Polymorphisms and ambiguous sites present in DNA sequences of *Leishmania* clones: Looking closer

Mariana Côrtes Boité *, Taíse Salgado de Oliveira, Gabriel Eduardo Melim Ferreira, Marcos Trannin, Barbara Neves dos Santos, Renato Porrozzi, Elisa Cupolillo

Laboratório de Pesquisa em Leishmaniose, Instituto Oswaldo Cruz (IOC), Fundação Oswaldo Cruz (FIOCRUZ), Rio de Janeiro, Brazil

**A R T I C L E   I N F O**

Article history:
Received 26 August 2013
Received in revised form 9 April 2014
Accepted 13 April 2014
Available online 21 April 2014

Keywords:
*Leishmania*
Polyclonal
Aneuploidy
Heterozygosis

**A B S T R A C T**

In genetic studies of *Leishmania* parasites, co-dominant markers are chosen for their ability to detect heterozygous polymorphisms, to infer the occurrence of inbreeding and to resolve genetic variability. The majority of DNA sequence based reports perform conventional dye terminator cycle sequencing where perfectly ambiguous sites or double peaks in the chromatogram are interpreted as heterozygous strains. However, molecular peculiarities of the parasite such as aneuploidy, mixed populations and homologous recombination advise that data from regular DNA sequence analysis should be carefully evaluated. We report here a closer look at ambiguous sites observed in 6pgd DNA sequences obtained for a multilocus sequence analysis project on *Leishmania* (*Viannia*) strains. After comparing 286 DNA sequences from biological and molecular clones of six *L. (Viannia)* strains we could distinguish events that contribute to genetic variation in *Leishmania* (recombination, mutation, chromosomal mosaics). Also, the results suggest how diversity might not be completely revealed through regular DNA sequence analysis and demonstrate the importance for molecular epidemiology research to be aware of such possibilities while choosing samples for studies.

© 2014 Elsevier B.V. All rights reserved.

1. Introduction

The *Leishmania* parasite is the causative agent of the disease leishmaniasis, which affects millions of people annually and represents the third highest global burden of disease among the neglected tropical diseases. Despite observed morphological homogeneity across most specimens, more than 20 genetically distinct species have been described for the *Leishmania* genus. Phenotypic diversity can be observed between species, even among clones (Garin et al., 2001), which may be a causative agent in the presentation of different clinical forms of the disease and the variation in responses to conventional drug treatment. Therefore, ability to identify *Leishmania* species and to characterize genetic variants could increase physicians’ ability to target treatment and therapy. With that in mind, many approaches have been developed, improved and applied aiming to contribute to population genetics, parasite identification and phylogeny. Among these approaches are multilocus enzyme electrophoresis (MLEE) (Cupolillo et al., 1994) and molecular markers based on DNA sequence analysis, such as multilocus sequence analysis (MLSA) (Zemanova et al., 2007; El Baidouri et al., 2013; Boité et al., 2012) and multi locus microsatellite typing (Alam et al., 2009; Oddone et al., 2009; Rougeron et al., 2010).

Traditionally, co-dominant markers used for these genetic based analyses are chosen by their ability to detect heterozygous polymorphisms, infer the presence of inbreeding, and determine genetic variability. The majority of DNA sequence based protocols utilize conventional dye terminator cycle sequencing, which assume standard procedures for interpreting inconsistencies in results, such as double vs single peaks. In MLSA sequences, double peaks in the chromatogram are normally considered to result from heterozygous strains. However, four major aspects of *Leishmania* biology raise concern over the interpretation of such DNA sequence based analysis. First, *Leishmania* presents mosaic aneuploidy in vitro (Sterkers et al., 2010; Rogers et al., 2011), which has also been detected in clinical isolates (Rogers et al., 2011). Mosaic aneuploidy creates cells with different karyotype contents and may result in overlapping the reads of different nucleotides, actually present in different cells, leading to misinterpretation or loss of information in the sequence analysis. Next, the occurrence of polyclonality and infrapopulation mixture in *Leishmania* isolates could cause the observation of more than one sequence in the
chromatogram. This can occur simply by natural mixing or can be created by karyotype diversity (Dujardin et al., 2007) generated after asymmetric chromosomal allotment (ACA) (Sterkers et al., 2012). Third, clonal propagation is considered the predominant mode of reproduction of this parasite. This phenomenon may overlook cryptic sexuality and could explain the presence of the double peak rather than being representative of the clone. Finally, the clonal concept in *Leishmania* is more plausible from a biological point of view rather than a genetic one (Tibayrenc et al., 1990; Tibayrenc and Ayala, 2012).

