Molecular characterization of the MRPA transporter and antimony uptake in four New World Leishmania spp. susceptible and resistant to antimony

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ABSTRACT

ATP-binding cassette (ABC) transporters have been associated with drug resistance in various diseases. The MRPA gene, a transporter of ABC subfamily, is involved in the resistance by sequestering metal-thiol conjugates in intracellular vesicles of Leishmania parasite. In this study, we performed the molecular characterization of the MRPA transporter, analysis of P-glycoprotein (Pgp) and aquaglyceroporin-1 (AQP1) expression, and determination of antimony level in antimony-susceptible and -resistant lines of L. (V.) guyanensis, L. (L.) amazonensis, L. (V.) braziliensis and L. (L.) infantum. PFGE analysis revealed an association of chromosomal amplification of MRPA gene with the drug resistance phenotype in all SbIII-resistant Leishmania lines analyzed. Levels of mRNA from MRPA gene determined by real-time quantitative RT-PCR showed an increased expression of two fold in SbIII-resistant lines of Leishmania guyanensis, Leishmania amazonensis and Leishmania braziliensis. Western blot analysis revealed that Pgp is increased in the SbIII-resistant L. guyanensis and L. amazonensis lines. The intracellular level of antimony quantified by graphite furnace atomic absorption spectrometry showed a reduction in the accumulation of this element in SbIII-resistant L. guyanensis, L. amazonensis and L. braziliensis lines when compared to their susceptible counterparts. Interestingly, a down-regulation of AQP1 protein was observed in the SbIII-resistant L. guyanensis and L. amazonensis lines, contributing for decreasing of SbIII entry in these lines. In addition, efflux experiments revealed that the rates of SbIII efflux are higher in the SbIII-resistant lines of L. guyanensis and L. braziliensis, that may explain also the low SbIII concentration within of these parasites. The BSO, an inhibitor of γ-glutamylcysteine synthetase enzyme, reversed the SbIII-resistance phenotype of L. braziliensis and caused an increasing in the Sb intracellular level in the LbSbR line. Our data indicate that the mechanisms of antimony-resistance are different among species of Leishmania analyzed in this study. © 2013 The Authors. Published by Elsevier Ltd. All rights reserved.

1. Introduction

Leishmaniasis is a complex of diseases caused by different species of protozoan parasites belonging to the genus Leishmania. This neglected tropical disease comprises clinical manifestations that range from self-healing cutaneous (CL) and mucocutaneous (MCL) skin lesions to a visceral (VL) form, which is lethal if untreated (Ashutosh et al., 2007). In the New World, L. (Leishmania) infantum (syn. L. (L.) chagasi) (Kuhl et al., 2007) is the causative agent of VL, whereas L. (L.) amazonensis and L. (Viannia) guyanensis cause CL, and L. (V.) braziliensis causes CL and MCL (Marzochi and Marzochi, 1994; Murray et al., 2005; Shaw, 2006). The disease is endemic in 98 countries, especially in northern Africa, Asia, the Mediterranean and Latin America, with more than 350 million people at risk (Ashford et al., 1992; Alvar et al., 2012). There are an estimated 12 million people infected worldwide (World Health Organization, 2012). The estimated incidence of leishmaniasis is approximately 0.2–0.4 million VL cases and 0.7–1.2 million CL cases per year. More than 90% of global VL cases occur in six countries: India, Bangladesh, Sudan, South Sudan, Ethiopia and...
Brazil. Whereas Afghanistan, Algeria, Colombia, Brazil, Iran, Syria, Ethiopia, North Sudan, Costa Rica and Peru account for 70–75% of global estimated CL incidence (Alvar et al., 2012).

