Review on the Molecular Tools for the Understanding of the Epidemiology of Animal Trypanosomosis in West Africa

G Duvallet, S de La Rocque, JM Reifenberg, P Solano, T Lefrançois, JF Michel, Z Bengaly, I Sidibe, D Cuisance, Cuny

CIRAD-EMVT, B.P. 5035, 34032 Montpellier Cedex 1, France *CIRDES, B.P. 454, Bobo-Dioulasso, Burkina Faso **ORSTOM, B.P. 5045, 34032 Montpellier Cedex 1, France

The epidemiology of animal trypanosomosis around Bobo-Dioulasso (Burkina Faso, West Africa) benefited a lot in the last years from the progress of molecular tools. The two most used molecular techniques were the polymerase chain reaction (PCR) for the diagnosis of the disease in cattle and the characterization of the trypanosomes in the host and the vector; and the microsatellite DNA polymorphism in tsetse flies to study the intraspecific genetic variability of the vector on the other hand. The results obtained in the Sideradougou area during a recent two year survey with these techniques, associated with many other georeferenced informations concerning vector and cattle distribution, natural environment, landuse, ground occupation, livestock management, were combined in a Geographical Information System. This new approach of a complex pathogenic system led to a better evaluation of the risk of trypanosome transmission.

Key words: African animal trypanosomosis - epidemiology - polymerase chain reaction - microsatellite DNA - Glossina - Trypanosoma sp.

Animal trypanosomosis represents a serious problem in most of Africa and also in certain zones of South America and Asia. The control methods currently in use require a good knowledge of the epidemiology of the disease for a better effectiveness.

The epidemiology of animal trypanosomosis around Bobo-Dioulasso (Burkina Faso) benefited a lot in the last years from the progress of molecular biology tools. The most employed techniques were (1) the polymerase chain reaction (PCR) for the diagnosis of the disease in cattle and the characterization of the trypanosomes in the host and the vectors; (2) the microsatellite DNA polymorphism in tsetse flies for the study of the intraspecific genetic variability of the vectors.

The PCR combines the advantages of a greater sensibility than the parasitological or immunological techniques with the possibility of differentiating trypanosomes which have a similar morphology but a very different economic impact.

This work was funded by CIRAD-EMVT (Action thématique programmée ‘Santé-Environnement’), CNRS (Programme Interdisciplinaire ‘Environnement, Vies et Sociétés’), AUPELF-UREF (LAF no. X7.10.02/4581-306) and CIRDES.

*Corresponding author. Fax: +33-4-6759.3797. E-mail: duvallet@cirad.fr

Received 12 November 1998
Accepted 19 January 1999

The study of the microsatellite DNA polymorphism in tsetse flies appears to be of prime importance to assess the consequences of intraspecific genetic variability on the epidemiology of trypanosomosis.

All the results obtained in the Sideradougou area during a recent two year survey with these techniques, associated with many other georeferenced informations concerning vector and cattle distribution, natural environment, landuse, livestock management, were combined in a Geographical Information System (GIS). This new approach of a complex pathogenic system must lead to a better evaluation of the risk of trypanosome transmission.

MATERIALS AND METHODS

Area - Sideradougou area is a pastoral zone situated in the southwestern part of Burkina Faso, where a multidisciplinary work aims to identify the most important factors determining the presence of tsetse flies which transmit the trypanosomes to cattle (de La Rocque & Cuisance 1997). It lies in the south sudanian savannah zone (4°20'W, 10°55'N). More than 120 km of forest galleries were prospected in 1996 and 1997 for the catching of tsetse flies with biconical (Challier & Laveissiere 1973) traps.

Tsetse flies - A total of 2,211 non teneral Glossina tachinoides and G. palpalis gambiensis were dissected among 4,357 captured flies. The instruments of dissection were carefully cleaned between
The study combined field survey and high-resolution remote sensing data (SPOT images). The following different layers of information (all georeferenced) have been entered into a GIS: entomology (1 trap every 100 m along a 120 km hydrographic network), parasitology (standard methods and PCR results) from vector and cattle (sentinel herds), natural environment (70 parameters concerning vegetation, waterstreams and frequency of use), valley-type landuse, ground occupation and its dynamics (agricultural fragmentation), cattle distribution (exhaustive census, producer typology), and the different pastoralist customs (livestock range management, drinking places). All the details are in de La Rocque (1997).

RESULTS

PCR

In tsetse flies - Among the 2,211 non teneral G. tachinoides and G. p. gambiensis dissected, the classical parasitological techniques gave the following results: (1) total infection rate: 14.6% in G. tachinoides, 12.1% in G. p. gambiensis (not statistically different); (2) Nannomonas type infection (proboscis + midgut): 13.6% in G. tachinoides, 7.4% in G. p. gambiensis (not different); (3) immature infection (midgut alone): 39% in G. tachinoides and 68.6% in G. p. gambiensis (statistically different); (4) Duttonella type infection (proboscis alone): 45.8% in G. tachinoides and 21.5% in G. p. gambiensis (statistically different).

