Molecular Preadaptation to Antimony Resistance in *Leishmania donovani* on the Indian Subcontinent

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ABSTRACT Antimonials (Sb) were used for decades for chemotherapy of visceral leishmaniasis (VL). Now abandoned in the Indian subcontinent (ISC) because of *Leishmania donovani* resistance, this drug offers a unique model for understanding drug resistance dynamics. In a previous phylogenomic study, we found two distinct populations of *L. donovani*: the core group (CG) in the Gangetic plains and ISC1 in the Nepalese highlands. Sb resistance was only encountered within the CG, and a series of potential markers were identified. Here, we analyzed the development of resistance to trivalent antimonials (SbIII) upon experimental selection in ISC1 and CG strains. We observed that (i) baseline SbIII susceptibility of parasites was higher in ISC1 than in the CG, (ii) time to SbIII resistance was higher for ISC1 parasites than for CG strains, and (iii) untargeted genomic and metabolomic analyses revealed molecular changes along the selection process: these were more numerous in ISC1 than in the CG. Altogether these observations led to the hypothesis that CG parasites are preadapted to SbIII resistance. This hypothesis was experimentally confirmed by showing that only wild-type CG strains could survive a direct exposure to the maximal concentration of SbIII. The main driver of this preadaptation was shown to be MRPA, a gene involved in SbIII sequestration and amplified in an intrachromosomal amplicon in all CG strains characterized so far. This amplicon emerged around 1850 in the CG, well before the implementation of antimonials for VL chemotherapy, and we discuss here several hypotheses of selective pressure that could have accompanied its emergence.

IMPORTANCE The “antibiotic resistance crisis” is a major challenge for scientists and medical professionals. This steady rise in drug-resistant pathogens also extends to parasitic diseases, with antimony being the first anti-*Leishmania* drug that fell in the Indian subcontinent (ISC). Leishmaniasis is a major but neglected infectious disease with limited therapeutic options. Therefore, understanding how parasites became resistant to antimonials is of commanding importance. In this study, we experimentally characterized the dynamics of this resistance acquisition and show for the first time that some *Leishmania* populations of the ISC were preadapted to antimony resistance, likely driven by environmental factors or by drugs used in the 19th century.

KEYWORDS *Leishmania*, antimonials, drug resistance mechanisms, genomics, metabolomics

Leishmaniasis is one of the 17 neglected tropical diseases (NTDs) recognized by the WHO and is responsible every year for the death of about 30,000 persons in 98 countries where the disease is endemic, where 350 million people are at risk of...
contracting the disease (1). On the Indian subcontinent (ISC), leishmaniasis is present in its more lethal form, visceral leishmaniasis (VL). Its etiological agent, *Leishmania donovani*, is a dimorphic parasite transmitted by a phlebotomine vector (1, 2). Since 1923, pentavalent antimonials (Sb V) have been used to treat VL in the ISC and in other regions of the world. Sb V is an immunomodulator active on the macrophage (3), and its reduced form, Sb III, is directly toxic for the parasite (4, 5). However, antimonials have lost their efficacy in the ISC toward the end of the 20th century, among others, because of drug resistance (DR) (6). In the ISC, antimonials were then replaced by miltefosine (MIL), but the efficacy of this drug is also declining (7): the first cases of MIL resistance were recently reported (8). A few drugs remain available (amphotericin B and paromomycin), and in the expectation of new compounds, it is essential to protect the current arsenal, among others, by using new formulations or combination therapy. Moreover, it is important to understand in depth the mechanisms and dynamics of DR.

A recent phylogenomic study of 204 *L. donovani* isolates shed new light on the emergence and spreading of antimonial-resistant (Sb-R) parasites in the ISC (9). First, the study described a natural history frame for the VL epidemics: the main population of *L. donovani* emerged about 150 years ago in northeast India, and after a succession of bottlenecks and expansions, it generated a series of subpopulations, currently defined as ISC2 to ISC10 and gathered within a main core group (CG). In addition, another genetic group (ISC1), which diverged much earlier from the CG, was discovered in the highlands of Nepal. Second, the study highlighted a series of features relevant in the context of Sb resistance. (i) All 191 isolates (Sb-R or Sb sensitive [Sb-S]) from the CG harbored two intrachromosomal amplifications (ICAs) of sets of adjacent genes, called the M and H loci; these genes were not amplified in isolates from the ISC1 subpopulation. The H locus was reported elsewhere to be a major driver of Sb III resistance because of the presence of one gene, *MRPA*, encoding a member of the multidrug resistance protein family that sequesters Sb-thiol complexes into intracellular vesicles (10–12). (ii) Within the CG, a particular genetic group, ISC5, concentrates most of the strains isolated from antimonial treatment failure patients. All 52 strains of that group harbor a 2-nucleotide (nt) insertion in the AQP1 gene, creating a nonfunctional variant of this transporter known to be involved in the uptake of Sb III (13, 14). (iii) Other genomic adaptations to Sb III, involved in detoxification mechanisms and increased levels of thiols reported in experimental resistance (15), had not been encountered (yet) in the natural population under study.

Considering the anthroponotic nature of VL in the ISC and the high rate of transmission documented in the Gangetic plains, we may assume that the CG represents the parasite population that has been under strong antimonial pressure over decades. The situation might be different in the highlands, where ISC1 was documented. In this region, VL cases were sporadic and *L. donovani* transmission has only been reported recently (16); hence we may hypothesize that antimonial pressure was historically lower in these regions than in the lowlands. In this context, the present study aimed at answering two main questions. First, do the dynamics of drug resistance emergence differ depending on the genetic background, known to be different between ISC1 and CG strains? Second, do we observe similar molecular adaptations when selecting antimonial resistance in the laboratory compared to natural selection in the field, and is the succession of events comparable?

