Markers for the population genetics studies of *Triatoma sordida* (Hemiptera: Reduviidae)

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**Abstract**

**Background:** *Triatoma sordida*, a vector of *Trypanosoma cruzi*, is native of Brazil, Bolivia, Paraguay, Argentina, and Uruguay, and occurs primarily in peridomiciles. Currently, it is the species most frequently captured by the Chagas Disease Control Program in Brazil. For this reason, population genetic studies attract great interest, as they can provide further information about the dispersal and household invasion processes of this species. In the absence of suitable markers, the objective of this study was to test the cross amplification of microsatellite primers.

**Findings:** 23 primers were tested for microsatellite loci already described for other species of the genus *Triatoma* sp. Forty four specimens of *T. sordida* captured in the north of Minas Gerais were used to validate the use of standardized loci for population genetic analyses. It was possible to amplify 10 of the 23 loci tested for *T. sordida*.

**Conclusions:** This is the first study that provides 10 microsatellite markers for population analysis of this triatomine species. Cross-amplification of primers can be used among other phylogenetically related species whose loci are already available for study.

**Keywords:** Triatominae, *Triatoma sordida*, Microsatellites, Population genetic, Chagas disease

**Findings**

**Background**

*Triatoma sordida* (Stål, 1859) is the triatomine species most frequently captured by Brazil’s Chagas Disease Control Program [1] and in neighboring countries [2]. It occurs in Bolivia, Paraguay, Argentina and Uruguay [2], and Brazilian Cerrado [1]. This vector has broad ecological valence, and so it can live in several ecotopes and use different food sources [3]. It shows high rates of active dispersal [4], and it can also be passively introduced into artificial environments, possibly in firewood piles transported from forests to households, or from one household to another [5], or attached to bird feathers [3]. In an artificial environment, *T. sordida* is often associated with peridomiciles; in addition, it uses birds as its preferred hosts. Thus, this vector has secondary epidemiological importance [1].

Microsatellites are established as a valid technique to study diversity for more than a decade. Because they have highly polymorphic loci, they are widely used to investigate the genetic structure of natural populations. These genetic markers possess high reproducibility, multiallelic nature, codominant inheritance, abundance and wide distribution throughout the genome [6].

The use of microsatellites has not been widely applied in triatomines yet, but they are very sensitive markers and yield promising results in the population analyses of this subfamily. Microsatellite markers have been identified and characterized for the following triatomines: *Rhodnius pallescens* [7], *Triatoma dimidiata* [8], *Triatoma infestans* [9, 10], *Rhodnius prolrixus* [11, 12], *Triatoma pseudomaculata* [13] and *Triatoma brasiliensis* [14]. Microsatellites described for particular species may sometimes be used to characterize others, e.g., markers for *R. prolrixus* that are used for other species of the genus. However, this requires specific research that is certainly associated with the degree of similarity among species [11].

The analysis of gene flow between populations of the wild environment, intradomiciles and peridomiciles can help understand the factors that favor the infestation/reinfestation of the household, and hence provide guidelines for Chagas disease control programs [15, 16]. Considering
this fact, the lack of microsatellite primers described for *T. sordida*, and also the great effort required for identifying and characterizing such markers, the objective of this study was to test the amplification of microsatellite loci in *T. sordida* using primers described for other triatomine species.

**Methods**

Primers were tested for 23 loci already described for other species of the genus *Triatoma*: *T. dimidiata* [8], *T. infestans* [10], *T. psedomaculata* [13] and *T. brasiliensis* [14]. The insects used in the tests were provided from a mixed colony maintained in the insectarium of the Laboratory of Triatomines and Chagas Disease Epidemiology at the René Rachou Research Center. They were originally captured in several northern regions of Minas Gerais. Genomic DNA was extracted from one of the legs of three *T. sordida* adult specimens using the Wizard Genomic DNA Purification Kit (Promega). Quantitation of the DNA was performed in a NanoDrop® ND-1000 spectrophotometer, and the material was kept at −20 °C until processing.

Polymerase chain reactions (PCR) were performed in a final volume of 10 μL containing: 1 unit of Taq DNA Polymerase, Recombinant (Invitrogen), 1x buffer, 1.5 mM or 3 mM of MgCl₂, 1 mM of dNTP, 5 pmoles of each primer, 2 ng of DNA and ultrapure water. Reactions were performed in an Eppendorf Mastercycles® Gradient thermocycler with the following cycle: initial denaturation at 94 °C for five minutes, 30 cycles at 94 °C for 30 s, temperature gradient dependent on annealing temperature in the primer description (±5 °C) for 30 s and 72 °C for 30 s, followed by final extension at 72 °C for three minutes. The amplified products were visualized in a polyacrylamide gel at 8 % in the mini-gel system (BIO-RAD), stained with 0.2 % silver nitrate.

