of the population in various developmental stages. Short feeding intervals increase the total population density of *T. cruzi* in the vector in contrast to starvation, which reduces the total number of flagellates and the number and percentage of trypomastigotes [63]. In the domestic environment, chicken coops are very attractive to triatomines. Additionally, in rural areas, poultry shelters are frequently constructed very close to human dwellings. This practice certainly results in an increase in the local triatomine population size and may either act as barrier against domiciliary invasion by the insects or may increase the chances of contact between triatomines, domestic mammals and humans.

2.3. Triatomine bugs

The Triatominae (Hemiptera, Reduviidae), still named kissing bugs, are blood-sucking insect vectors of *T. cruzi*. Their life cycle

consists of five nymph stages before they reach sexual maturity. This process requires about six months, depending upon the species. Their size ranges from a few millimeters after hatching to more than 3 cm for adults, in some species. Currently, 139 triatomine species are listed, belonging to 6 tribes and 18 genera [18, 43, 68]. Most of these species are widespread in the Americas, except for the Indian genus *Linshcosteus*, and the tropicopolitan *Triatoma rubrofasciata* and its related Asian species. The species richness of the New World Triatominae increases from the poles towards the equator, peaking over the 5°–10° S latitudinal band [97]. Several pieces of evidence strongly support a polyphyletic origin for the Triatominae though the classic assumption of monophyly persists [56, 104]. The Triatominae subfamily is assumed to have evolved from Reduviidae predator groups dispersed in several areas of the world rather than stemming from a cladistic class of individuals sharing a common ancestry [21]. In human and veterinary medicine, the importance of three genera, namely *Triatoma*, *Rhodnius*, and *Panstrongylus*, lies in the fact that some of their members feed on humans and synanthropic mammals and may thus transmit *T. cruzi* [119]. However, wild populations of all triatomine species (except those that are strictly ornithophilous) also maintain an enzootic cycle involving wild mammals in a variety of terrestrial or arboreal biotopes. Habitat trends also exist in the different triatomine genera. As such, *Rhodnius* species are primarily associated with palm trees, *Panstrongylus* species are predominantly found within burrows and tree cavities, and the genus *Triatoma* is found in rocky habitats or hollow trees [19, 45]. Some species display a close relationship with one type of habitat (*Rhodnius brethesi* with the palm tree *Leopoldina piassaba*, *Psammolestes* species with nests of Furnariidae), while others exhibit a great range of terrestrial and arboreal ecotopes (*Panstrongylus megistus*, *T. dimidiata*). Particularly interesting are the cases of *T. infestans* and *Triatoma sordida*, in which wild populations at high altitudes live in rock-piles while those in lowlands are found in hollow trees [85]. Such species provide good models for studying the pressure

of habitat on morphology, genetic traits and behavior in Triatominae.

All species of Triatominae were originally sylvatic and, among them, only a small number were successful in establishing domestic colonies, thus potentially becoming involved in transmission of *T. cruzi* to humans. The anthropogenic environmental changes and ensuing damages to triatomine natural biotopes seem to favor the synanthropic process. However, the basic mechanisms of adaptation of these insects to artificial ecotopes remain poorly understood. We do not know if only some of the sylvatic genotypes/haplotypes may be successful in the process of adaptation. Nevertheless, the trends toward domesticity seem to be associated with behavioral plasticity, a reduction of the genetic repertoire and increasing developmental instability, making a triatomine species/population a more efficient vector as well as a more vulnerable target for control measures [17, 119].

The classification of the Triatominae based on synanthropic process is most often adopted [119]. Among the neotropical species that have proven adaptable to human dwellings are *T. infestans* in the Southern Cone countries (Fig. 7), *R. prolixus* in Colombia, Venezuela, and also in parts of Central America, and *T. dimidiata*, a species distributed from northern South America to Central America and Mexico. Proof of the epidemiological importance of these insects lies in the 65% reduction in the incidence of Chagas disease in Latin America that followed the elimination of synanthropic populations of these insects over large areas [119]. Long considered exclusively domestic species or species with limited wild populations, *T. infestans* and *R. prolixus* actually have a large sylvatic distribution in Bolivia and Venezuela, respectively [41, 84]. Wild *T. infestans* inhabits tree holes in the Chaco and can be found under large piles of rocks in the Andean valleys [25, 84]. Over its entire range, this vector colonizes rural as well as urban areas and has proven highly adaptive to human dwellings. *R. prolixus* is an arboreal species living in palm trees and can reach very high population densities in houses [105]. There is clear evidence of genome reduction in



