Attempts to Improve Xenodiagnosis: Comparative Test of Sensibility Using *Rhodnius neglectus*, *Panstrongylus megistus*, *Triatoma vitticeps* and *Triatomad infestans* in Endemic Areas of Brazil

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From June 1984 to July 1992, 392 xenodiagnostic tests were applied on 264 patients with chronic Chagas disease from Brazilian endemic areas of Virgem da Lapa and Coronel Murtiá situated in the Jequitinhonha Valley, in the State of Minas Gerais.

The susceptibilities of *Rhodnius* neglectus, *Panstrongylus megistus*, *Triatoma vitticeps* and *Triatoma infestans* were compared.

Most of the time 20 nymphs (fourth instar) of each species were applied to 161 women and 103 men aged between 5 and 83 years of age. The tests were prepared to compare the susceptibilities of two species at a time, using the same patients for each test.

Results showed a xenopositiveness of 26.28% (103 tests) being 27.98% in women (68 positive in 243 applied tests) and 23.49% in men (35 positive in 149 applied tests). The relative frequency of xenopositiveness displayed a great superiority of *P. megistus* and *T. vitticeps*. In tests from type I, for example, *P. megistus* was the unique responsible for 10.73% of positive xenodiagnosis vs. only 0.98% in *T. infestans*. Other parameters analyzed in this work confirm this superiority, and corroborate that *T. infestans* can be replaced by *P. megistus* and/or *T. vitticeps* in order to upgrade the efficacy of xenodiagnosis.

**Key words:** Chagas disease • xenodiagnosis • insect susceptibility • *Rhodnius neglectus* • *Panstrongylus megistus* • *Triatoma vitticeps* • *Triatoma infestans* • Brazil

Chagas disease represents the main cause of morbidity and mortality in Latin America (*sensu lato*).

Up to now, xenodiagnosis used in screening chagasic patients, confirming the diagnosis of seropositive individuals, treatment control and stock isolation, is the most specific and sensitive technique currently available for diagnosis. However, this well documented technique described by Brumpt (1914) is considered a low sensibility method. In fact, the index of serologically positive patients is confirmed to be higher than the para- and/or serologically positive ones (Avilla et al. 1993).

Segura (1987) demonstrated that xenodiagnosis applied to positive chronic patients produced positive results in only 17 to 70% of the cases. This phenomenon can be explained by the fact that in the majority of cases, triatomine bugs were fed in chronic patients having low numbers of circulating parasite forms.

Various studies showed different results depending on the species used (Cerisola et al. 1971, Schenone et al. 1974, Alvarenga & Bronfen 1984, Bronfen et al. 1984, Bronfen 1989, Pereira et al. 1993). Questions arose from these studies about the vector reliability as xenodiagnostic agents.

Some authors had already predicted that the efficiency of xenodiagnosis could be increased by using large numbers of bugs per patient (Meckelt 1964, Schenone et al. 1974, Cerisola et al. 1974); by repeatedly examining the patients (Castro et al. 1983) and by using significant epidemiological vectors in the same area where the patient lives (Dias 1940, Ryckman 1965, Little et al. 1966, Zeledorí 1974, Minter et al. 1978). However, interpreting results is difficult due to lack of standardization.

Cerisola et al. (1975) suggested for the first time the standardization of xenodiagnostic procedures, recommending the use of 40 third instar nymphs of *Triatoma infestans* as this species is associated with the domestic transmission of Chagas disease.
in Brazil. The above preference agreed with the polemic statement by Dias (1940), attributing greater susceptibility of the indigenous vector from the same geographical area as the patients. The exam would be performed after a 60 day post infection period (p.i.). Cerisola made efforts to minimize the large number of variables that could have affected the results such as the species utilized, methodology of application and examination.

P.-Szumlewicz and Muller (1987) described a variety of species which were identified as successful xenodiagnostic agents in testing hosts with chronic infection by Y. Trypanosoma cruzi strain (Panstrongylus megistus, T. pseudomaculata, T. rubrovaria, Rhodnius neglectus). This species could make xenodiagnosis more efficient and reliable.