The control of leishmaniasis relies mainly on chemotherapy. The first line choice of treatment against all forms of the disease is based on the administration of pentavalent antimony-containing compounds, such as sodium stibogluconate (SSG) (Pentostam®) and N-methyl-glucamine (Glucantime®). The mechanisms of action of antimony are not fully elucidated. It is known that SbV acts on the amastigote form, inhibiting enzyme activity and the oxidative pathway of fatty acids (Herwaldt, 1999). It is generally agreed that trivalent antimony (SbIII) is the active form of the drug against Leishmania amastigote and promastigote forms (Frézard et al., 2009). Murray and Nathan (1988) demonstrated that macrophage activation has a significant effect on intracellular parasite killing. It has been reported that SbV induces the macrophage to produce leishmanicidal molecules like nitric oxide (NO) and reactive oxygen species (ROS) by activation of signalling pathways, leading to the elimination of intracellular Leishmania donovani amastigotes (Basu et al., 2006). In an animal infection model, the mode of action of SbV is dependent on a number of factors including T cell subsets and cytokines (reviewed by Murray, 2001). Furthermore, SbV was found to be a potent inhibitor of protein tyrosine phosphatases, leading to an increase in cytokine responses (Pathak and Yi, 2001). Thus, these results suggest that SSG may kill the parasites by both direct and indirect mechanisms. Studies suggest that SbIII causes disturbances in the thiol redox potential of the parasite by inducing the efflux of intracellular thiols and by inhibiting trypaonthione reductase, leading the cell to death by oxidative stress (Wylie et al., 2004; Moreira et al., 2011). It has been shown that antimony kills the parasite by a process involving DNA fragmentation and externalization of phosphatidylserine on the outer surface of membrane (Sereno et al., 2001; Lee et al., 2002; Sudhandiran and Shaha, 2003).

Drug resistance is one of the major clinical problems for the treatment of various diseases ranging from bacteria and parasite infections to cancer. Treatment failure with pentavalent antimonials has been reported recently in several countries. In India, over 60% of patients with VL do not respond to treatment with pentavalent antimonials, due to acquired resistance (Sundar et al., 2000). The mechanisms by which species of Leishmania acquire resistance to antimonials have been subject of intensive research for several decades. It has been described that resistance is an interplay between uptake, efflux and sequestration of active molecules (reviewed by Croft et al., 2006). A down-regulation of aquaglyceroporin-1 (AQPI) is correlated with lower SbIII uptake, decreasing the drug concentration within the cell (Marquis et al., 2005). On the other hand, amplification of DNA segments has been observed in several Leishmania species selected for drug resistance (Beverley, 1991; Ouellette and Borst, 1991).

Different mechanisms of drug resistance have been identified in Old World Leishmania species (reviewed by Croft et al., 2006). Although there is considerable evidence showing variability in the response to antimony chemotherapy in New World pathogenic Leishmania species (Moreira et al., 1998; Romero et al., 2001), the mechanisms of drug resistance in these species have not been extensively analyzed. Therefore, understanding the biological diversity and responses to chemotherapy of different New World Leishmania species is necessary to ultimately overcome current limitations in anti-parasitic drug treatment. Thus, in this study we performed the molecular characterization of the MRPA transporter and determination of antimony level in antimony-susceptible and -resistant lines, which were experimentally induced, in four New World Leishmania species: Leishmania guyanensis, Leishmania amazonensis, Leishmania braziliensis and Leishmania infantum. Promastigote forms were characterized for: chromosomal location, analysis of amplification and mRNA levels of the MRPA gene; Pgp and APQ1 protein expression; measurement of intracellular antimony level, efflux rates of SbIII and susceptibility of L. braziliensis lines to BSO, an inhibitor of γ-glutamylcysteine synthetase (GCS) enzyme.

2. Materials and methods

2.1. Leishmania strains

In this study, we used promastigote forms of four different New World Leishmania species: L. guyanensis, L. amazonensis, L. braziliensis and L. infantum (Table 1). These lines were selected in vitro to trivalent antimony (SbIII) by step-wise drug pressure and the resistance index varied from 4 to 20-fold higher than of their wild-type counterparts (Liarte and Murta, 2010). This previous study showed that the SbIII-resistant lines of L. amazonensis, L. braziliensis and L. infantum have cross-resistance to paromomycin. Parasites were grown at 26 °C in M199 medium (Liarte and Murta, 2010).