Figure 7. Sylvatic form of *Triatoma infestans*, the main vector of Chagas disease in the Southern Cone countries. (A color version of this figure is available at www.vetres.org.)

derived populations of *T. infestans* from the lowlands of the Southern Cone countries compared with their putative ancestral populations from the Bolivian Andes [89]. Similarly, *R. prolixus* in Central America has a reduced genome compared to ancestral populations in Venezuela and Colombia [35]. *T. dimidiata* displays a great diversity in its biology and behavior across its geographic range and has the ability to transmit *T. cruzi* to humans at a very low density [34]. Human intervention has led to the passive dispersal and subsequent colonization of new areas by synanthropic populations of these vectors [34, 35, 90]. Wild populations of these three vectors may be potential sources of reinfestation of treated areas with epidemiological consequences [34, 41, 85]. Another major obstacle to the control of these main vectors may stem from a reduction in the effectiveness of the insecticides used. For example, in *T. infestans*, high resistance to pyrethroid insecticides has been recently associated with ineffective field treatments in northern Argentina and southern Bolivia [92, 114].

Besides these primary vectors, other triatomine species are displaying synanthropic trends. Based on their geographical distribution, ability for colonizing artificial structures and vectorial capacity, they are named secondary species (*Triatoma brasiliensis*, *T. barberi*, *P. megistus*...) or vector candidates (*T. sordida*, *Triatoma pseudomaculata*...) [84]. The secondary species are capable of colonizing dwellings in the absence of a primary species whereas the vector candidates are poorly adapted to living in houses and instead build colonies in the peridomestic environment [32]. Traditional control approaches involving household insecticide spraying are unlikely to be effective against these vectors that keep a sylvatic habitat.

Cryptic species seem to be evenly distributed among Triatominae as they are among other major metazoan taxa [91]. Cryptospeciation was described in the genera *Triatoma* (*T. brasiliensis*, *T. dimidiata*, *T. sordida*) and *Rhodnius* (*R. robustus*, *R. ecuadoriensis*) [1, 79, 80, 83, 90]. More surprising, some triatomine species display considerable phenotypic variability. In *T. infestans*, four morphs exhibiting obvious chromatic and morphological differences were observed [25, 48, 82]. As the different morphs occupy distinct biotopes (hollow tree, palm tree, rock) and biogeographic regions, the plasticity displayed by the wild *T. infestans* supports the theory suggesting the occurrence of rapid morphological divergence in response to different ecological factors [36].

3. INTERACTIONS WITHIN THE SYSTEM

3.1. Interactions between *T. cruzi* and triatomine bugs, and vectorial capacity

The localization of *T. cruzi* in the insect vector gut entails a series of morphological and physiological transformations that take place at different sites along the digestive tube of the vector (Fig. 8). Morphological changes involve passage from the unreplicative blood trypomastigote form, which is ingested with a blood meal from an infected host, through the replicative epimastigote stage, to reach the