Bronfen (1989) suggested the simultaneous use of at least two species of triatomine bugs associated with xenoculture and bloodculture for strain isolation and diagnosis respectively.

P.-Szumlewicz et al. (1988, 1990) emphasized the superiority of silvatic vectors over those from residences as xenodiagnostic agents in experimental hosts with acute and chronic infections caused by seven strains of T. cruzi, establishing that susceptibility varied according to the species used, giving top priority to P. megistus as an xenodiagnostic agent.

The results obtained from this study indicate the superiority of P. megistus, T. rubrovaria and T. vitticeps exhibiting the best rates of positivity and density of parasites, followed by T. pseudomaculata and R. neglectus. In the first place, we tested P. megistus, T. vitticeps and R. neglectus against T. infestans and P. megistus against T. vitticeps.

**MATERIALS AND METHODS**

*Xenodiagnosis* - Sixteen experiments were performed from June 1984 to July 1992 according to the following scheme: 9 tests between *P. megistus* and *T. infestans* (type I); 3 tests between *T. vitticeps* and *T. infestans* (type II); 2 between *R. neglectus* and *T. infestans* (type III) and 2 between *P. megistus* and *T. vitticeps* (type IV).

The tests were intended to compare the susceptibilities of two species at a time, using the same patients for each test. Some patients were tested more than once at different times.

The nymphs were divided in wooden boxes and covered with cheese clothes (10 nymphs/box) and kept without food for three weeks; 40 fourth instar nymphs (20 of each different species tested) were fed for 30 min on each patient’s forearm, except in the first and second tests between *P. megistus* and *T. infestans* where 10 and 20 respectively were used for the former and 30 for *T. infestans*.

Fifteen days after feeding, the bugs received a supplementary feeding of chicken blood.

A total of 15,600 bugs were utilized in experiments. From this total 13,281 were examined, 6,503 for *T. infestans* (total of 14 tests), 3,788 for *P. megistus* (11 tests), 2,318 for *T. vitticeps* (5 tests) and 672 for *R. neglectus* (2 tests).

The examination started after a 45 day post infection period employing optical microscopy (400X) of the suspension of the entire digestive tract which had been ground into a physiological saline. The suspension was prepared after cutting the extreme rear portion of bugs’ abdomen, pulling out the digestive tract and grinding it in two or three drops of physiological saline. Thereafter two or more wet films were made to complete the exam.

Dead dried nymphs were not examined.

A test of difference of proportions (p <0.01) was used for statistical analysis.

*Patients* - A total of 161 women and 103 men were tested. The patients were from endemic areas of the Jequitinhonha Valley, in the State of Minas Gerais. In the municipality of Virgem da Lapa 254 patients (16 tests) were tested and in Coronel Murta 10 patients (1 test). The age of patients varied from 5 to 83 years (38.5 ± 16.8 years).

Only 24 patients proved negative for IgG against *T. cruzi*, shown in immunofluorescent tests and indirect blood aglutination.

**RESULTS**

From a total of 392 applied xenodiagnosis 26.28% were positive. In 161 women 243 tests were applied and in 103 men 149 xenos with positive rates of 27.98% and 23.49% respectively. These rates did not show significant differences (p<0.01). From 161 female patients 52 were positive (32.30%) and from 103 male patients 33 were positive (32.03%). These rates did not show significant differences (p<0.01) either.

*Xenopositiveness* rates are presented in Table I. In tests from type I there were significant differences (p<0.01) among rates of *P. megistus* (10.73%) and *T. infestans* (0.98%). Table I also shows the rates produced when both species tested were positive. The unique case where the percentage of positive patients from the two species tested was statistically inferior to the rates of species tested separately, occurred in tests from type IV where *P. megistus* and *T. vitticeps* positive patients produced the rate of 32.35%, significantly higher than the rate of *T. vitticeps* (5.88%).