2.2. Pulsed field gel electrophoresis (PFGE)

Chromosomal DNA from Leishmania lines (2.0 × 10⁹ cells/mL) was prepared in low-melting temperature agarose plugs as described by Smith et al. (1988). The agarose blocks containing intact Leishmania chromosomes were separated by PFGE in 1% agarose gels in 0.5x TBE (45 mM Tris-HCl, 45 mM boric acid, 1 mM EDTA, pH 8.3) using the Gene Navigator TM System (Amersham Pharmacia, Buckinghamshire, UK). The running buffer (0.5x TBE) was maintained at 9 °C throughout the electrophoresis. Saccharomyces cerevisiae chromosomes were used as size markers (Bio-Rad Inc., Hercules, CA, USA). Electrophoresis conditions were standardized to allow the separation of the greatest number of parasite chromosomes in a single gel, as follows: 90 s for 18 h, 200 s for 18 h, 400 s for 22 h and 600 s for 7 h at 90 V. A range from 365 to 2200 kb was used for wide separation of Leishmania chromosomes. After electrophoresis, gels were stained with ethidium bromide (10 μg/mL) and bands were then transferred onto nylon membrane (Sambrook et al., 1989). The MRPA gene was identified by incubation of the membrane with the 32P-labelled MRPA gene probe. For this probe, it was used a 452 bp MRPA fragment (LbrM23_V2.0280) amplified with the primers: forward 5’-TCGTAGATATTCGGGTGCCTTT-3’ and reverse 5’-ACGCTTCACGCTTTCAATTTT-3’.

2.3. Southern blot

Genomic DNA was isolated from the antimony-susceptible and -resistant Leishmania species according to the protocol described by Sambrook et al. (1989). Approximately 10 μg total DNA of
Leishmania species used in this study were digested with the restriction enzyme BamHI (Invitrogen, Carlsbad, CA, USA). The fragments were separated by electrophoresis on a 1% agarose gel and transferred to nylon membrane (Sambrook et al., 1989). The blots were hybridized with [α-32P]dCTP labeled MRPA gene probe as described above. Leishmania DNA polymerase gene (LbrM.16.1600) was used as quantitative control. A fragment of 483 bp from this gene, after amplification using primers: forward 5'-GAAGACAGAGAAGGATGCGA-3' and reverse 5'-GAGAGCGGCGCCACACCG-3', was used as probe. The band intensities were analyzed using the software CP ATLAS 2.0 (http://lazarsoftware.com/download.html).

2.4. Quantitative real-time RT-PCR

Reverse transcription reactions for first strand complementary DNA (cDNA) synthesis were carried out as described below. Each reaction contained 2 μg total RNA, 0.5 μg oligo d(T), 1x first strand buffer, 10 mM DTT, 0.5 mM dNTP, 40 U RNasin and 200 U Superscript II reverse transcriptase in a final volume of 20 μL. All the reagents used were obtained from Invitrogen (Life Technologies, Carlsbad, CA, USA). All reactions were allowed to proceed for 1 h at 42 °C before being stopped by incubation at 70 °C for 20 min. The obtained cDNA was then diluted 15x in water and 5 μL of the reaction product was amplified by real-time PCR using specific primers. Amplification reaction was carried out using the 7000 System SDS Software (PE Applied Biosystems, Foster City, CA, USA). The primers, MRPA forward: 5'-AAGTGGGCAGCGACTCAAA-3' and MRPA reverse: 5'-CCAGTTCCAGCTTCCGT-3', were the same described by Torres et al. (2010) and Adaui et al. (2011). The reaction contained 2 μL of cDNA template, 1x SYBR Green reaction mix (Applied Biosystems, Life Technologies, Carlsbad, CA, USA) and 5 μL of template DNA. The PCR conditions were as follows: an initial denaturation step at 95 °C for 10 min followed by 40 cycles of denaturation at 95 °C for 15 s and annealing/extension at 60 °C for 1 min. Standard curves were prepared for each run using known quantities of TOPO PCR 2.1 plasmids (Invitrogen, Life Technologies, Carlsbad, CA, USA) containing the MRPA and DNA polymerase genes. Estimates of DNA levels were obtained using the Sequence Detection System data analysis software. Values were normalized to those obtained for DNA polymerase for each sample.

2.5. Western blot analysis

Total proteins and membrane protein fractions from different Leishmania lines were extracted according to the protocols described by Gamarro et al. (1994) and Grogl et al. (1991), respectively. Protein extracts were separated by electrophoresis on a 12% SDS polyacrylamide gel and electrotransferred onto nitrocellulose membranes (Bio-Rad, Hercules, CA, USA). The blots were blocked by incubation with 5% instant non-fat dry milk in PBS supplemented with 0.05% Tween 20 (PBS-T) for 1 h. The blots were then washed twice in PBS-T and incubated for 16 h at 4 °C in the presence of antibodies: monoclonal antibody C219 (1:100) (Abcam, Cambridge, UK) or rabbit polyclonal antibody anti-AQP1 (1:5) (kindly provided by Andrade et al. – in preparation). The blots were washed and incubated with anti-IgG mouse HRP-conjugate (1:2,000) (GE Healthcare) or HRP-conjugated anti-rabbit IgG (1:5,000) (GE Healthcare) for 1 h at room temperature. After, the blots were exposed to Amersham ECL Plus detection agent according to the manufacturer’s instructions and exposed to an X-ray film (Amersham, Buckinghamshire, UK). The results were normalized using the anti-α-tubulin monoclonal antibody (Sigma, St. Louis, USA). The intensity of the bands was analyzed using the software CP ATLAS 2.0.

