First report of Leishmania RNA virus 1 in Leishmania (Viannia) braziliensis clinical isolates from Rio de Janeiro State - Brazil

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BACKGROUND Leishmania parasites carry a double-stranded RNA virus (Leishmania RNA virus - LRV) that has been divided in LRV1 and LRV2.

OBJECTIVES Leishmania (Viannia) braziliensis clinical isolates were assessed in order to determine LRV presence.

METHODS Two-round polymerase chain reaction (PCR and nested PCR) was performed to detect LRV1 or LRV2 in L. (V) braziliensis clinical isolates (n = 12).

FINDINGS LRV1 was detected in three clinical isolates which was phylogenetically related to other sequences reported from other American tegumentary leishmaniasis (ATL) endemic areas of Brazil. Patients infected with L. (V) braziliensis LRV-negative showed only cutaneous lesions while LRV-positive reported different manifestations.

MAIN CONCLUSION Data presented here show for the first time that LRV1 is circulating in L. (V) braziliensis clinical isolates from Rio de Janeiro State in Brazil.

Key words: Leishmania RNA Virus 1 - Rio de Janeiro - Leishmania (Viannia) braziliensis - clinical isolates

American tegumentary leishmaniasis (ATL) is the term used to describe cutaneous lesions caused by Leishmania parasites of Viannia and Leishmania subgenus, exclusively found in the American continent. The clinical spectra of ATL is broad, including cutaneous leishmaniasis (CL), severe diffuse CL, disseminated CL (DCL), metastatic and mucosal leishmaniasis (ML), and mucocutaneous leishmaniasis (MCL). Eleven species have been reported as ATL causative agents and, in Brazil, ten of them are related to ATL: Leishmania (Viannia) braziliensis, L. (V) guyanensis, L. (V) panamensis, L. (V) lainsoni, L. (V) naffii, L. (V) shawi, L. (V) utingensis, L. (V) lindenbergi, L. (Leishmania) amazonensis and L. (L) mexicana. The diverse clinical manifestations of ATL vary according to Leishmania spp., immune state of the mammalian host and parasite virulence factors. Leishmania parasites carry a double-stranded RNA virus (LRV) that has been divided, according to genetic distances between LRV types found on infected Leishmania strains, in LRV1 and LRV2. To date, LRV1 has been detected in clinical isolates from Bolivia, Colombia, Costa Rica, Ecuador, French Guiana and Peru, while LRV2 only in isolates from Middle eastern and African countries, showing LRV geographical distribution. Specifically, in Brazil, LRV1 has been detected in clinical isolates of L. (V) braziliensis, L. (V) panamensis, L. (V) guyanensis, L. (V) lainsoni, L. (V) naffii and L. (L) amazonensis from Minas Gerais, Rondônia and Amazonas states. Interestingly, RNA virus in some Leishmania spp. is considered as a virulence factor associated with the development of severe forms of ATL. For instance, L. (V) guyanensis metastatic strains had a higher rate of LRV1 positivity than non-metastatic strains and, during macrophages infection, caused over-expression of proinflammatory mediators such as TNF-α and IL-6. Furthermore, these authors also showed that macrophages treated with purified LRV1 had a similar phenotype compared to the ones infected with the metastatic strains, expressing not only higher levels of TNF-α and IL-6 but also α-chemokine and β-chemokines, which compose a typical immune profile of patients developing MCL. Similarly, using L. (V) guyanensis, it was shown that LRV1 can be transmitted through exosomes that are secreted to the extracellular environment from multivesicular bodies and/or the parasite flagellar pocket. This is interesting...
since it was demonstrated, on in vivo models, that co-inoculation of L. (L.) mexicana and L. (V.) panamensis with their respective exosomes increased lesion size but co-inoculation with L. (V.) guyanensis LRV positive exosomes exacerbated lesion development.\(^{(28)}\) Another study revealed that LRV1 positivity frequency in L. (V.) guyanensis and L. (V.) braziliensis LRV positive exosomes from patients with MCL was higher than in patients with CL.\(^{(23)}\) However, the data on the subject is contradictory, as another study showed a similar LRV1 detection rate among L. (V.) braziliensis, L. (V.) guyanensis and L. (V.) peruviana metastatic and non-metastatic strains in a longitudinal cohort of ATL patients from Peru.\(^{(29)}\) Also, in a cohort of ATL patients from the southeast, north and northeast regions of Brazil, less than 5% of strains were LRV1 positive and the severity of the disease was related to other factors such as age, gender and immune status of the hosts.\(^{(22)}\) In another study with 40 L. (V.) braziliensis isolates from Minas Gerais State, no sample was positive for LRV1. In this context, we report for the first time that L. (V.) braziliensis clinical isolates from ATL patients living in Rio de Janeiro State can be infected with LRV1. In fact, the results presented here contribute to reinforce the heterogeneity previously seen for these clinical isolates.\(^{(31,32)}\)