The nymphs’ positive reaction to *P. megistus*, *T. vitticeps* and *R. neglectus* are compared with *T. infestans* (Table II). There were significant variation (p<0.01) between the rate of 4.59% by nymphs of *P. megistus* and 32.6% in *T. vitticeps* positive patients.
Amblvoomma cayennense | micrographs of leg II of larva. Fig 1: distal portion | Tibia: BTA basitarsus; TTA telotarsus. Arrow: slit cuticular structure. Fig 2: detail of slit cuticular structure. PTA: pretarsus.

Amblvoomma cayennense | electron micrographs of leg II. Figs 3, 4: larva. Figs 5, 6: nymph. Figs 7, 8: adult. Arrow: slit cuticular structure; TTA telotarsus. BTA basitarsus.
The contamination rates from each species separately (overall) and the mortality rates are presented in Table IV. The former varied from 1.55% in *T. infestans* (101 in 6,503 examined) to 6.13% in *P. vitticeps* (142 positive in 2,318 examined). There were significant differences (p < 0.01) between the rates of *T. vitticeps* (6.13%) and *P. megistus* (5.44%) in comparison to *T. infestans*’ rate (1.55%). The rates of mortality varied from 13.52% in *P. megistus* (592 dead in 4,380 sent to endemic area) to 18.95% in *T. vitticeps* (542 dead in 2,860 sent). The rate demonstrated by *T. vitticeps* is significantly higher than the rate in *P. megistus* (p < 0.01).

Only one patient (woman) negative in serology exhibited positive xenodiagnosis with *P. megistus* (0.38%).

TABLE IV

<table>
<thead>
<tr>
<th>Vector species</th>
<th>Mortality (overall)</th>
<th>Infectivity (overall)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Triatoma infestans</em></td>
<td>13.98 (7560)</td>
<td>1.55 (6503)</td>
</tr>
<tr>
<td><em>Panstrongylus megistus</em></td>
<td>13.52 (4380)</td>
<td>5.44 (3788)</td>
</tr>
<tr>
<td><em>Triatoma vitticeps</em></td>
<td>1X.95 (2860)</td>
<td>6.13 (2318)</td>
</tr>
<tr>
<td><em>Rhodnius neglectus</em></td>
<td>16.00 (800)</td>
<td>1.64 (672)</td>
</tr>
</tbody>
</table>

a: number in parenthesis indicates the number of nymphs sent to endemic areas
b: number in parenthesis indicates the total number of examined nymphs

DISCUSSION

Xenodiagnosis has been well documented in literature. Its low sensibility and divergent results obtained, raised questions about the efficacy of this technique. According to some investigators it could be increased by the use of large numbers of bugs per patient, by repeated exams (previously mentioned) and by the use of vectors capable of supporting rapid development and vigorous multiplication of the few parasites ingested (P.-Szumlewicz & Muller 1987).

Aiming to improve xenodiagnosis, comparative works were performed to search for a suitable insect model with the above mentioned characteristics (Alvarenga et al. 1977, Barreto et al. 1978, Minter et al. 1978, Marsden 1986, Pereira et al. 1993).

Experimental studies in acute and chronic Chagas disease were developed by P.-Szumlewicz and Muller (1982, 1987) using Y strain of *T. cruzi*. The authors demonstrated that efficiency could be associated with the triatomine species used, enumerating factors of parasite density, giving an accurate portrayal of parasite development. This should be incorporated in screening bugs for xenodiagnosis. They also listed some measures to achieve a reliable xenodiagnostic procedure: the importance of biweekly supplementary feeding (after infection) that could triple the population density of parasites in infected bugs minimizing the probability of recording a false negative result, and stipulated a routine of 45 day post-infection interval in the examination of bugs.

Later, P.-Szumlewicz et al. demonstrated the superiority of silvatic vectors over residential ones in acute (1988) and chronic (1990) infections from seven strains of *T. cruzi*, calling attention to the phenomenon of vector/parasite relationship, showing the species’ capability to respond to chronic infections using different parasite stocks and indicating the most efficient vectors within species tested: *P. megistus*, *T. rubrovaria* and *T. vitticeps*. In this study the rates of infected nymphs for *P. megistus* and *T. vitticeps* were significantly higher than for *T. infestans*. These results are in accordance with those obtained by P.-Szumlewicz et al. (1990) and Pereira et al. (1993, 1996).