2.6. Antimony transport assays

2.6.1. Uptake assay

Before performing the uptake and efflux assays, the SbIII-resistant lines were maintained at least two passages in M199 medium in the absence of SbIII to eliminate the residual drug. The antimony uptake assay was based on two protocols previously described (Roberts and Rainey, 1993; Wang et al., 2003). Log phase Leishmania promastigotes were washed twice in a HEPES/Glucose (HG) buffer (20 mM HEPES, 0.15 M NaCl, 10 mM glucose, pH 7.2) and resuspended in this buffer at a density of 1.0 × 10^6 cells/mL. The volume of 1 mL of this parasite suspension was aliquoted into tubes in quadruplicate: tubes containing only parasites (blank) and tubes with parasites in the presence of 540 μM SbIII (potassium antimonyl tartrate) (Sigma, Saint Louis, MO, USA). Cells were then incubated at 26 °C for 1 h under agitation. Subsequently, the cells were centrifuged at 1816 g for 5 min at 4 °C and washed three times with HG buffer. Pellets were then resuspended with 100 μL HG buffer. One aliquot of 10 μL of each tube was used for normalization (parasite quantification) and the remaining volume (90 μL) was submitted to digestion with nitric acid (65%). Antimony concentration was quantified by graphite furnace atomic absorption spectrometry (Perkin Elmer AAnalyst 600). Each uptake assay was performed three times and the signal from blanks was used for background subtraction.

In addition, the antimony-susceptible and -resistant lines of the species L. guyanensis and L. braziliensis were evaluated for the kinetics of incorporation of Sb. The parasites were incubated with 540 μM SbIII and, at different time intervals (0.5, 1, 2 and 3 h), 1 mL aliquots of the parasite suspension were taken and they were submitted to the protocol described above. This assay was also performed to identify the time points at which the WTS and SbR lines exhibit the same intracellular concentration of Sb, to be explored in the efflux protocol.
2.6.2. Efflux assay

Log phase antimony-susceptible and -resistant *L. guyanensis* and *L. braziliensis* cells were washed twice with HG buffer and resuspended in this buffer at a density of 1.0 × 10⁶ cells/mL. Aliquots of 1 mL containing only parasites (blank) were taken and the remaining cells were incubated with 540 µM SbIII for each 1 mL (potassium antimony tartrate) (Sigma, Saint Louis, MO, USA) at 26 °C under agitation. The WTS lines of *L. guyanensis* and *L. braziliensis* were incubated during 2 min and 30 min, respectively. Both SbR lines of these species were incubated for 70 min. After incubation, the cells were centrifuged at 890g for 5 min at 4 °C and washed two times with HG buffer. The pellets were then resuspended with HG buffer at the original density and the cells were incubated at 26 °C under agitation. Aliquots of 1 mL were taken from the parasite suspension at the times 0, 1 and 2 h. Subsequently, these aliquots and the blanks were centrifuged at 890g for 5 min at 4 °C and washed three times with HG buffer. After the last centrifugation, the pellets were resuspended with 100 µL HG buffer. One aliquot of 10 µL of each tube was used for normalization (parasite quantification) and the remaining volume (90 µL) was submitted to digestion with nitric acid (85%). Antimony concentration was quantified by graphite furnace atomic absorption spectrometry (Perkin Elmer AAnalyst 600). Each efflux assay was performed three times in triplicate and the SbIII dosage from blanks was used for background subtraction.