These questions were answered, as a first step, in the context of Sb III exposure. We first measured the intrinsic Sb III susceptibility of promastigotes from 10 *L. donovani* strains representing the genetic diversity in the ISC. We then identified three reference strains among these (one from ISC1 and two from the CG) and experimentally selected Sb III resistance at the promastigote stage. We analyzed two parameters: (i) the time to resistance (i.e., the number of weeks required upon stepwise selection to achieve resistance) and (ii) the type of resistance (i.e., the nature and dynamics of molecular adaptations developed by resistant parasites). This was done by untargeted genomic and metabolomic analyses and targeted transcript-level analyses. Validation of the main drivers of resistance was done by overexpression of candidate genes.
RESULTS

Intrinsic promastigote SbIII susceptibility of L. donovani strains from ISC1 is higher than that of the core group. We first assessed the intrinsic SbIII susceptibility of promastigotes representative of the genomic diversity encountered in Nepal (9). For this, we chose three cloned strains of the ISC1 group and seven strains of the CG (from the ISC3, -4, -5, -6, and -9 groups). The in vitro susceptibility assay revealed a similar 50% effective concentration (EC50) to SbIII among CG promastigotes, with a mean value of 97.8 ± 19.9 μM; this contrasted with the different sodium stibogluconate (SSG) susceptibility profiles observed for intracellular amastigotes of these strains shown in previous studies (9). In contrast, promastigotes from ISC1 were more susceptible to SbIII than promastigotes from the CG, with a mean value of 24.4 ± 4 μM (Fig. 1; see Fig. S1 in the supplemental material). The difference observed between CG and ISC1 strains was statistically significant ($P = 3.2 \times 10^{-8}$ by analysis of variance [ANOVA]).

Stepwise selection of SbIII resistance in promastigotes. We aimed to compare the dynamics of SbIII resistance selection in three L. donovani strains differing in (i) intrinsic promastigote SbIII susceptibility, (ii) intracellular amastigote SbV susceptibility (sensitive BPK026/282 versus resistant BPK275), and (iii) genomic background (see Table S1 in the supplemental material). The selection process was performed in quadruplicates (each replicate called line A, B, C, or D), starting with 47.75 μM SbIII for both CG strains, a concentration close to their EC50. For BPK026, the selection process was initiated with 6 μM SbIII, as its EC50 was much lower. The two CG strains showed similar selection dynamics, with the four replicates of each strain successfully reaching the highest SbIII concentration (382 μM) within four rounds of selection (Fig. 2A). These lines will be further called BPK282 SbIII-R and BPK275 SbIII-R. The selection dynamics for BPK026 were rather different: one replicate (line D) did not survive the first selection round, line A was cleared during the third round of selection (48 μM), and line C did not survive the sixth round (191 μM). Thus, one replicate only (line B) was successfully selected to survive the highest SbIII concentration (further called BPK026 SbIII-R) (Fig. 2B). Time to resistance was defined as the time needed for each line to display a wild-type (WT) growth curve in the presence of 382 μM SbIII, following the stepwise selection process. Accordingly, time to resistance was estimated to 20 weeks for the two CG strains, whereas 35 weeks were required for BPK026 line B to reach the same level (Fig. 2C). In order to evaluate the stability of the resistance phenotype, all the SbIII-resistant lines as well as their parental strains were maintained for 20 weeks without SbIII. The withdrawal of the drug pressure did not impact their growth rate nor their SbIII susceptibility (EC50 > 382 μM) (data not shown).

To fully characterize the antimony resistance profile of the selected lines, we assessed the SbV susceptibility of intracellular amastigotes by applying SSG pressure on
infected macrophages. Analysis of parental lines confirmed previous results: (i) intracellular amastigotes of both BPK026 and BPK282 clearly showed a dose-response curve typical of SbV-sensitive strains, with EC50s of 42.46 and 27.51 μM, respectively (see Fig. S2 in the supplemental material); (ii) in contrast, for BPK275, similar infection indices
were observed from 0 to 246.4 μM, characteristic of Sb\textsuperscript{V}-resistant amastigotes. After Sb\textsuperscript{III} resistance selection of promastigotes, BPK275 Sb\textsuperscript{III}-R amastigotes remained resistant to Sb\textsuperscript{V}; BPK026 Sb\textsuperscript{III}-R and BPK282 Sb\textsuperscript{III}-R both showed a more than 6-fold-increased EC\textsubscript{50} to Sb\textsuperscript{V} (276.7 and 170 μM, respectively) (Fig. S2).

**Untargeted genomic analysis during stepwise Sb\textsuperscript{III} resistance selection.** For all parasite lines, samples were collected at each selection round for whole-genome sequence analysis of nuclear DNA (see Fig. 2A and B for details). We first looked at the DNA sequence itself to detect single nucleotide polymorphisms (SNPs) and small nucleotide insertions or deletions (indels). The average coverage of all the analyzed samples was 13.5 x ± 3.9, ranging from 7.4 to 24.9 x, and >98% of the genome was covered. All along the selection process, the two CG strains did not show any de novo SNP or indel in the whole nuclear genome; among 303 heterozygous sites in the CG strains, no significant changes in allele frequency were detected. In BPK026 Sb\textsuperscript{III}-R line B, we did not observe any new indel during the selection process, but in two different genes, a heterozygous SNP appeared at round 4, under an Sb\textsuperscript{III} pressure of 48 μM. These two de novo SNPs, respectively, introduced a stop codon at position 1699 of LdBPK\_210017400 (formerly LdBPK\_211220.1), coding for an intraflagellar transport protein, and a nonsynonymous mutation, H1104T, in LdBPK\_120011000 (formerly LdBPK\_120530), coding for a fusaric acid resistance protein-like protein. In subsequent selection rounds, the frequency of the mutated alleles progressively increased together with the Sb\textsuperscript{III} concentration (Fig. 3A), from 0.067 to 0.667 (LdBPK\_210017400) and from 0.5 to 1 (LdBPK\_120011000). Among the 11,189 heterozygous sites of BPK026, no change in allele frequency was observed during the selection process.