**MATERIALS AND METHODS**

In this study, RNA was obtained from stationary-phase promastigotes \((10^7\) to \(10^8\) parasites/mL) of L. (V.) braziliensis clinical isolates \((n = 12)\) and positive control sample [IOC/L0565 (MHOM/BR/1975/M4147) L. (V.) guyanensis], cultured in vitro as previously described.\(^{(31)}\) Each sample was lysed in TRIzol containing chloroform, and RNA was extracted using RNeasy Mini Kit (QIAGEN, Germany). Then, RNA samples were converted into cDNA, using High-Capacity cDNA Reverse Transcription kit (Applied Biosystems, USA), to detect LRV1 and LRV2 with specific primers in a two-round polymerase chain reaction (PCR) (Table I). Both PCR and nested PCR were conducted in a final volume of 25 \(\mu\)L of reaction containing 1X PCR buffer, 3 mM of \(\text{MgCl}_2\), 2.5 U of Taq DNA Polymerase (Invitrogen Life Technologies, Brazil), 200 mM of triphosphate deoxyribonucleotides dNTP (Invitrogen Life Technologies, Brazil) and 0.2 \(\mu\)M of each primer, and the PCR assay conditions were performed as described in Table I. The amplicons obtained from nested PCR were purified using the NucleoSpin\textsuperscript{TM} Gel and a PCR Clean-up kit (Macherey-Nagel GmbH & Co. KG, Germany), with a minor change in incubation (increased to five minutes). The purified products were subjected to sequencing in both directions using the ABI Prism\textsuperscript{TM} BigDye Terminator Cycle Sequencing kit (Applied Biosystems, USA) on an ABI 3730 automatic DNA sequencer at Fiocruz facilities [Capillary Electrophoresis DNA Sequencing Platform (SANGER) - RPT01A].\(^{(33)}\) The obtained sequences were deposited in GenBank under accession number ON409677-ON409679. Electropherograms were analysed using Chromas 2.4, while percent identity with sequences producing significant alignments was performed using the Basic Local Alignment Search Tool using nucleotide (BLASTn). Nucleotide sequences were aligned by the CLUSTAL W algorithm from Mo-
TABLE II
Clinical characteristics of patients infected with *Leishmania (Viannia) braziliensis*