2.6.3. Test of susceptibility to BSO and SbIII uptake in *L. braziliensis* lines pre-treated with this inhibitor

Approximately 2.0 × 10⁶ cells/mL of *L. braziliensis* were grown in M199 medium containing various concentrations (2.5–30 mg/mL) of buthionine-sulphoximine (BSO) (Sigma) during 48 h at 26 °C. BSO is an inhibitor of γ-glutamlycysteine synthetase (γ-GCS), a rate-limiting enzyme in thiol biosynthesis (Griffith and Meister, 1979). The drug concentration that decreases cell growth by 50% (IC₅₀) was determined by counting the parasites in the presence and absence of the BSO. After pre-treatment with BSO, the uptake of Sb was investigated in both susceptible and resistant *L. braziliensis* lines. These parasites were first grown in M199 medium for 48 h in the presence BSO at their respective IC₅₀. Subsequently, the cells were washed twice with HG buffer, resuspended in this buffer at a density of 1.0 × 10⁶ cells/mL and submitted to the uptake assay.

2.7. Statistical analysis

Data were analyzed by Student’s t test performed using the software GraphPad Prism 5.0. A p value of <0.05 was considered statistically significant.

3. Results

3.1. MRPA amplification in SbR Leishmania parasites

The molecular karyotype of antimony-resistant and -susceptible *Leishmania* lines, obtained through PFGE is presented in Fig. 1A. Overall, the pattern of the physical chromosomal map obtained for Sb-resistant and -susceptible *Leishmania* parasites is similar except for LgsB, which, in comparison with its parental WT strain, showed a supplementary band of approximately 1900 kb (Fig. 1A white arrow). The pattern of hybridization of the chromosomes showed that the MRPA gene probe recognized a chromosomal band of approximately 800 kb in all lines of *Leishmania* analyzed (Fig. 1B) which fits with the size of chromosome 23 (795 kb). In a preliminary analysis, the intensity of this band was increased in the SbIII-resistant line of *L. amazonensis*. An interesting observation in the antimony-resistant line of *L. braziliensis* (LbS) is that the MRPA gene probe recognized in this sample a DNA smear and two other bands of approximately 200 kb and 1500 kb (Fig. 1B). This result indicates an extrachromosomal amplification of MRPA gene in LbsB. In agreement with this observation, another band of approximately 700 kb was observed in the resistant sample of *L. infantum* (Fig. 1B).

Southern blot assays were carried out using samples of genomic DNA from different *Leishmania* lines previously digested with the endonuclease BamHI, which has one restriction site within MRPA gene (LbrM23_V2.0280) in a conserved region. Hybridization of nylon membranes containing BamHI-digested DNA against a MRPA-specific probe revealed a major band of 11.0 kb and other faint band of 5.0 kb in all samples analyzed (Fig. 2A). Considering the 11 kb band, after normalization using a DNA polymerase gene probe (Fig. 2B), the densitometry revealed an increased intensity of three- and ten-fold in antimony-resistant *L. amazonensis* and *L. braziliensis* lines, respectively, when compared to their susceptible counterparts. This data confirms MRPA gene amplification in these antimony-resistant *Leishmania* lines. No difference was observed in the other *Leishmania* lines analyzed.

![Fig. 1. Chromosomal location of the MRPA gene in the antimony-susceptible (WTS) and -resistant (Sb-R) lines of Leishmania. (A) Chromosomal bands from the Leishmania species were separated by PFGE and stained with ethidium bromide. (B) Profile of the chromosomal bands hybridized with a ^32P-labelled MRPA-specific probe. Whole chromosomes from Saccharomyces cerevisiae were used as molecular weight markers. The white arrow shows an additional band of approximately 1900 kb in the LgS line.](image-url)
3.2. Increased MRPA transcripts in SbR Leishmania

MRPA mRNA levels in the Leishmania lines were evaluated by real-time RT-PCR. Measurements of total RNA were normalized by comparison with those obtained for the housekeeping gene DNA polymerase (LbrM.16.1600). Then, a standard curve was obtained using serial dilutions (10^3–10^6 molecules) of plasmids containing the DNA polymerase and MRPA genes cloned into pCR2.1 TOPO vector. We obtained a standard curve with good linearity for both genes (r^2 = 0.998 to DNA polymerase and r^2 = 0.973 to MRPA). Our data showed a slope of ~3.78 and ~3.60 for the DNA polymerase and MRPA genes, respectively, indicating high efficiency of PCR (results not shown). The specificity of PCR was analyzed by plots of the temperature-dependent dissociation of the SYBR GREEN dye from the MRPA and DNA polymerase PCR products.

The results revealed that fluorescence was only emitted at one temperature, suggesting that only one PCR product was generated in each reaction (results not shown).