Given that misinterpretation of sequencing results may have an important impact on all *Leishmania* typing and characterization studies, we investigated ambiguous sites represented by double peaks in 6pgd DNA sequences obtained by MLSA for *Leishmania* (*Viannia*) strains (Boité et al., 2012) to observe how the above described parasite characteristics may interfere in DNA sequences generated for genetic variability analysis.

2. Material and methods

2.1. Strains, biological cloning and MLEE characterization

Six *L. (Viannia)* strains (Table 1) presenting double peaks in previously obtained 6pgd DNA sequences (Boité et al., 2012) were retrieved from the Coleção de *Leishmania* do Instituto Oswaldo Cruz (CIOC – WDCM 731). More details about the samples can be obtained by accessing the CIOC catalogue (http://clioc.fiocruz.br). The parasites were cultivated in Schneider’s medium and the culture volume divided for both procedures: biological cloning and direct DNA isolation. For biological cloning 10^6 parasites in 600 μl of Schneider’s medium were added to 2.4 ml of Low Melting Point Agarose (Sigma) 1% warmed solution. The mixture was distributed over a Petri Plate containing NNN medium. After 7 days, the colonies were harvested and cultured separately in tubes containing biphasic medium (NNN + Schneider’s) and processed for DNA isolation and MLEE assay. Enzymatic MLEE systems were assayed for the characterization of biological clones as previously described (Cupolloilo et al., 1994).

2.2. Molecular procedures and analysis

DNA purification, PCRs for the 6pgd locus and PCR product purification was carried out as described by Boité et al. (2012). Sequences were obtained from two parasite states: non-cloned culture (NC) and culture from biological clone (BC) as pointed in Fig. 1; and through two distinct approaches: PCR products subjected to direct sequencing (DS) and to molecular cloning followed by sequencing (MC) (Fig. 1A). These conditions combined generated the following DNA sequences: non-cloned directly sequenced (NCDS), biological clones directly sequenced (BCDS) and biological clones subjected to molecular cloning (BC_MC) (Fig. 1B).

![Fig. 1. Procedures to obtain the DNA sequences compared in the present study.](Image)

**Table 1.** *Leishmania* (*Viannia*) strains analyzed for a fragment of 6pgd DNA sequence of 716 bp, including species identification as determined by MLEE, number of biological clones and molecular clones, different alleles (sequence types) obtained and type of event observed.

<table>
<thead>
<tr>
<th>Species by MLEE</th>
<th>IOC/L</th>
<th>Total of DNA sequences compared</th>
<th>BCDS</th>
<th>MC</th>
<th>Sequence types</th>
<th>Events observed</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>L. braziliensis</em></td>
<td>918</td>
<td>50</td>
<td>8</td>
<td>40</td>
<td>10</td>
<td>Hot spot sites</td>
</tr>
<tr>
<td><em>L. braziliensis</em></td>
<td>2538</td>
<td>52</td>
<td>9</td>
<td>41</td>
<td>9</td>
<td>Homologous recombination</td>
</tr>
<tr>
<td><em>L. braziliensis</em></td>
<td>2823</td>
<td>134</td>
<td>20</td>
<td>112</td>
<td>14</td>
<td>Infrapopulation of nonequivalent heterozygous cells</td>
</tr>
<tr>
<td><em>L. guyanensis</em></td>
<td>2966</td>
<td>7</td>
<td>5</td>
<td>0</td>
<td>2</td>
<td>ACAs; polymorphisms</td>
</tr>
<tr>
<td><em>L. lainsoni</em>/L. naiffi</td>
<td>2490</td>
<td>23</td>
<td>4</td>
<td>17</td>
<td>3</td>
<td>Monoclonal heterozygous</td>
</tr>
<tr>
<td><em>L. guyanensis</em></td>
<td>2057</td>
<td>20</td>
<td>4</td>
<td>15</td>
<td>2</td>
<td>ACAs</td>
</tr>
</tbody>
</table>

MLEE = multilocus enzyme electrophoresis; BCDS = biological clones directly sequenced; MC = molecular clones; ACAs = asymmetric chromosomal allotments.

* Including the previous and recent NCDS.

* Sequence types were determined considering a 6pgd gene region as described previously (Boité et al., 2012).
that mosaic aneuploidy generated by ACA occurring during parasite culture played an important part in the interpretations of results. These results have a considerable impact on past and future molecular epidemiologic studies of leishmaniasis since we can no longer assume heterozygosis as the unique cause of ambiguous sites in sequencing based methods.