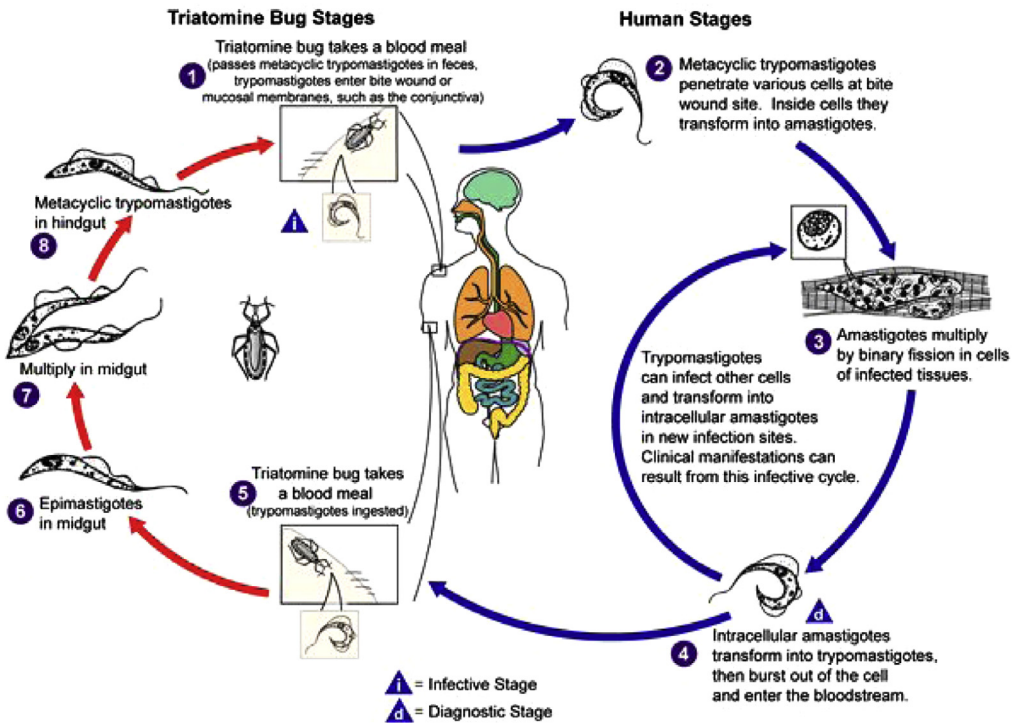


Figure 8. *T. cruzi* life cycle. From the DPDx Parasite Image Library of the CDC (<http://www.dpd.cdc.gov/dpdx>). (A color version of this figure is available at www.vetres.org.)

infective metacyclic trypomastigote stage. The blood trypomastigote to epimastigote transformation takes place in the anterior part of the midgut, and subsequently, epimastigotes undergo binary divisions mainly in the posterior part of the midgut. Finally, epimastigotes reach the rectum and attach to the rectal cuticle, where they transform into metacyclic trypomastigotes, which will be eliminated with the insect feces and will be able to infect mammals [115]. At each one of these steps, vector-parasite interactions occur and parasites are exposed to a series of factors that will influence their viability. It has been postulated that viability of *T. cruzi* within the digestive tract of the vector, largely depends on the host nutritional state, the parasite strain, trypanolytic compounds, digestive enzymes, lectins, resident bacteria in the gut, and the endocrine system of the insect vector [44].

Two main approaches have been used to study interactions between vector and parasite in Chagas disease. The first one of these, that can be called the “molecular and biochemical approach”, involves studies that deal with biochemical and molecular factors that may be involved in the mechanisms determining both the viability and the morphological-physiological changes that *T. cruzi* undergoes within the insect vector. The second approach, that can be called the “transmissibility approach”, involves a wide range of studies that deal with the differential capacity of triatomine species to transmit *T. cruzi*.

From the “molecular and biochemical approach”, it must be considered that through *T. cruzi* passage and multiplication within the digestive tube of the insect vector, the parasites are exposed to hostile environments, and also to

factors that may act as signals to induce morphological and physiological changes. The mechanisms by which parasites evade potentially harmful insect factors are not well understood yet, however the widely diversified gene families that encode for surface molecules, some of which are differentially expressed in the different morphological stages of the parasite, are hypothesized to be involved [119]. Notably, some of the different insect factors to which parasites are exposed are well-described. Crop hemolytic factor, which lyses the erythrocyte membrane and releases free hemoglobin for digestion, and lectins have been suggested as possible modulators of multiplication and transformation of *T. cruzi* [44]. Differential agglutinin and lysin activities in the gut of *R. prolixus* toward different strains of *T. cruzi* have also been reported [77]. Furthermore, correlations among infectivity, differences in the surface carbohydrates of each parasite strain, and both agglutination and lytic characteristics for these strains have been observed, suggesting a strong influence of some parasite surface molecules on their infectivity to the insect vector [77]. Cysteine proteases and a cDNA encoding for a lysozyme from the gut of *T. infestans* have also been described, but their role as possible vector-parasite interaction factors has not yet been elucidated [64, 65]. Another factor that could be involved in the success of vector infection by *T. cruzi*, at least indirectly, is the presence of symbionts in the digestive tube of triatomines. It is possible that many vitamins and nutrients absent in the diet of feeding blood insects, such as triatomines, are supplied by symbionts that live in their intestines. This was demonstrated with the bacterium *Rhodococcus rhodnii*, which supplies nutrients to the insect *R. prolixus* [10]. The presence of symbionts could reduce the needed concentrations of components of the vector humoral immune system, such as defensin molecules, in the digestive tube of triatomines [73]. This environment of lower humoral immune components could consequently be advantageous for *T. cruzi* development. During development within the insect vector, *T. cruzi* must attach to different epithelial cell surfaces along the digestive tube. These processes of attachment

involve recognition between parasite-vector cell surface molecules. Some of these molecules have been identified and found to be differentially expressed during the metacyclogenesis process wherein epimastigotes are transformed into metacyclic trypomastigotes. For example, it has been shown that the gene encoding a 7-kDa protein belonging to a chitin-binding-like protein gene family displays maximal synthesis levels in differentiating epimastigotes [27]. This and other related data suggest an elaborate molecular system involved in attachment of parasites to the rectal wall, which is thought to correlate with metacyclogenesis [2, 62].