The amount of cDNA amplified in the samples of Leishmania spp. was determined by linear regression analysis using the Ct values obtained from the standard curve generated with known amounts of the plasmids of the MRPA gene, normalized with DNA polymerase values. The results showed that mRNA level from MRPA gene in Leishmania lines is increased two-fold in the antimony-resistant L. guyanensis, L. amazonensis and L. braziliensis lines compared with their respective susceptible pairs. No difference in the MRPA gene expression between antimony-resistant and -susceptible lines of L. infantum was observed (Fig. 3).

3.3. Expression level of Pgp

We also determined the levels of Pgp in the antimony-susceptible and -resistant Leishmania lines by Western blot analysis using a monoclonal antibody (C219). According to manufacturer’s instructions (Abcam), this antibody recognizes an internal, highly conserved amino acid sequence: VQEALD and VQAALD, corresponding to the C-terminal and N-terminal regions, respectively, found in both MDR1 and MDR3 isoforms of P-glycoprotein of mammals. The results revealed that the C219 antibody recognized a polypeptide of 170 kDa in some Leishmania samples analyzed (Fig. 4B). According to literature data, this polypeptide corresponds to the expected size of Pgp (Cornwell et al., 1987). Interestingly, this antibody detected Pgp only in antimony-resistant L. guyanensis and L. amazonensis lines, but not in the susceptible ones. This result indicates that Pgp is more expressed in these antimony-resistant Leishmania lines, whereas the Pgp levels in the susceptible lines were not high enough for detection. On the other hand, this antibody recognized Pgp in both antimony-susceptible and -resistant L. infantum lines. Densitometry using α-tubulin levels for normalization showed the same level of Pgp expression between both L. infantum lines (Fig. 4C). Additionally, we observed that the C219 antibody also detected another polypeptide of about 48 kDa in all antimony-susceptible and -resistant Leishmania lines analyzed. However, the intensity of this polypeptide was very faint for both L. braziliensis lines (Fig. 4B). The presence of this 48 kDa polypeptide can be due to antibody recognition of a common epitope of another protein of Leishmania or an original fragment of Pgp. It is described in the literature that this same anti-Pgp C219 antibody recognizes smaller polypeptides named as “Pgp-like” components (Grogol et al., 1991; Murta et al., 2001; Anacleto et al., 2003).

It is important to emphasize that the anti-Pgp antibody did not detect the protein of 170 kDa in any of the two L. braziliensis lines. This result may be due to the absence of overexpression of Pgp in the LbSbR line or changes may have occurred in the amino acid sequence of the epitope region of the protein, preventing its
obtained for LbWTS and LbSbR lines was 2 mg/mL and after pre-treatment with the
was separated by SDS-PAGE on 12% gel and stained with Coomassie blue. (B) Western blot analysis using the monoclonal antibody C219. (C) The membrane was normalized with the α-tubulin antibody.

3.4. Intracellular accumulation of antimony

In order to compare the level of antimony uptake between antimony-susceptible and -resistant Leishmania lines, the parasites were incubated for 1 h in the presence of 540 μM antimony and, after washing, the intracellular antimony level was quantified by graphite furnace atomic absorption spectrometry. Fig. 5 shows that the uptake of antimony in the resistant lines was significantly reduced in three out of four species analyzed. The data indicate that the antimony levels were approximately seven-, five- and two-fold lower in the antimony-resistant L. guyanensis (Fig. 5A), L. amazonensis (Fig. 5B) and L. braziliensis (Fig. 5C) lines, when compared to their susceptible pairs, respectively. However, the susceptible and resistant lines of L. infantum (Fig. 5D) showed no statistically significant difference in the antimony incorporation. The graphics also show marked differences in the level of antimony incorporation between the different species studied.

In addition, these results of antimony intracellular accumulation demonstrate that there is an inverse correlation between the intrinsic antimony susceptibility of the Leishmania wild-type lines used in this study and their ability to accumulate antimony. For example, the LgWTS line, which is relatively more Sb sensitive when compared to LiWTS (IC_{50} 0.09 μM versus IC_{50} 0.33 μM, respectively, a 3.7-fold difference), accumulated 14-fold less Sb. The same remark can be made for other lines of Leishmania. The LgWTS line accumulated 2.8-fold less Sb than the LaWTS line (IC_{50} 0.28 μM, a 3-fold difference) and the LbWTS line (IC_{50} 0.15 μM) accumulated 5-fold less Sb than the LiWTS line (a 2.2-fold difference in IC_{50}).