In spite of these results various authors continued recommending the use of *T. infestans* in xenodiagnosis. Chiari (1992) in the “Relatório da VIII Reunião de Pesquisa Aplicada em Doença de Chagas, Uberaba, MC”, suggested the use of first and third instar nymphs of *Dipetalogaster maximus* and *T. infestans*, respectively. The enthusiasm for *D. maximus* as a potential xenodiagnostic agent has been linked to its large size which allows the ingestion of great quantities of infected blood, but up to the moment there is no confirmed relation between infection rates, high density rates of parasites and great quantities of imbebed blood, in chronic Chagas disease.

P.-Szumlewicz et al. (1988) demonstrated that the third instar nymphs of *T. infestans* could ingest more blood than the first instar nymphs of *D. maximus* and that, maintenance of a *P. megistus* colony in the laboratory is easier (fast development, slow locomotion, low mortality and easy to breed under changing environments).

Nirschl et al. (1994) analized the susceptibility of *T. infestans* infected by seven strains of *T. cruzi* derived from chagasic patients from Triângulo Mineiro, MG, and concluded that this species had low susceptibility, which could explain the low rates of infectivity in xenodiagnosis in the same geographical area.

Results from Tables I-III confirm the existence of significant differences (p < 0.01) in rates from xenopositiveness, contagion of nymphs individu-
ally and general contagion of nymphs (overall) among three species tested in this work. These results demonstrated that the nymphs of *T. infestans* could be replaced by nymphs of *P. megistus* and/or *T. vitticeps*, the best hitherto tested in endemic areas, by our laboratory, in order to upgrade the reliability of xenodiagnosis.

Pereira et al. (1996) also recommended the replacement of *T. infestans* nymphs by *P. megistus*, which according to the authors increased considerably the xenopositivity in nymphs on the fourth stage. Vector resistance could be a parameter for the species' choice. Data shown in Table IV demonstrate that *P. megistus* has a mortality rate significantly lower than *T. vitticeps*.

Additional data from Table I indicated that the association between two different species contributes, in some cases, to the improvement of results. These improvements appear to be species dependent, related to the phenomenon of parasite specificity in parasite/vector relationship. We think that species considered good vectors with different sensitivities to infection from strains of *T. cruzi*, should be used, therefore becoming indispensable to screen a significantly large number of good vector species in order to justify the generalization of these procedures.

It was also observed that the positiveness in nymphs of *P. megistus* and *T. infestans* in the positive xenodiagnosis of both species was significantly higher than that positive result obtained exclusively from these species separately, which leads us to think about the possibility of the existence of *T. cruzi* subpopulations with different degrees of development in both species.

In the near future xenodiagnosis, which lacks sensibility, could be replaced by K-DNA PCR (Polymerase Chain Reaction Amplification of Kinetoplast Minicircle DNA) in direct parasitological evaluation of chronic chagasic patients. This technique allows characterization of *T. cruzi* nucleotide sequences and the replication of synthetic chains giving support to the identification of the parasite with high degrees of sensitivity (Avila et al. 1991, Romana 1992). According to Avila et al. (1993) this new technique has 100% sensitivity compared with serologic tests. Maybe the application of xenodiagnosis could be restricted only to strain isolation. Even so, the use of different species used until now have not given satisfactory results (for example *T. infestans*). Another that can produce better results must be found to enhance the sensibility of xenodiagnosis and consequently its quality. Careful studies on vector-parasite relationship are also urged which require the use of biochemical methods.

**ACKNOWLEDGEMENTS**

To Dr Jose Borges Pereira for testing our triatomine species in his patients, to Shirley Maria Cordeiro for typing the manuscript and to Prof. Lídia de Andrade Ribeiro for her assistance in English revision.

**REFERENCES**


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