In a second stage, we looked at genome structure changes, more specifically somy and local copy number variations (CNVs), both known to occur when *Leishmania* is under selective pressure. The karyotype variability contrasted sharply with the conservation of the nucleotidic sequence itself reported above. However, this variability was much higher in BPK026 than in the two CG strains, with 153 significant somy changes among all lines of BPK026 versus 31 and 28 significant changes in BPK282 and BPK275, respectively (Fig. 3B, C, and D). Interestingly, at each round of selection, karyotypes of distinct replicate lines were rather similar for the CG strains, while different "scenarios" of aneuploidy were followed by the three replicate lines of BPK026. Five chromosomes (8, 18, 23, 25, and 31) showed an increase in the statistical significance of chromosome copy number (S value) in one or more lines of each strain, the most extensive increases being observed for chromosomes 23 and 31 (Fig. 3; see Data Set S1 in the supplemental material).

Interestingly, the S value increase of chromosome 23 in BPK026 Sb\textsuperscript{III}-R was due to a strong increase in the read depth of a large chromosomal segment (from positions 10048 to 567768 (Fig. 3E)): at 382 μM, this fragment showed a copy number of 7.1 ± 1.3, while the small segment (from positions 569158 to 768383) showed a copy number similar to that of the WT (3.0 ± 0.4). These additional copies of the large segment of chromosome 23 resulted in the specific amplification of 140 open reading frames (ORFs), of which 52.85% are putatively annotated and 47.15% correspond to hypothetical proteins. Interestingly, among these genes, 6 (3.8%) were coding for proteins involved in redox pathways, while none of these were found in the small fragment. In addition, several genes of the large fragment have been previously reported to be involved in (i) drug resistance or (ii) virulence. (i) The large fragment genes associated with drug resistance were LdBPK\_230007200 and \_7300 (previously LdBPK\_230230.1), coding for pentamidine resistance protein, LdBPK\_230007700 and _7800 (previously LdBPK\_230280 and _290, respectively), associated with the terbinafine resistance locus and MRPA within the H locus, and LdBPK\_230009800 (previously LdBPK\_230500), coding for trypanothione synthetase. (ii) The large fragment genes associated with virulence were LdBPK\_230018500 and _18600, encoding hydrophilic acylated surface proteins a and b (previously not assembled), and LdBPK\_230020400 (previously LdBPK\_231430), coding for membrane-bound acid phosphatase 2. All other protein...
FIG 3  Genomic analysis of the selected Sb<sup>III</sup>-resistant strains. (A) Evolution of allele frequency over the Sb<sup>III</sup> resistance selection process of BPK026 for a fusaric acid resistance protein-like protein (LdBPK_120011000-H1104T) and the intraflagellar transport protein (LdBPK_210017400-Δ1699). (B and C) Heat map representing the karyotype dynamics across the Sb<sup>III</sup> resistance selection of BPK282 and BPK275. (D) High hierarchical representation of BPK026 karyotype evolution. Heat maps show median normalized read depths of chromosomes found within each cell population for each of the 36 chromosomes (y axis) and each sample (x axis), replicate populations called A, B, C, and D are shown for each selecting Sb<sup>III</sup> concentration (from 6 to 382 μM), and the samples “WT” framed in a black box represent the parental Sb<sup>III</sup>-sensitive population for each strain. The color key for panels B and C shows the normalized chromosome read depth (S). S ranges are as follows: monosomy, S ≤ 1.5; disomy, 1.5 ≤ S < 2.5; trisomy, 2.5 ≤ S < 3.5; tetrasomy, 3.5 ≤ S < 4.5; pentasomy, 4.5 ≤ S < 5.5. In panel D, the ranges are as follows: monosomy, S ≤ 1.5; disomy, 1.5 ≤ S < 2.5; trisomy, 2.5 ≤ S < 3.5; tetrasomy, 3.5 ≤ S < 4.5; pentasomy, 4.5 ≤ S < 5.5.
categories analyzed by Gene Ontology were represented in comparable proportions in both fragments (Data Set S2A and B).

Local CNVs like the H and M loci, which are constitutive of the CG (9, 17), did not vary in copy number in the CG lines: they also did not appear as intra- or extrachromosomal amplifications in BPK026 SbIO-R (Fig. 3F). Altogether, the untargeted genomic analysis revealed that along the selection process, BPK026 developed more molecular adaptations than the two CG strains, including amplification of genes known to be involved in drug resistance and virulence.

Targeted transcript-level analysis of SbIO-resistant selected lines and validation of main drivers. Given the general amplification of the H and M loci in the CG strains (9, 17), as well as the observed changes in chromosome 23 during the SbIO selection process, we performed quantitative reverse transcription (RT)-PCR assays to compare the expression levels of genes present in the H and M loci in the parental and SbIO-resistant lines generated here. With respect to the M locus, we observed a higher expression of the two tested genes (Hyp36 and MPK1) in parental (WT) CG lines versus BPK026, which reflected the intrachromosomal amplification of the M locus; however, SbIO resistance in both BPK026 and the CG lines was not accompanied by a further increase of the expression of these genes (Fig. 4A). The H locus was the most interesting target; as for the M locus, levels of MRPA, HTBF, Hyp23, and ASS transcripts were slightly higher in WT CG lines than in WT BPK026 (1.5- to 3.0-fold increase), which also reflected...
the intrachromosomal amplification of the H locus in CG lines. After selection of SbIII resistance, there were no significant changes in the expression of these genes within the CG lines, but in BPK026, the levels of the four genes increased 1.5- to 3.5-fold, reaching the expression levels of the CG lines (Fig. 4A). This fit with the increased copy number observed for the large fragment of chromosome 23 (harboring among others the H locus) mentioned above.