3.5. Susceptibility of L. braziliensis lines to BSO and the effect of this inhibitor in the SbIII uptake

We also investigated the effect of BSO, an inhibitor of γ-glutamylcysteine synthetase (GCS) enzyme, in the intracellular accumulation of antimony by LbWTS and LbSbR lines pre-treated with this inhibitor. Interestingly, the BSO susceptibility assay showed that the LbSbR line is more susceptible to BSO than its susceptible counterpart LbWTS. The IC_{50} obtained for LbWTS and LbSbR lines were 15 mM and 2.5 mM, respectively. Then, these lines were pre-treated with BSO during 48 h, incubated with SbIII by 1 h and then submitted to antimony uptake measurement (uptake assay). Our results revealed that LbSbR line accumulated more SbIII compared to its susceptible pair (LbWTS) (Fig. 5E). On the other hand, without BSO, the LbSbR line accumulates less SbIII (Fig. 5C). These results suggest that BSO probably decreases the intracellular concentration of thiols, interfering directly on the SbIII-thiol complex formation and leading to the SbIII accumulation. In addition, we determined the SbIII IC_{50} for the LbWTS and LbSbR lines in the absence and presence of BSO. The LbWTS and LbSbR lines were pre-treated with 15 mM and 2.5 mM BSO, respectively, during 48 h. Subsequently, we incubated these parasites with different SbIII concentrations for 48 h. The IC_{50} obtained for the LbWTS line in the absence of BSO was 0.025 mg/mL and in the presence of BSO was 0.00625 mg/mL (4-fold lower). On the other hand, for the LbSbR line the IC_{50} was 2 mg/mL and after pre-treatment with BSO it was 0.0625 mg/mL (32-fold lower), indicating that the BSO reversed the SbIII resistance phenotype.

3.6. Influx and efflux of antimony

Initially, we performed influx kinetics analysis for the lines of L. guyanensis and L. braziliensis. Our results showed that both SbR lines exhibited lower rate of influx of SbIII than their respective susceptible counterparts, the difference being much more pronounced in the case of L. guyanensis species (Figs. 6A and B). In order to compare the efflux of Sb between susceptible and resistant lines, cells were first loaded with about the same level of antimony. Subsequently, the parasites were washed with HG buffer and aliquots were taken at the times 0, 1 and 2 h to analyze the efflux of antimony by graphite furnace atomic absorption spectrometry.

![Fig. 4. Pgp expression levels in Leishmania species. (A) Electrophoretic profile of proteins from antimony-susceptible (WTS) and -resistant (Sb-R) Leishmania lines. Proteins were separated by SDS-PAGE on 12% gel and stained with Coomassie blue. (B) Western blot analysis using the monoclonal antibody C219. (C) The membrane was normalized with the α-tubulin antibody.](image)
The results demonstrate that the rates of SbIII efflux were higher in both antimony-resistant lines of *L. guyanensis* and *L. braziliensis*, when compared to their susceptible pairs (Fig. 6C and D, respectively). This efflux data explains, at least in part, the lower SbIII concentration found in both SbIII-resistant lines, as presented in the Fig. 5A and C, respectively.

### 3.7. Decreased expression of AQP1 in SbR Leishmania

In order to investigate whether the lower level of intracellular antimony in the SbIII-resistant lines was due to down-regulation of AQP1 protein, we evaluated the expression level of this protein in the four *Leishmania* species studied. Western blot analyses with membrane proteins fractions were performed using a rabbit polyclonal antibody anti-AQP1. Our results showed that this antibody recognized a polypeptide of approximately 35 kDa in all antimony-susceptible and -resistant lines of *Leishmania* analyzed (Fig. 7). After normalization using an α-tubulin antibody, quantification of the band intensity demonstrated that AQP1 expression level is decreased 1.7 and 3-fold in SbIII-resistant lines of *L. amazonensis* and *L. guyanensis*, respectively, when compared to their respective susceptible pairs (Fig. 7). These data are consistent with the lowest SbIII accumulation presented by these two lines among the species studied (Fig. 5). On the other hand, the expression level of AQP1 was similar between the SbIII-susceptible and -resistant lines of *L. braziliensis* and *L. infantum* showing band intensities ratio (WT/SbR) values of 0.9 and 0.8, respectively (Fig. 7).