3.1. Loss of ambiguous site by asymmetric chromosomal allotments (ACA)

The double peak previously detected for two strains IOC/L 2957 and 2966 in site 602 was no longer observed, and NCDS and BCDS were identical, suggesting a monoclonal population (Table 2). The loss of heterozygous cells and the alternative allele once detected can be explained by ACAs during the axenic culture (Sterkers et al., 2012). The clonal propagation of the parasites presenting the homozygous chromosomal arrangement might have leaden to the detection of only DNA sequences without the double peaks in the chromatogram. It is not possible, however, to be certain about ploidy of the cell since ACAs explain the monomorphic DNA sequences among BCDS and MC, but ACAs also generates karyotype diversity.

In addition to the double peak site, polymorphisms were noticed between the previous sequence and the new sequences (sites 640, 672, 685, Table 2). These polymorphisms could be the result of point mutations during the culture procedure. Double peak and polymorphic sites, associated with preponderant clonal evolution (PCE) (Tibayrenc and Ayala, 2013) represents one more complicating factor in Leishmania population structure analysis; nevertheless some strictly sexual organisms might also challenge models dependent on whether ability to prevent whole chromosome elimination is present (Benatti et al., 2010).

3.2. Less frequent alleles can only be detected after molecular cloning in conventional dye terminator sequencing methods

The previously observed double peak for the strain IOC/L 2823 represented an equivalent proportion of nucleotides (1:1) but

<table>
<thead>
<tr>
<th>IOC Sequence origin (origin)</th>
<th>Site position</th>
<th>T A C C T A T A T T T Y A A A A A A</th>
</tr>
</thead>
<tbody>
<tr>
<td>IOC/L 2957 Previous NCDS: (NCDS)1; (BCDS)4: (MC)17</td>
<td>R</td>
<td>A</td>
</tr>
<tr>
<td>L 2957 Previous NCDS: (NCDS)1; (BCDS)4: (MC)17</td>
<td>A</td>
<td>G</td>
</tr>
<tr>
<td>L 2823 Previous NCDS: (NCDS)1; (BCDS)4: (MC)17</td>
<td>T</td>
<td>C</td>
</tr>
<tr>
<td>L 2538 Previous NCDS; (BCDS)4</td>
<td>R</td>
<td>S</td>
</tr>
<tr>
<td>L 2490 Previous NCDS; (NCDS)1; (BCDS)4</td>
<td>R</td>
<td>K</td>
</tr>
<tr>
<td>L 918 Previous NCDS; (NCDS)1</td>
<td>T</td>
<td>G</td>
</tr>
</tbody>
</table>

Table 2: Sequence polymorphisms observed in a 6pgl gene region for non-cloned culture, and for biological and molecular clones of six Leishmania strains.

"Previous NCDS" = sequence from Boité et al. (2012); NCDS = non cloned directly sequenced; BCDS = biological clone directly sequenced; BC_MC = molecular clones of a biological clone; R = A (adenine) or G (guanine); Y = C (cytosine) or T (thymine); S = G or C; K = G/T.
was no longer identified among BCDS (Table 2, site 460); nonetheless one MC presented the alternative nucleotide “C”. From that is possible to infer a change has occurred in allele’s proportion, and only biological and molecular cloning allowed both nucleotide possibilities to be detected.

The heterozygous concept applied for *Leishmania* is the 1:1 nucleotides ratio in chromatogram. However, for polyploidy organisms, it is appropriated to determine a heterozygous individual by the identification of different copies for a given target gene or locus (van Belkum et al., 2001). The present result represents exactly such situation. The detection of less frequent alleles in non-cloned culture can be performed either through molecular cloning or, very cautiously, in chromatogram (Fig. 2). For the last option, however, the approach should be first validated since the smaller peak can be misinterpreted as background. Because of this, in the present study we only considered the sequences from molecular cloning to determine different copies of the targeted allele.

Aside the ambiguous sites targeted as the main objective of study, polymorphisms were also noticed along the alignment of all sequences obtained for this strain. In site 45, for instance, three BCDS and one BC_MC differ from their correspondent MCs and BC (Table 2). This suggests that biological clones represent a heterozygous infrapopulation, and the alternative allele (T) is among the respective BC_MC, but not in equal proportion to generate overlapped peaks in BCDS or NCDS. The polymorphisms can be explained either by: (i) point mutations, which have generated infrapopulation during the culture or (ii) initial aneuploidy, with different heterozygote cells: the sequences with the polymorphism in position 45 would represent the least frequent alleles and therefore could be detected only after biological and molecular cloning. Although mutations might occur frequently, option (ii) is more plausible because it is corroborated by the detection of MC presenting alleles different from their corresponding BCDS. Homozygous cells might also be present (after MC many presented just one allele). Nonetheless these findings demonstrate how less frequent alleles might often be undetected or excluded from DNA sequence based analysis. It also points that the removal of samples presenting multiple peaks would not avoid this bias (Camara et al., 2010).