In order to confirm the importance of the H locus and its amplification for SbIII resistance (through intrachromosomal amplification or partial increase of chromosome 23’s genomic material), the full H locus was overexpressed in BPK026 promastigotes using a cosm id carrying that region; similar overexpression of the M locus was done as control. This experiment revealed that overexpression of the H locus was associated with an increase of SbIII tolerance of about 3-fold \( P = 6.6 \times 10^{-3} \) by ANOVA, in contrast to overexpression of the M locus, which did not impact drug susceptibility. We further dissected the importance of the different genes constituting both H and M loci, by overexpressing them separately in BPK026. For one gene only, MRPA, overexpression was associated with increased SbIII tolerance (3-fold, \( P = 1.71 \times 10^{-3} \) by ANOVA) comparable to that of the CG isolates (Fig. 4B; see Fig. S3 in the supplemental material).

**Untargeted metabolomics analysis of SbIII-resistant selected lines.** An untargeted metabolomics analysis was performed on the three parental lines, their respective fully resistant lines (382 \( \mu \)M), and several intermediate resistant lines (12 and 24 \( \mu \)M) of BPK026 line B. In total, we detected 300 metabolites (see Data Set S3A in the supplemental material). Principal-component analysis (PCA) was carried out as a first exploratory analysis of this metabolomics data set (Fig. 5A). The two first principal components (PC1 and -2) explained 33.02 and 16.03% of the total variation and clearly separated two experimental sets: (i) the BPK026 parental and resistant lines and (ii) the BPK275 parental and resistant lines B, C, and D, together with BPK282 parental and resistant lines B, C, and D. For BPK026, the two intermediate-resistant populations clustered at an intermediate position between the parental strain and the fully resistant line. PCA results were confirmed by analysis of individual metabolites. The numbers of metabolites significantly differing between parental and the fully resistant lines ranged as follows: 52 (BPK026), 24 (BPK282), and 17 (BPK275), highlighting the larger amount of changes occurring in the BPK026 SbIII-R strain (Fig. 5B; Table 1). A comparison of all the resistant lines showed the metabolic similarity of the CG strains BPK282 and BPK275 (Fig. 5A and B; Data Set S3A and C). This general trend was confirmed when analyzing specific classes of metabolites; among them two classes represented significant common changes during all the comparisons we performed—glycerophospholipids (GPLs) and amino acids.

GPL composition evolved differently over the selection process of the different strains: we observed 13 changes between BPK026 WT and BPK026 SbIII-R, while only four changes were detected in BPK282 SbIII-R compared to the BPK282 WT, and lipid composition was unchanged between all BPK275 lines. When the GPL composition of the BPK026 WT was compared to the two WT strains of CG taken together, we observed smaller amounts of 10 GPLs in the former (three unsaturated/low unsaturated and seven highly unsaturated), and 3 were more abundant (low unsaturated). When doing the same comparison between resistant lines of the three strains, no difference was observed (Table 2). This indicated that upon SbIII resistance selection of BPK026, the levels of 13 GPLs evolved toward levels encountered in CG lines.

Amino acids were also significantly altered in the current experimental set. Again, more amino acids changed during SbIII resistance selection of BPK026 \( (n = 4) \) than in BPK282 \( (n = 2) \) and BPK275 \( (n = 2) \). These 4 amino acids deserved particular attention due to their variation in abundance as well as their biological importance. Two amino acids varied in all resistant lines versus their WT parental strain: (i) proline abundance increased 6.76-fold in BPK026 SbIII-R, 2.8-fold on average in BPK282 SbIII-R, and 2-fold in average in BPK275 SbIII-R lines; (ii) leucine abundance decreased 2-fold in BPK026 SbIII-R, 1.9-fold in BPK282 SbIII-R lines, and 1.7-fold in BPK275 lines. Arginine was less...
FIG 5 Metabolomics analysis of the selected SbIII-resistant strains. (A) Principal-component analysis based on 300 metabolites and including the initial parental strains (WT) BPK026 (green), BPK275 (red), and BPK282 (blue), the final populations resistant to 382 μM SbIII (BPK026 382B, BPK275 382B, -C, and -D, and BPK282 382B, -C, and -D) as well as BPK026 intermediates (12B and 24B, respectively, resistant to 12 and 24 μM SbIII). (B) Venn diagrams showing metabolites significantly different between strains. (C) High hierarchical representation of amino acid abundance in the different strains, including the parental strains (BPK026 WT, BPK275 WT, and BPK282 WT) and the selected populations resistant to 382 μM SbIII (BPK275 382B, -C, and -D, BPK282 382B, and BPK282 382B, -C, and -D).
| Metabolite class | No. and name(s) of metabolites from $a$ | BPK206 Sb III-R/BPK206 WT | BPK282 Sb III-R/BPK282 WT | BPK275 Sb III-R/BPK275 WT |
|-----------------|---------------------------------------|---------------------------|---------------------------|---------------------------|
| **Increased**   |                                       |                           |                           |                           |
| Amino acids, peptides, and amino acid conjugates | 9: arginine, proline, isoleucine, glutamylamino-butanooate, $\text{N}^\beta\text{-succinyl-ornithine, glutamylcysteine, prolyl-proline, phenylalanine, glutamylalanine}$ | 1: proline | 2: acetyl-lysine, prolylhydroxyproline |                           |
| GLUTs           | 7: 1 saturated or low unsaturated, 6 highly unsaturated | 3: 1 saturated or low unsaturated, 2 highly unsaturated |                           |                           |
| Thiols          | 1: glutathione                         |                           |                           |                           |
| Nucleobases and nucleosides | 23: flumipropyn; magnoshinin; dobenpropit; PC(20:4(5Z,8Z,11Z,14Z)/18:4(6Z,9Z,12Z,15Z)); 2'-hydroxyfurano [2'-3',4',3']chalcone; coenzyme M 7-mercaptopotheanoyl/threonine-phosphate heterodisulfide; PC(22:5(4Z,7Z,10Z,13Z,16Z)/22:5(4Z,7Z,10Z,13Z,16Z)); S-hydroxy-6,7,3',4',5'-pentamethoxyflavanone 5-O-rhamnoside; $N^\beta$-phosphoagmatine; (S)-AMP; 5-glutamylglycine; uplandicine; 1-pyroline; piperidine; 4-aminopyridine; histamine; $\gamma$-Glu–Thr; carnitine; dihydro-4,4-dimethyl-2,3-furandione; heptitol; 3,4,5-trihydroxy-hexanoic acid, rhamnose, deoxygalactopyranose, fucose 1,5-anhydroxyorbitol; 2,2-dimethylsucinic acid, aceto-hydroxybutanoic acid; xylitol, arabitol; 4-guanidinobutanal | 7 | 17 | 13 |
| **Total**       | 40                                    |                           |                           |                           |
| **Decreased**   |                                       |                           |                           |                           |
| Amino acids, peptides, and amino acid conjugates | 2: proline-betaine; trimethyl-lysine |                           | 4: glutamylalanine; succinylidiaminopimelate, $N^\beta$-succinyl-arginine; X-amino-pentanoic acid, valine | 3: isoleucine, glutamylcysteine, argininosuccinate |
| GLUTs           | 5: 4 saturated or low unsaturated, 1 highly unsaturated |                           | 1: trypanothione-disulfide | 1: phosphorylthanolamine |
| Thiols          |                                       |                           |                           |                           |
| Nucleobases and nucleosides |                           |                           | 12: allantoin; 2-ethyl-2-hydroxybutyric acid, 2-hydroxy-3-methylpentanoic acid, hydroxy(iso)caproic acid, X-hydroxy-hexanoic acid; $N$-acetylglutamine; $\gamma$-Asp–Asp–Pro; 5-$\gamma$-glutamylglycine; dobenpropit; guvacine; (S)-AMP; cholinephosphate; fagomine; dehydroalanine; SP(34:0) | 1: IMP |
| Others          | 5: creatine; cyclohexane-1,3-dione; spisulosine, 1-deoxy-sphinganine; 2-keto-3-methyl-valerate, 2K-4CH₃-pentanoic, X-oxo-X-methyl-pentanoic acid, 2-ketoscaproate, X-oxo-hexanoic acid; 5-adenosylmethionine |                           |                           |                           |
| **Total**       | 12                                    | 17                        | 13                        |                           |