| Parasite isolate | Age/Sex | Occupation | Local of infection* | Clinical manifestation | Clinical response** | Subsequent treatments*** | LRV1 | LRV2 |
|------------------|---------|------------|---------------------|-----------------------|--------------------|-------------------------|------|------|
| 1                | 41/M    | Farmer     | Campo Grande/RJ/RJ  | CL                    | NR                 | Meglumine antimoniate (1 round) | -    | -    |
| 2                | 59/M    | Garbage recycler | Campo Grande/RJ/RJ | CL                    | NR                 | Meglumine antimoniate (3 rounds) | -    | -    |
| 3                | 31/M    | Military   | -/Manaus/AZ         | CL                    | NR                 | Meglumine antimoniate (1 round) | -    | -    |
| 4                | 24/M    | Self employed | Mazomba/Itaguaí/RJ | CL                    | R                  | NST                     | -    | -    |
| 5                | 55/F    | Nurse      | Santíssimo/RJ/RJ    | CL                    | R                  | NST                     | -    | -    |
| 6                | 48/M    | Housekeeper | Cidade Jardim Marajoara/Japeri/RJ | CL | R | NST | - | - |
| 7                | 48/M    | Self-employed | -/Barra Mansa/RJ    | DCL 24 lesions        | NR                 | Meglumine antimoniate (1 round) | +   | -    |
| 8                | 25/F    | Housewife  | Caçador/Itaguai/RJ  | CL                    | NR                 | Meglumine antimoniate (1 round) | -    | -    |
| 9                | 38/M    | Civil construction | Campo Grande/RJ/RJ | CL                    | R                  | Meglumine antimoniate (1 round) | -    | -    |
| 10               | 52/F    | Housekeeper | Itaipava/Petrópolis/RJ | CL | NR | Meglumine antimoniate (1 round) | -    | -    |
| 11               | 35/M    | Civil construction | -/Carangola/MG    | ML nose               | R                  | Meglumine antimoniate (4 rounds) | +   | -    |
| 12               | 21/F    | Student    | Campo Grande/RJ/RJ  | CL                    | NR                 | Meglumine antimoniate (1 round) | +   | -    |

*As labelled in; *neighborhood/city/state; **response after 1 round of intramuscular Meglumine antimoniate 5 mg/kg/day for 30 days until clinical cure (epithelisation); ***all patients were first treated with intramuscular Meglumine antimoniate 5 mg/kg/day for 30 days until reaching clinical cure. Who did not respond was submitted to subsequent treatments until reaching clinical cure, Meglumine antimoniate (same dose) and/or Amphotericin B (lipid complex: total dose of 1800 mg; deoxycholate: total dose of 1000 mg). AZ: Amazonas; CL: cutaneous leishmaniasis; DCL: disseminated cutaneous leishmaniasis; F: feminine; LRV1: *Leishmania* RNA virus 1; LRV2: *Leishmania* RNA virus 2; M: masculine; MG: Minas Gerais; ML: mucosal leishmaniasis; NR: non-responder; NST: no subsequent treatments; R: responder; RJ: Rio de Janeiro.
The presence of LRV1 was identified in three L. (V.) braziliensis isolates from patients living in Rio de Janeiro by gel electrophoresis (Fig. 1) and confirmed by sequencing. Phylogenetic analysis indicates that the LRV1 have greater identity compared to other LRV1 identified in L. (V.) braziliensis isolated from patients of other Brazilian cities, such as Porto Velho and Candeias of Rondonia State (Fig. 2).

**RESULTS**

In accordance with LRV reported by other authors in Latin America, LRV2 was not identified in this study. Although all the individuals were residents of Rio de Janeiro, one of the patients was infected by L. (V) braziliensis presumably in the municipality of Carangola in Minas Gerais State (Table II), where the presence of LRV has been previously reported in patients with CL.\(^{36}\) The infection by L. (V) braziliensis of the other two patients in the study occurred in the state of Rio de Janeiro (municipalities of Barra Mansa and Rio de Janeiro) (Table II), where circulation of LRV has not been reported so far. In addition, to confirming LRV1, it was possible to observe the following single-nucleotide polymorphisms among the isolates: T/A (position 35, isolates 7 and 11 versus 12), T/G (position 37, isolates 7 and 11 versus 12), T/C (position 65 isolates 7 and 12 versus 11), G/A (position 71, isolates 7 and 12 versus 11). Moreover, it is important to mention that these three clinical cases have a more exuberant disease profile, and at least two needed subsequent treatments, while all the negative cases are associated to patients with CL (Table II).

In conclusion, the findings presented here are original and serve as an alert that LRV1 is circulating in L. (V) braziliensis in Rio de Janeiro. Additional studies with more clinical isolates are needed to assess a possible correlation between LRV1-infected parasites, clinical manifestations and treatment response, as described for ATL in other endemic areas.

**DISCUSSION**

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AUTHORS' CONTRIBUTION
AZ-P, MF and GD-L conceived, designed the analyses and collected the data; JLS, LM, LR, MIP and FCS supplied clinical isolates information; AZ-P, MF, GD-L and CRA performed the manuscript conceptualisation and final review.

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