Data gathered using the “transmissibility approach” suggest that there are differences in vectorial capacity among different species of triatomines, increased transmissibility when *T. cruzi* strains and triatomine bugs from the same region are involved, and differences in transmissibility for each parasite lineage in the same vector species (*T. infestans*) [16, 30, 31]. Serial xenodiagnosis performed on dogs showed that vectors from a given area are more susceptible to local strains of *T. cruzi* [31]. More recently, a higher capacity of the Chilean triatomine *Mepraia spinolai* to host and foster the reproduction of different *T. cruzi* lineages (*T. cruzi* I, IIb, IIc and IIe) was demonstrated compared to that of *T. infestans*. The *M. spinolai* used in that study was a wild vector from the same area where the *T. cruzi* isolates were obtained [16]. The data cited above suggest some degree of interaction among triatomine species and *T. cruzi* lineages. However, more studies are needed to elucidate whether these interactions do occur, and to understand the nature of these phenomena.

3.2. *T. cruzi* transmission to animals and humans

The classical theory regarding the natural history of Chagas disease proposes that humans were included in the *T. cruzi* transmission cycle (i) as a consequence to the acquisition of sedentary habits brought about by domestication of plants and animals or (ii) as a consequence of the colonization by triatomine vectors of human dwellings built after European colonization [100]. Either way, the new

ecological environment favored domestication of *T. infestans* and, as a consequence, transmission of *T. cruzi* to humans. Nevertheless, recent paleoepidemiological studies reported the antiquity of human *T. cruzi* infections in the Andean region and more recently in Brazil, in organic remains of hunter and gatherer people [5, 69]. Megacolon, a typical lesion of chronic Chagas disease, has been observed in Chilean and Peruvian mummies up to 9 000 years old [5]. This diagnosis was confirmed by the recovery of *T. cruzi* kDNA in these mummy samples. Infection by the TCI genotype of a 7 000 years old human sample was also shown, in Minas Gerais State, Brazil, where contemporary cases of Chagas disease are now predominantly TCII [69]. Hunters and gatherers were described as consumers of raw meat since remnants of raw bones and rodent fur have been detected in human coprolites [94]. Therefore, it is clear *T. cruzi* infection and Chagas disease preceded European colonization of the continent, and that this parasite has infected humans since their entry in the Americas. These data also suggest that the current microepidemic outbreaks of Chagas disease in the Amazon basin reflect a reemergence of this ancient epidemiological cycle in the Americas since the oral route of infection is very efficient and the acidic environment of the stomach enhances the infection competence of *T. cruzi* [122]. Among wild mammals, this route is more probable than infection through metacyclic forms voided in feces of triatomines that must pass through the dense fur of the animal. Furthermore, in the last three years, orally acquired *T. cruzi* infections attributed to the ingestion of food contaminated by metacyclic forms have become more frequent and several human cases have been reported.

The study of wild and domestic mammals in human Chagas disease outbreak areas characterized by oral infection has produced distinct enzootical pictures. The first one of these is a classical picture of acquisition of Chagas disease by entrance of humans into the *T. cruzi* transmission cycle in the wild where patches of forest are still preserved, with all representatives of the local wild fauna infected by the parasite. Here, a panzootic scenario is observed

and high parasitemia levels are detected in wild mammals of all local ecological strata. The second picture is of disease outbreaks resulting as the consequence of the reduction of mammalian fauna and the selection of new suitable reservoir hosts for *T. cruzi*. As a common point between the two pictures, it was observed that domestic animals were highly exposed to infection, showing that these animals should be considered as sentinels for *T. cruzi* transmission in the environment [98].

3.3. Interaction between Triatominae and their hosts

All members of the Triatominae subfamily are hematophagous, which is considered to be a recent characteristic in evolutionary terms. Likewise, the five nymphal instars and the adult stage are generally hematophagous. Triatomines probably evolved from predatory reduviids (catching and eating other arthropods) in response to the availability of vertebrate food source in animal refuges and nesting sites, and underwent modifications in their buccal and digestive apparatuses during this process [21, 106]. However, primitive alimentary behavior still occurs in some triatomine species, which can feed on arthropods (Fig. 9) [78, 102]. Such a behavior seems to be more related to hemolymphagy than predation, where predigestion of tissues occurs followed by the death of the prey. A cleptohematophagous behavior (sucking blood from recently fed triatomines) could be a transitional stage between predation and hematophagy. Such observations suggest that adaptation to hematophagy is still occurring in Triatominae [20].