*aOnly the biologically (2-fold changes) and statistically ($p < 0.05$) significant differences are indicated.*
Table 2: Changes in glycerophospholipid abundance

| GPL type                        | Change in abundance for comparison showna |
|---------------------------------|------------------------------------------|
|                                 | BPK026 SbIII-R vs BPK026 WT             | BPK282 SbIII-R vs BPK282 WT | BPK275 SbIII-R vs BPK275 WT | BPK026 WT vs BPK282 WT | BPK026 WT vs BPK275 WT | BPK026 SbIII-R vs CG SbIII-R |
| Saturated and low unsaturated (≤3) | Up 0 Down 1 | Up 0 Down 1 | Up 0 Down 3 | Up 0 Down 0 | Up 0 Down 0 | Up 0 Down 0 |
| High unsaturated (≥3)           | Up 8 Down 4 | Up 3 Down 0 | Up 4 Down 3 | Up 3 Down 3 | Up 0 Down 0 | Up 0 Down 0 |

a“Up” represents a log2 FC of >1, and “Down” represents a log2 FC of <1.

bBPK026 SbIII-R population B.
cThe values shown for BPK282 and BPK275 SbIII-R are the average values from the three resistant replicates (populations B, C, and D).
dThe value shown for CG SbIII-R is the average of the values for BPK282 and BPK275 SbIII-R replicates.

abundant in the BPK026 WT compared to the CG strains (5.7-fold decrease compared to the BPK282 WT and 8.8-fold decrease compared to the BPK275 WT); interestingly, the levels of arginine did not change significantly during the SbIII selection of CG lines, but it increased 5.7-fold in BPK026 SbIII-R versus the WT parental strain (Fig. 5C; Data Set S3A), thus reaching the levels encountered in CG lines. Finally, lysine was more abundant in the BPK026 WT than in the CG WT strains and dropped drastically (9.2-fold) in BPK026 SbIII-R, while levels did not change in CG resistant lines.

**Adaptation of *L. donovani* strains to a high concentration of SbIII.** The previous results suggested that promastigotes of CG strains are better prepared than BPK026 to respond to SbIII exposure. To test this preadaptation hypothesis under extreme conditions, we assessed the possibility for the parasites to survive the highest concentration of SbIII without the stepwise selection previously undertaken. The 10 strains described above and representing ISC1 (BPK026, BPK156, and BPK031) and CG (BP067, BP206, BP275, BP173, BP282, BP178, and BP294) (Fig. 1; Table S1) were used for a “flash selection,” meaning direct exposure of a wild-type strain to 382 μM SbIII during 5 weeks, passaging the parasites every 7 days and performing a growth curve for 5 weeks. Interestingly we observed from the first day a drastic reduction of parasite numbers for all strains and already a complete loss of BPK031 (ISC1). The two other ISC1 strains suffered of a drastic decrease, but living parasites could still be observed after 7 days in at least one replicate. After 5 weeks, no living parasites were detected in any of the three ISC1 strains. In sharp contrast, all seven CG strains recovered their original growth rate after four passages only (Fig. 2A and C and 6).

**DISCUSSION**

The objective of our study was to analyze experimentally the dynamics of emergence of SbIII resistance in ISC strains from different genetic backgrounds, to characterize molecular adaptations developed by resistant lines, and to compare them to the adaptations encountered in a clinical context. Our study was performed in vitro with promastigotes and the reduced form of antimony, SbIII. As such, it does not *a priori* take into account resistance mechanisms related to the immunomodulatory effect of SbV nor the reduction of SbV to SbIII, but mechanisms related to transport and detoxification of SbIII. This study therefore constitutes a first step in the analysis of *Leishmania* adaptations to antimonials but should be complemented in the future by similar work with amastigotes and SbV.