### 4. Discussion

The phenomenon of resistance to antimonials in *Leishmania* is complex, multifactorial and involves several pathways, which have
similar features with other microorganisms. These pathways include the entry, metabolism, efflux and/or sequestration, as well as cell death through the drug action (Jeddi et al., 2011). MRPA is one of the most studied ABC transporters in Leishmania and its role in the resistance has been well established in vitro (Ouellette et al., 1990). However, the majority of these studies were performed using Old World drug-resistant Leishmania lines.

Our PFGE results showed that the MRPA gene is located in an 800 kb band corresponding to chromosome 23 (795 kb) of Leishmania, corroborating literature data (Leprohon et al., 2009; Monte Neto et al., 2011). Interestingly, the MRPA gene probe also recognized two other bands of different sizes, 200 and 1500 kb, only in the antimony-resistant L. braziliensis line. These supplementary bands indicate that MRPA is rearranged or present as an extrachromosomal amplification in LbSbR (Fig. 1B). Additionally, the smear observed exclusively in this resistant line, provide evidence of circular DNA amplification as indicated by this particular migration pattern in PFGE possibly corresponding to various topoisomers of the circles (White et al., 1988). According to Beverley (1988), large supercoiled circular DNAs appear to exhibit unusual electrophoretic mobility in PFGE, when compared to the large linear chromosomes. Then, an explanation for the presence of approximately 1500 kb chromosomal band in the antimony-resistant L. braziliensis line may be related to the integration of this band into large linear chromosomal DNA or the formation of large concatenated networks of circular DNA. Moreover, Grondin et al. (1998) demonstrated the formation of extrachromosomal circular DNA amplification derived from precursors linear amplicons in methotrexate-resistant L. tarentolae. This possibility could explain the presence of additional bands concomitant with the smear detected by MRPA gene probe in the same LbSbR line. It is also important to note that the MRPA gene probe recognized a band of approximately 700 kb only in the antimony-resistant L. infantum line. It is described in the literature that the variability in the chromosomal...
the MRPA transporter is overexpressed in axenic amastigotes of *L. braziliensis* (Teixeira, 1998). It was shown by targeted DNA microarray that transcriptional control that decreases the RNA levels of this gene its susceptible counterpart. This could also be due to post-translational modification of the MRPA protein (El Fadili et al., 2001). Increased expression between antimony-susceptible and -resistant lines of *L. amazonensis* and *L. braziliensis* lines (Fig. 5). Literature data demonstrated that antimony-resistant *L. infantum* and *L. panamensis* lines incorporated less antimony, when compared to their susceptible pairs (Brochu et al., 2003). It is possible that another transporter may be involved in the low accumulation of antimony in resistant *Leishmania* lines. On the other hand, no significant difference of antimony incorporation was observed between the antimony-susceptible and -resistant lines of *L. infantum*. This can be explained due to the low resistance index of this antimony-resistant line. These data of antimony accumulation suggest the presence of different antimony-resistance mechanisms among these *Leishmania* species analyzed.

We investigated the SbIII efflux rates in lines of *L. guyanensis* and *L. braziliensis*. The results revealed that the efflux rates are higher in the antimony-resistant lines of these parasites, contributing to the lower accumulation of SbIII in these *Leishmania* species studied. Although, the SbIII-resistant *L. braziliensis* line present low antimony concentration, the level of expression of AQPI was similar between both SbIII-susceptible and -resistant lines, suggesting that AQPI is not involved in the antimony-resistance phenotype. However, the SbIII-resistant *L. braziliensis* line showed an increased rate of antimony efflux, which could explain in part the lower SbIII accumulation in this line.

Interestingly, susceptibility test to BSO, an inhibitor of γ-glutamylcysteine synthetase (GCS), showed that this inhibitor reversed the SbIII-resistance phenotype of *L. braziliensis*. The LbSbR line was more susceptible to BSO than its susceptible counterpart LbWTS. More importantly, with BSO pre-treatment, the resistant parasites accumulated more antimony than its wild-type counterpart, suggesting this inhibitor interferes in the polyamine metabolism, changing the intracellular concentration of thiols and antimony. Indeed, in vitro studies have shown that BSO, a specific inhibitor of GCS, an enzyme involved in glutathione and trypanothione biosynthesis, can reverse resistance to trivalent antimony in the parasite *L. tarentolae* (Grondin et al., 1997).

In conclusion, our data show that the antimony resistance mechanisms are different in the New World *Leishmania* species analyzed in this study. Functional analysis studies will be performed to investigate the involvement of the MRPA gene in our *Leishmania* samples.
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