To determine the size of the loci, new PCRs were performed with forward primers tagged with bioluminescent probe. The PCR products were diluted at 1:10 in ultrapure water and genotyped in a MEGABace (Amersham Biosciences) sequencer. The size of the PCR products was estimated in comparison with a standard size marker (ET-400, GE Health Care), and the genotypes were read using the software Fragment Profiler™.

Specimens from the municipality of Coração de Jesus (16°41′15″S, 44°18′45″W), in the north of Minas Gerais, were used to test the use of standardized loci for *T. sordida*. These specimens were captured by agents of the Chagas Disease Control Program and not part of the mixed colony used for the selection of microsatellites. Forty four specimens of *T. sordida* were caught in the peri-domicile of five neighboring localities: seven in Barriguda,

### Table 1

| Locus   | Primer sequences (5′-3′) T (°C) MgCl₂ 50 mM(%) bp N | bp | N |
|---------|---------------------------------------------------|----|---|
| Tb 830  | F: GTCAGATGCAATGGTGATAC 48 3 119-122(265–292) 4(8) |
| Tb 8112 | F: GAATCCGCTTTCCACAG 54 3 78-80(78–96) 3(6)       |
| Tb 8124 | F: GCCACGTGTGTCTCACTCC 59.5 3 209-248(209–253) 12(10) |
| TDM53   | F: TCAGATGCAACAGGTTGGATG 63 3 137-143(129–146) 3(7) |
| TDM54   | F: CAGTTGTCTCATCAAGGGAATGAATC 54 6 160(150–186) 1(20) |
| Tinfest_ms23 | F: CTCTGCTGTGTTGCTGGACTG 64 6 156-173(148–177) 10(5) |
| Tinfest_ms42 | F: GCCAATTTGGGTTAGTCTATG 66 6 205-224(206–246) 9(15) |
| Tp59    | F: ACTTGAATGAGGATGAA 53 3 121-149(120–128) 9(4)  |
| Tp20    | F: ACTGACTCCGAGAAGTGG 57 3 125-144(170–206) 7(15) |
| Tp544   | F: TGTTAGAATGAGGACTGA 55 3 142-213(148–172) 7(8)  |

T: annealing temperature; bp: allele size range; N: number of alleles. Data in brackets refer to the original description (*Tb*, Harry et al. 2009 [14]; TDM5, Anderson et al. 2002 [8]; Tinfest_ms, Marcet et al. 2006 [10]; Tp, Harry et al. 2008 [13])
nine in Boa Vista, 10 in Bom Jesus, 10 in Jataí I, and eight in Jataí II.

DNA was extracted from the wing muscle of adult insects [16]. Each muscle was homogenized individually in 100 μL of 1X STE solution (0.01 M NaCl, 0.1 M Tris–HCl and 1 M EDTA), incubated at 90°C for 10 min, and centrifuged at 13,000 rpm for one minute, and the supernatant was recovered. The quantitation and storage of DNA were carried out as described hereinabove.

The samples underwent standard PCR, the annealing temperature and the amount of MgCl₂ for each primer are described in Table 1. The primers were tagged with bioluminescent probe; genotyping was also performed with standard conditions. For each locus, calculations were made of a number of alleles using the software Arlequin 3.1.

Results and discussion

Ten of the 23 pairs of tested primers showed satisfactory amplification for *T. sordida* in specific conditions. The size of the alleles obtained was similar to the original descriptions, except for loci Tb830 and Tp20, which were smaller for *T. sordida* (Table 1).

The number of alleles per locus ranged from 1 (TDMSA) to 12 (Tb 8124) with a mean of 6.5. Only TDMS4 locus showed no heterozygous individuals (Table 1).

Harry *et al.* [11] considered that cross-species amplification should be used with caution because microsatellite null alleles may occur. Their presence should be considered when population analyses are performed, as they may underestimate population diversity [17]. This study could validate the use of the 10 microsatellite markers.

Some of the primers described were tested for cross amplification in other species [10, 11, 13, 14], including *T. sordida* [9]. However, this is the first study that provides microsatellite markers for the population analysis of this triatomine species. Cross-amplification of primers among species was a low-cost, effective strategy which was faster than isolation, identification, and development of primers. Thus, it can be used among other phylogenetically related species whose loci are already available for study.

Due to the high sensitivity of the microsatellite markers, this study provides a new tool for the assessment of gene flow between populations of *T. sordida* also in microgeographic scale.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

CJB participated in the design and standardized the study, data analysis, performed the statistical analysis, and drafted the manuscript. GCDP participated in the design, data analysis, and manuscript review. LSD, PFS, and ACLR participated in the samples preparation and processing. LSD also assisted in data analysis. LD conceived and coordinated the study, participated in the design of the study and manuscript review. All the authors read and approved the final manuscript.

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