We showed here that (i) SbIII susceptibility of promastigotes was higher in ISC1 than in the CG (Fig. 1; Table S1); (ii) following a classical drug resistance selection scheme, time to SbIII resistance (Fig. 2C) was higher for ISC1 parasites (35 weeks) than for CG strains (20 weeks); (iii) untargeted genomic and metabolomic analyses revealed that molecular changes associated with the acquisition of SbIII resistance were more numerous in the ISC1 strain than the CG strain; and (iv) during selection, genomic changes were very similar in the replicates of each CG strain, while very different scenarios were observed among replicates of ISC1, only one of them leading to full resistance.
Altogether, these observations led to the hypothesis that CG promastigotes are pre-adapted to SbIII resistance. This hypothesis was experimentally tested by a “flash selection”—i.e., a direct exposure of WT parasites to the maximal concentration of SbIII (Fig. 6; Table S1). The fact that none of the ISC1 strain survived the flash selection while all the CG strains could recover growth comparable to that of the WT after 5 weeks confirmed the preadaptation hypothesis.

Interestingly, based on our phylogenomic study of heterochronous isolates, we estimated the emergence of the Indo-Nepalese populations of the CG around 1900: this was revealed by a Bayesian phylogenetic model in an explicit temporal and spatial framework (9) and supported by historical reports (18). This corresponds more or less to the first use of SbIII in clinical practice (1919), before being replaced by SbV in 1923 because of toxicity (9). In the anthroponotic context of L. donovani transmission in the ISC, it is possible that these 4 years of application of SbIII provided the selective pressure to the Indo-Nepalese ancestors of the CG. However, there are two alternative explanations. On one hand, the widespread contamination of drinking water by arsenic, shown to occur in the ISC (19), could have provided a strong selective pressure. Arsenic and antimonials share the same sequestration mechanism via MRPA overexpression (12). Furthermore, it was demonstrated that L. donovani could develop resistance to sodium stibogluconate (Pentostam) after exposure to arsenic (20). On the other hand, considering (i) the importance of the H locus amplification in the preadaptation here reported, (ii) the occurrence of this intrachromosomal amplification throughout the whole CG (India, Nepal, and Bangladesh), and (iii) the estimated emergence of the whole CG around 1850, we may suggest that this amplification was already present in 1850, at the onset of the first reported epidemics of kala-azar. It is noteworthy that Leishmania was only discovered half a century later, and for decades, this epidemic disease of the ISC was considered “quinine-resistant malaria” (21). In other words, L. donovani was for decades under pressure from other drugs, such as quinine, and this could also have played a role in the preadaptation.

The untargeted genomic approach here used revealed the importance of genome structure variation for parasite adaptation to drug pressure. Despite our efforts to find new genes or mechanisms implicated in SbIII resistance using whole-genome sequencing (WGS) data, we did not identify any SNP/indel, gene copy number variation, or a consistent somy pattern in the 2 CG strains analyzed (BPK282 and BPK275 [Fig. 3B and C]). On the other hand, two SNPs were found in BPK026 as well as significant modulations of the karyotype occurring rather early during the selection process and partial chromosomal amplifications (Fig. 3D, E, and F). This fits perfectly with the findings of Shaw et al. (22), who showed that during experimental selection of miltefosine resistance, aneuploidy was the first adaptation of the parasite, before the appearance and fixation of SNPs. Similar results were encountered in the context of SbIII resistance in
Leishmania major, Leishmania guyanensis, and Leishmania amazonensis (11, 14, 23). Karyotype modulation has been described as a powerful strategy of gene dosage regulation (24), playing a role in environmental stress and in drug resistance (25).

Relative to the two CG strains, aneuploidy changes were more abundant in BPK026 SbIII-R as expected for a less-preadapted or nonpreadapted strain. Among affected chromosomes, two deserved a particular attention, as they were also showing somy changes in one or more CG lines, chromosomes 23 and 31. In BPK026, the increase in the S value of chromosome 23 was due to the increase in copy number of a large fragment, containing 140 genes, among others those of the H locus involved in drug resistance and virulence (Data Set S2A). This is in line with extrachromosomal linear amplification, as reported several times in a subgenus of Leishmania (26). However, the same fragment of chromosome 23 was shown to be amplified during in vitro selection of SbIII-resistant Leishmania (Viannia) guyanensis, which is known to be unable to generate intra- or extrachromosomal amplifications (11). Interestingly, the position of the chromosomal breakpoint corresponded to a strand-switch region and to the localization of the centromere of chromosome 23, as described recently for L. major (27). Accordingly, other explanations for the partial amplification of chromosome 23 in our L. donovani strain should be considered, like increase in somy of the whole chromosome, followed by deletion of a small segment. In BPK282 and BPK275, somy changes of chromosome 23 were less pronounced, and preadaptation likely stems from the already present amplification of the H locus. We experimentally confirmed the link between the amplification of the H locus and a higher tolerance to SbIII by overexpression and suggested that among the genes present in the H locus, MRPA was the driver (Fig. 4B). The case of chromosome 31 is also interesting, as it contains the gene encoding AQP1, known to be the transporter responsible for the uptake of SbIII (13) and inactive in all parasites of ISC5 because of a 2-nt indel (9). Such an indel was not observed in BPK282 and BPK026 during the SbIII resistance selection, and we did not find any other downregulation mechanism at the genomic level (like the subtelomeric deletion of AQP1 as reported elsewhere [28]) or transcriptomic level (quantitative PCR; qPCR [data not shown]). Further work will be required to understand the meaning of the somy increase of chromosome 31 (25). Bringing together our experimental and phylogenomic data (presence of H locus intrachromosomal amplification in all strains of CG), we believe that MRPA is likely the earlier driver of SbIII resistance. Later on, other genomic adaptations likely appear, such as AQP1 inactivation, which is present only in ISC5 and had not emerged (yet) under our experimental conditions.