3.3. Detection of polyclonal heterozygous sample and homologous recombination

Previous NCDS of IOC/L 2538 presented four double peaks and identical sequence types were observed, after biological cloning, among three BCDS (Table 2). However, the pattern was not kept for all other sequences: the NCDS differed in site 64 from the previous obtained sequence; the BCDS were either (i) identical to the previous sequence; (ii) identical to the NCDS or (iii) a third variation, with no double peak in the site 466 (Table 2). These results suggest at least three possible heterozygous cell populations occur within the IOC/L 2538 strain. However, when each site is observed, only positions 391 and 460 appear as classical heterozygous, with the possible nucleotides detected among the MCs. Such pattern would be expected for all sites since they are located in the same locus. For sequence types from case (ii), the corresponding MCs presented both nucleotide possibilities in the four sites, including position 64, which had no double peak. Therefore, it might represent a heterozygous infrapopulation in the sample, containing alleles with different nucleotides only in position 64. This confirms
a third heterozygous allele within the culture observed only after BC and MC.

Upon molecular cloning in a diploid organism, the four ambiguous sites could generate at most 16 possible haplotypes. Here, a total of six different haplotypes were detected among 42 MCs, obtained from nine BCs from the same strain IOC/L 2538. This number of alleles demonstrates the intra-strain sequence type diversity. Although PCR recombination and point mutation cannot be excluded, the origin of the supernumerary heterozygous alleles could also be consequence of homologous recombination. In most organisms, the primary function of homologous recombination is to allow genome protection by the faithful repair of DNA double-strand breaks and search for sequence homology. However, Trypanosoma brucei and Leishmania use homologous recombination as a beneficial mechanism for antigenic variation (Boothroyd et al., 2009) or for drug resistance (Ubeda et al., 2008), respectively.

These results reflect the aneuploidy state and homologous recombination. As an aneuploidy organism, a gene previously described as one-copy can in fact present a variable number of copies. Moreover, even between BCs, ACAs do occur during parasite growth (mitosis). Thus, if a BC is a heterozygous cell initially, this state may not be sustained during the necessary period of culture to obtain DNA. Nevertheless, as pointed out by Sterkers et al. (2012), the alleles will still be present in the same initial proportion (the process is stable instead of fluctuating) although the number of heterozygous cells may decrease. This means if the heterozygote was not detected at first, it will be after MC (see point 3.2).

3.4. Monoclonal heterozygous strain profiling in DNA sequences cannot exclude karyotype diversity

The strain IOC/L 2490 preserved the two double peaks in NCDS and BCDS (Table 2). The NCDS and all four BCDS identical to the previous sequence suggest, therefore, a monoclonal heterozygous strain. However, the clonal concept in Leishmania is only suitable from a biological point of view, but not genetically (Dujardin et al., 2007) since karyotype diversity is generated from one parasite cell (Sterkers et al., 2012, 2011). Indeed, the clonal theory for Leishmania does not refer to any precise cytological mechanism, but instead to the genetic consequences of clonality. In the present study we cannot guarantee the cells present the same karyotype, but it is possible to guarantee the equivalent proportion of alleles through the perfect 1:1 overlapping peaks observed (Fig. 2). Considering the two polymorphic sites (Table 2) and recombination occurrence, four possible haplotypes at most could be detected for a diploid cell. Three alleles were detected within 17 MCs, suggesting more than two homologous chromosomes are present.

3.5. The double peaks and polymorphisms presented coincident sites between the strains analyzed

Strain IOC/L 918 NCDS and previously NCDS were identical, with the three double peaks still present (Table 2). For the BCDS, two sites (391, 466) kept the ambiguity. Two other sites (42, 64), not double peaks initially, were included in the analysis because, curiously, among the respective MCs both nucleotide alternatives were observed. Moreover, these sites were also double peaks for other strains (IOC/L 2494 and 2538).

Five haplotypes (Table 2) where detected among the eight BCDS. This may represents the occurrence of populations with different heterozygous karyotypes. Additionally, it suggests the polymorphic sites, which were double peaks for other strains but not for IOC/L 918, might be related to hot spots of mutation/recombination.