Triatomines belong to the group of vessel-feeding insects that use their proboscis as a needle to cannulate a host blood vessel from where they suck the blood. Obtaining a large blood meal is a time consuming and challenging task, and requires a great adaptation. Hematophagous adaptive evolution is reflected in the composition of the saliva, where various compounds are able to counteract the host hemostatic response and overcome vasoconstriction, coagulation and platelet aggregation. Antiplatelet and vasodilator activity levels are correlated



Figure 9. *Triatoma brasiliensis* nymphal instar feeding on the hemolymph of an immobilized scolopendra. (A color version of this figure is available at www.vetres.org.)

with blood feeding efficiencies in different triatomine species [95]. Triatomines are not that different from Reduviidae predators in habitat and forms. However, they ingest blood through a painless bite whereas the bite from Reduviidae predators tends to be very painful [106]. An action of triatomine saliva on sodium channels could be associated with loss of host sensibility at the bite site during the blood feeding process [28]. Hematophagy also requires a rapid compensation for the enormous amount of blood that triatomines ingest. The insect therefore immediately excretes great amounts of water and salts immediately to reduce its weight after a meal. Another adaptation to hematophagy is the erythrocytic rupture and hemolytic process that occurs at the beginning of digestion [21]. Sometimes, Chagas disease vectors can induce significant blood loss in their hosts in cases of massive domestic infestation [103].

A primary close association between *Rhodnius* species and marsupials, and between *Triatoma* species and rodents/edentates, was suggested in studies of the associated trypanosomes strains [45]. Similarly, it is suggested that the *Panstrongylus* species might be primarily associated with marsupials whereas the terrestrial species may also have been associated with edentates. Later, host transfers occurred to other sylvatic and synanthropic animals, which may

or may not be considered reservoir hosts of *T. cruzi*. Some triatomine species are highly adapted to a single host and are able to survive only in the microhabitat occupied by this particular host. Other species are more eclectic and can adapt more easily to different hosts. Unfavorable environmental changes and the subsequent rarefaction that occurs in wild fauna may conduce triatomines to move to man-made habitats, which are highly stable and offer a variety of hiding places and an abundance of food throughout the year [119].

The identification of feeding sources is very helpful in characterizing feeding habits of triatomine populations in wild, peridomestic or domestic environments, allowing for the recognition of synanthropic animal species that may favor colonization and long-term infestation of human habitats, and assessment of the importance of contacts between humans and vectors [11]. Nevertheless, faulty sampling may bias the studies and the conclusions about the blood sources of a triatomine species often refer exclusively to domestic/peridomestic colonies, ignoring the preferences of these species in natural environments.

To avoid predatory vertebrates in the nests and burrows, Triatominae adopted cryptic behaviors and inverse activity patterns, feeding while the vertebrates are asleep. Most triatomines exhibit typically nocturnal habits in discrete temporal windows, such as responsiveness to host odors early in the night [66, 74]. The heat radiated by warm-blooded animals appears to be the major cue used by Triatominae to locate a host. Besides the thermal cue, carbon dioxide has been shown to attract triatomine bugs similar to almost every blood-sucking arthropod [9]. Other chemical cues related to host detection have also been identified in Triatominae. Among the constituents of vertebrate odor, isobutyric acid, ammonia and aliphatic aldehydes have been shown to play an important role in host location by triatomines [47, 109].

4. CONCLUSIONS

The common pattern of *T. cruzi* vectorial transmission in humans is being supplanted

by emerging patterns involving sylvatic triatomines that display synanthropic trends. In terms of wild animal hosts, an increase in the population of highly competent mammalian reservoirs is currently observed. In both cases, unfavorable environmental changes and the subsequent rarefaction that occurs in wild fauna diversity favor these processes. These epidemiological modifications are inducing new specific problems such as challenges in the assessment of new transmission risks, the possible emergence of an atypical physiopathology of the disease linked to infection by sylvatic parasite strains and, finally, the need for adjustments of the traditional control approaches and entomological vigilance. The currently available data show a great genetic diversity within *T. cruzi* and the existence of very divergent lineages, some of them associated with domestic cycles. Furthermore, all *T. cruzi* lineages and sublineages reported so far have been found in wild cycles. These cycles could act as a source of parasite for domestic transmission. In spite of the successful control of the intradomiciliary transmission of *T. cruzi*, the diversity of the infection mechanisms in addition to the numerous vector and mammalian host species makes the maintenance of sustainable surveillance activities a necessity. This will only be possible if all components of the transmission network are identified.

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