Untargeted metabolomics analyses provided results complementary to the genomic study. They clearly supported the preadaptation hypothesis of CG strains and showed that during the SbIII selection process, the metabolism of BPK026 evolved toward levels of CG WT strains. This was best illustrated by GPLs and amino acids. GPLs play an important role in membrane fluidity, as shown earlier (29, 30), and thus in drug resistance since it can act on the absorption of the drug. Amino acids were also modulated, notably proline and arginine (Fig. 5C). Proline levels actually increased in all lines following SbIII resistance selection. This is not unexpected since proline has been shown to enhance cell survival during environmental stresses: in particular, proline is required for volume recovery during osmotic stress responses (31). Increased levels of proline have been observed in other SbIII-resistant strains selected in vitro, and addition of proline to the culture medium led to increased parasite tolerance to SbIII—probably because of its protective role against drug-induced oxidative stress (30). Arginine levels, on the other hand, remained unchanged in the SbIII-R CG strains but were increased to the CG level in SbIII-R ISC1, perfectly fitting with the preadaptation hypothesis. Arginine is part of the urea cycle, which leads to the synthesis of ornithine, polyamines, and ultimately the antioxidant trypanothione. Increased levels of thiols and of other metabolites linked to thiol production have been associated with better protection against Sb-induced oxidative stress and were demonstrated in other studies on SSG-R parasites (15, 29, 32). Lysine levels also supported the preadaptation hypothesis, remaining unchanged in WT or SbIII-R CG strains but decreasing to CG levels in SbIII-R ISC1. Further
studies are however required to understand the implication of lysine in parasite susceptibility/resistance to SbIII.

In summary, we reported for the first time the preadaptation of a whole *Leishmania* population to drug resistance. The preadaptation to SbIII shown here does not mean that all the *L. donovani* parasites of the CG would cause clinical resistance to SSG, and many of them did indeed show susceptibility to SbIII (as shown in reference 9 and the BPK282 WT in this article). However, under adequate pressure, they could rapidly become resistant, which was not the case with ISC1 parasites. We believe that MRPA amplification (thus increased sequestration of SbIII) is a first and important line of defense of the parasites, providing them a grade of tolerance to SbIII. Additional adaptive layers—occurring later during the evolution of the CG—were likely required for a higher resistance to SbIII (e.g., AQP1 inactivation, leading to decreased uptake of the drug) or SbV (e.g., the presence of unique glycans driving the subversion of macrophages) and ultimately for full clinical resistance. Our findings highlight the importance of genetic diversity in the development of resistance and the need for close monitoring of parasite populations before and after implementation of a new drug. Similarly, Research and Development of new drugs should take into account this concept of preadaptation and include a challenge of new compounds to parasites with different preadaptation features.

**MATERIALS AND METHODS**

**Strains and culture conditions.** Ten strains representing the diversity of *L. donovani* in the ISC were used in the present study (Table S1). Promastigotes were maintained in HOMEM (Gibco) supplemented with 20% (vol/vol) heat-inactivated fetal calf serum (FCS) at 26°C. Growth curves of the strains were performed by daily manual counting for 7 days using Uriglass counting chambers (A. Menarini Diagnostics).

**Antimony susceptibility tests.** SbIV susceptibility tests were performed using potassium antimony tartrate (Sigma-Aldrich) as the source of SbIV (36.9%). The EC50s were determined on logarithmic-stage promastigotes after 24 h of exposure to SbIV with a resazurin assay as previously described (33). Briefly, promastigotes were exposed to a semilogarithmic concentration range of SbIV from 1 μM to 382 μM. EC50s were calculated with GraphPad Prism using a sigmoidal dose-response model with variable slope.

For the SbV susceptibility assay, primary peritoneal macrophages from Swiss OF1 mice (Charles River, Inc.) were infected with day 7 stationary-phase promastigotes as previously described (33). After 24 h of infection, infected macrophages were washed, and fresh medium containing SSG (Calbiochem) at a concentration range of SbV between 12.3 and 246.4 μM was added for 5 days before the final wash and Giemsa staining. The number of infected macrophages and the amount of amastigotes per infected macrophage were determined by manual counting. These numbers were used to establish the infection index (infected macrophages × amastigotes/infected macrophages). The percentage of parasite growth inhibition was calculated as [1 - (InfInd/InfInd0)] × 100, where InfInd0 and InfInd are infection indices of untreated infected cells and cells treated with different SSG concentrations, respectively. EC50s were calculated with GraphPad Prism using a sigmoidal dose-response model with variable slope.

**SbIII resistance selection.** For each strain, selection of resistant parasites was initiated in quadruplicates in 200 μl on a 96-well plate. Briefly, in each well, 10⁵ logarithmic-phase promastigotes were incubated for 7 days with SbIII at concentrations ranging from 0.4 to 382 μM following a 2-fold increase. This first step allowed the estimation of the highest concentration tolerated by the parasites at a given time point. The first concentration of the selection process was determined by the highest concentration at which at least 3 replicates were growing. Parasites were then transferred into 5 ml culture medium with the selected concentration of SbIII and maintained for 5 weeks before starting a new selection round. Selection rounds were performed successively with 6, 12, 24, 48, 96, 191, and 382 μM SbIII for BPK252 and 48, 96, 191, and 382 μM SbIII for BPK282 and BPK275. A rapid procedure for SbIII resistance selection (called SbIII flash selection) was also attempted. Wild-type parasites from different genetic groups described in reference 9 were cultured in parallel in triplicates with and without 382 μM SbIII (Table S1). Counting was performed every day for 5 weeks as described above. Parasites were subcultured at 5 × 10⁵ parasites/ml⁻¹ at day 7 when possible. When the parasite concentration was too low, cells were pelleted and resuspended in fresh medium at a lower SbIII concentration.