3.6. Recombination detection

After RDP analysis including 175 L. (Viannia) non-cloned sequences plus all 286 sequences obtained in the present study, one recombination event was detected for one Leishmania braziliensis strain (IOC/L 3089, from Bahia, Northeastern Brazil). The major and minor parental populations indicated were, respectively, IOC/L 3072 (L. braziliensis, from Rio de Janeiro), and the BCDS and NCDS (which were identical) of IOC/L 2490 Leishmania naiffi/Leishmania lainsoni – (hybrid profile in 6PGDH isoenzyme) by MaxChi and 3Seq algorithms (P < 0.05). The breakpoint beginning/end was wide enough to encompass almost the whole sequence, but the MaxChi graph representation indicates a window between positions 350 and 492, which includes the main polymorphic and double peaks sites evaluated in the present study.

Recombination is usually hard to detect within species, mainly because of low inter-strain diversity, or apparent low diversity due to inappropriate sampling (Prugnolle and De Meeus, 2010). However, some authors were able to present such data: population genetics studies have described inbreeding in L. braziliensis as well as in other Leishmania species (Rougeron et al., 2010; Ferreira et al., 2012). Recombination signals appear also in MLSA approaches (Boité et al., 2012) and in natural hybrids that were widely described (Akopyants et al., 2009; Belli et al., 1994; Brito et al., 2009) or even created in vitro (Coelho et al., 2012). RDP software recognizes IUPAC symbols, enhancing the ability of the algorithms to detect recombination. The possibility of PCR recombination cannot be excluded, but random PCR repetition of some sequences (data not shown) validated the polymorphisms detected. The present results indicate that recombination may occur between specific clones, and the molecular and biological cloning can contribute to the detection of a specific recombinant allele. Well-structured clonal complexes in Leishmania (Boité et al., 2012), could be the consequence of homologous recombination, as described in bacteria (Gonzalez-Gonzalez et al., 2013; Paul et al., 2013; Jinkerson et al., 2013) and other microorganisms (Vink et al., 2011).

4. Further comments

The main concern that has driven this study is the discussion regarding intra strain variation detected here in DNA sequence based analysis in Leishmania and elsewhere in Trypanosoma cruzi (Camara et al., 2010). The exact determination of allele frequency in individuals and, eventually, in populations is essential in association studies (Tibayrenc and Ayala, 2012, 2013); therefore, the quantitative accuracy of genotyping is critical. Multiple peaks in chromatograms, for instance, might point to: (i) different clones in the sample; (ii) heterozygotes or both. After the present results we observe that to exclude strains which present multiple peaks from the analysis is not the best way to deal with these facts since such diversity might be present even when the multiple peaks are not detected.

The occurrence of polyclonal samples is a constant if one considers the aneuploidy state described for Leishmania. The ACAs during mitosis seem to be quite frequent, generating daughter cells with different karyotype profiles, even after cellular cloning (infra-population) (Sterkers et al., 2012). Therefore, the (genetic) clonal concept is quite intricate to be applied in this parasite (Tibayrenc and Ayala, 2012). We demonstrate here that there is DNA sequence diversity within one strain kept in culture and/or cryopreserved; and changes can be detected along time, in different DNA batches. The allele frequency, however, usually does not change within the mosaic population. If multiple peaks are detected once and the two most prominent considered, that means the most frequent alleles (prevalent infrapopulation in the sample) are being distinguished.
Nevertheless, as presented in this project, some alleles will only be observed after BC and MC.

In terms of allele frequency, DNA sequence based approaches are able to reflect (partially) the biological reality, especially if there is a good sample representation in terms of diversity and geographic origin. A strain will still appear as a ‘heterozygote’ when global analysis methods are used because all the alleles present at first (in a real heterozygous cell) are still present in the population. This process seems to be stable, so, despite the mosaic of cells, a pattern can still be determined as representative for that population. From a population genetics point of view, if a heterozygous signal is detected, it does represent heterozygous strains at some point and reflects the two most frequent alleles in that sample. Nevertheless, the real diversity will always be underestimated (Prugnolle and De Meeus, 2010) because some alleles will remain undetected. The bias of underestimated diversity could be diminished through a proper representation of the circulating strains, such that the greatest number of different alleles would be detected. A deeper knowledge over the karyotypes profiles of the different strains is of major importance.

Acknowledgments

We thank the Plataforma de Seqüenciamento do Instituto Oswaldo Cruz (PDTS) for performing all the sequencing; Instituto Oswaldo Cruz – FIOCRUZ, CNPq/Fiocruz (PIBIC) and Faperj for financial support. We also would like to thanks Global Science Editing, UK, Mariel Marlow and Hugo Caire Castro de Faria Neto for English revision.

References