A total of 298 positive flies were submitted to PCR and the comparative results were the following:

- 72.7% of Nannomonas type infection could be characterized by PCR with: (1) 54.5% confirmed as Nannomonas (with 21.2% of mixed infections); (2) 15.2% identified as T. vivax; (3) 3% identified as mixed T. vivax/brucei.
- 15.8% of immature infections only could be characterized with PCR with: (1) 11.2% identified as T. congolense savannah type; (2) 3.9% identified as T. congolense forest type; (3) 3.9% identified as T. brucei.
- 87.9% of Duttonella type infection could be characterized with PCR with 86% confirmed as T. vivax.
- T. simiae has not been identified during this survey.

In cattle - Among the 120 cases of infection identified in cattle, 90% could be characterized by PCR. It is important to note that 50% of parasitologically negative animals with a haematocrit value under 25% were identified as positive with PCR.

Among the PCR positive animals: 63.3% were positive for T. congolense savannah type; 2.5% were positive for T. congolense forest type,
40% were positive for *T. vivax*; 9.2% were positive for *T. brucei*. 14.2% had mixed infection. 

*T. congolense* riverine forest type was found only in three parasitologically negative animals with a haematocrit under 25%. One of them had *T. congolense* forest type alone, and the other two had both types of *T. congolense*.

**Bloodmeal analyses** - A total of 233 midgut residues were analyzed for bloodmeal origin. Among all the results (Lefrançois et al. 1998) comparing tsetse species, sex and geographic position in the area, we retain here that (1) bloodmeals on ruminants were more frequent in *G. tachinoides* (46.3%) than in *G. p. gambiensis* (19.3%); (2) on the opposite, bloodmeals on reptiles were more frequent in *G. p. gambiensis*; (3) bloodmeals on reptiles were more frequent in males than in females, and in the western part of the area than in the eastern part.

**Microsatellite markers** - The study of the microsatellite DNA polymorphism of the *G. p. gambiensis* caught in the area showed at least two genetically different populations on the same hydrographic network. It is, to our knowledge, the first demonstration of a genetic variability of tsetse flies on a short distance (13 km). These two populations seem to be adapted to two different eco-epidemiological situations. These results are still being evaluated (Solano 1998).

**GIS**

The association of all the layers of information into a GIS led to a classification of favorable biotopes. For example, *G. tachinoides* was more abundant in large and open forest galleries, and *G. p. gambiensis* more abundant in dense and closed galleries. *G. tachinoides* was more abundant in sites frequented by man and domestic animals (pigs, goats and sheep); *G. p. gambiensis* in sites frequented by cattle or wild antelopes. Many other parameters have been taken into account to analyze the special differences in vector localization, type of trypanosome infection and hosts (de La Rocque 1997).

To briefly summarize the results, we could say that two different situations at least have been identified in the Sideradougou area for the *G. p. gambiensis* (de La Rocque 1997, Solano 1998):

- the Nyarafo situation in the western part with (1) high proportion of tsetse bloodmeals on reptiles; (2) high total infection rate of tsetse flies; (3) many cases of infection not identified by PCR with the primers used; (4) high infection rate of midgut.
- the Yeguere situation in the eastern part with (1) high proportion of tsetse bloodmeals on ruminants; (2) low total infection rates; (3) high infection rate of proboscis identified as *T. vivax* by PCR.

**DISCUSSION**

Molecular tools for the diagnosis and the epidemiology of animal trypanosomiasis have been introduced several years ago at CIRDES (Duvallet & Toure 1994, Reifenberg 1996, Reifenberg et al. 1997b). At the beginning, because of its cost, the PCR was limited to epidemiological studies in situations well known with classical methods. Now, with the decrease in the price of reagents and the possibility to use very low volumes and to run many samples at a time, the PCR can be envisaged as a diagnosis tool.

The high sensitivity of the PCR is illustrated in the Sideradougou survey by the fact that 90% of parasitologically infected and 50% of parasitologically negative animals (with classical tests) but with a haematocrite value under 25% were diagnosed as positive. *T. simiae* has not been identified in the Sideradougou area during this survey, but had been reported from other parts of Burkina Faso (Solano et al. 1996, Reifenberg et al. 1997a).

The PCR also made it possible to detect mixed infections and to characterize some of the immature infections. It is important to note that up to 45.5% of the presumed (classical parasitological tests) *Nannomonas* infections in tsetse flies were identified by PCR as *T. vivax*, *T. brucei* infections or were not identified.

Majiwa et al. (1994) already reported positive PCR results on 12.5% antigenemic but parasitaemic animals.

The correlation between the low identification rate by PCR in the midgut infection (even after DNA extraction), the higher proportion of reptile bloodmeals especially for *G. p. gambiensis*, led us to hypothesise that the midgut infections came from reptile trypanosomes. Furthermore when bloodmeal analysis could be conducted on PCR non-identified midgut infected flies, 75% were from a reptile origin (Lefrançois et al. 1998).

The study of the genetic variability of the tsetse flies using microsatellite DNA polymorphism markers is still in progress at CIRDES (Solano 1998).

The integration of PCR results (characterization of the parasites), with bloodmeal origins, microsatellite markers, and many different environmental factors led to the identification of different eco-epidemiological situations. This new approach of a complex pathogenic system led to a
ACKNOWLEDGMENTS

To Dr SM Toure, Director general, and all the technical staff of CIRDES.

REFERENCES


MEMÓRIAS DO INSTITUTO OSWALDO CRUZ
An International Journal of Biological and Biomedical Research