**DNA preparation for NGS and data analysis.** Parasites were pelleted after 5 weeks of culture with a given concentration of SbIII. DNA isolation was done using QIAamp DNA blood mini kit (Qiagen), and the DNA concentration was assessed with the Qubit DNA broad-range DNA quantification kit (Thermo Fisher). Sequencing libraries were prepared with the TrueSeq DNA library prep kit (Illumina) according to the manufacturer’s instructions. The libraries were quantified by qPCR using the KAPA Library quantification kit optimized for Roche LightCycler 480 (KAPA biosystems). The libraries were 2× 100-bp paired-end sequenced with the Illumina Hiseq 2000 platform by the Beijing Genomics Institute (BGI) according to standard protocols. Ploidy, single nucleotide polymorphisms (SNPs), and local copy number variations (CNVs) were determined as described by Imamura et al. in 2016 (9) using the BPK282v2 PacBio
reference genome accessible via ftp://ftp.sanger.ac.uk/pub/project/pathogens/Leishmania/donovani/LeDPK/AC201beta/. Sequences are available at GenBank Nucleotide Archive under accession no. PRJEB22849. Gene identity in the manuscript is given with LeDPK v2 as annotated elsewhere (19). However, LeDPK v1 gene identity, as used at GeneDB (http://www.genedb.org/Homepage), is also mentioned to facilitate comparisons with other studies. Biological and statistical significance of chromosome copy number (S value) comparison between samples was done according to two criteria as previously described (25). Briefly, the somy of a given chromosome in a population of cells is known to be a continuous variable with a multimodal distribution of S values because of aneuploidy mosaicism. Under this model, the S value of homologous chromosomes is thus considered to be significantly different if (i) S values differ from more than 0.5 with a somy distribution shift from one mode to another and (ii) the P value is $\leq 10^{-5}$.

**Metabolite extraction, LC-MS protocol, and data analysis.** For each analyzed sample, metabolites were extracted from four replicates of logarithmic-phase promastigotes (3 days of culture) using a 1:3:1 (vol/vol/vol) chloroform-methanol-water solvent as previously described (29). Samples were analyzed with an Orbitrap Exactive (Thermo Fisher Scientific) mass spectrometer coupled to a 2.1-mm ZIC-hydrophilic interaction liquid chromatography (HILIC) column (Sequent) at Glasgow Polynomials (University of Glasgow, Glasgow, Scotland) as previously described (30). All samples were analyzed in randomized order and in the same analytical batch. Several quality control samples, including solvent blanks, pooled samples, and authentic standard mixtures, were also measured in the same run to verify liquid chromatography-mass spectrometry (LC-MS) stability and allow the identification of contaminants (34). Data analysis was performed in R using the packages XCMS (35) and mzMatchR (36). The workflow was described previously (29). R was also used to generate heat maps and PCA plots based on all metabolites in the data set with measurements across all samples. For each metabolite, we computed log fold changes (FC): i.e., the log2 ratio of average metabolite abundance between two conditions for comparisons at the level of individual metabolites. The associated P value was obtained by performing a Student’s t test for each comparison, which was subsequently corrected with the Benjamini-Hochberg algorithm to limit the false-discovery rate (FDR) to 5%. Differences in the amounts of metabolites were considered significant if the log2 FC was $>1$ or $<-1$ and the corrected P value was $<0.05$.

**Single-gene overexpressers.** Each gene of the H and M loci was overexpressed in BPK026/0 cl5 after cloning in pLEXSY-hyg2 (Jena Bioscience) (Table S2). All genes except the MRPA gene were PCR amplified from BPK282 DNA using Phusion high-fidelity polymerase (NEB) with overhang primers (Table S2) for subcloning in pGEMT (Promega) before final cloning in pLEXSY-hyg2. MRPA was amplified using iProof high-fidelity DNA polymerase (Bio-Rad) with GC buffer plus 10% dimethyl sulfoxide (DMSO) and directly cloned in the pLEXSY-hyg2 using the In-Fusion HD cloning kit (Clontech). Every PCR-amplified sequence was sequenced at the VIB Genetic Service Facility (VIB Genetic Service Facility, Antwerp, Belgium). Promastigotes of BPK026/0 cl5 were then transfected using GenePulser Xcell (Bio-Rad) as described previously (37). Transfected parasites were selected with 50 µg·ml⁻¹ of hygromycin (Jena Bioscience) and then maintained with 100 µg·ml⁻¹.

**Cosmid isolation and transfection.** A cosmid bank of L. donovani MHOM/NP/03/BPK190 (ISC4) had already been prepared in the laboratory of Joachim Clos in Hamburg, Germany (38, 39). Cosmids carrying copies of the H locus (pcosTL-H-Locus) and the M locus (pcosTL-M-Locus) were isolated by colony lifting. Briefly, the cosmid bank was plated on LB agar-ampicillin plates, and a replica of each plate was taken on a nylon membrane (Amersham Biosciences, Inc.). Probes specific for LeDPK_360076800 (MPK1) and LeDPK_340035000 (SAT) (Biogazelle) using for normalization two genes previously shown to be very stable: SAT (LeDPK_340035000) (40) and LeDPK_240021200 (25). The expression of both normalization targets was confirmed by qBase+ as stable during the course of the experiment. Data were then rescaled to the BPK026 WT. An increase of 1.5-fold compared to the BPK026 WT was considered significant.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mSphere.00548-17.

**FIG S1,** TIF file, 0.5 MB.

**FIG S2,** TIF file, 0.7 MB.

**FIG S3,** TIF file, 2.9 MB.

**TABLE S1,** PDF file, 0.1 MB.
TABLE S2, PDF file, 0.1 MB.
DATA SET S1, XLS file, 0.1 MB.
DATA SET S2, XLS file, 0.1 MB.
DATA SET S3, XLS file, 0.5 MB.

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