Glyoxalase I Gene Deletion Mutants of Leishmania donovani Exhibit Reduced Methylglyoxal Detoxification

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Abstract

Background: Glyoxalase I is a metalloenzyme of the glyoxalase pathway that plays a central role in eliminating the toxic metabolite methylglyoxal. The protozoan parasite Leishmania donovani possesses a unique trypanothione dependent glyoxalase system.

Principal Findings: Analysis of the L. donovani GLOI sequence predicted a mitochondrial targeting sequence, suggesting that the enzyme is likely to be targeted to the mitochondria. In order to determine definitively the intracellular localization of GLOI in L. donovani, a full-length GLOI gene was fused to green fluorescent protein (GFP) gene to generate a chimeric construct. Confocal microscopy of L. donovani promastigotes carrying this chimeric construct and immunofluorescence microscopy using anti-GLOI antibodies demonstrated that GLOI is localized in the kinetoplast of the parasite apart from the cytosol. To study the physiological role of GLOI in Leishmania, we first created promastigote mutants heterozygous for GLOI by targeted gene replacement using either hygromycin or neomycin phosphotransferases as selectable markers. Heterozygous mutants of L. donovani display a slower growth rate, have lower glyoxalase I activity and have reduced ability to detoxify methylglyoxal in comparison to the wild-type parasites. Complementation of the heterozygous mutant with an episomal GLOI construct showed the restoration of heterozygous mutant phenotype nearly fully to that of the wild-type. Null mutants were obtained only after GLOI was expressed from an episome in heterozygous mutants.

Conclusions: We for the first time report localization of GLOI in L. donovani in the kinetoplast. To study the physiological role of GLOI in Leishmania, we have generated GLOI attenuated strains by targeted gene replacement and report that GLOI is likely to be an important gene since GLOI mutants in L. donovani showed altered phenotype. The present data supports that the GLOI plays an essential role in the survival of this pathogenic organism and that inhibition of the enzyme potentiates the toxicity of methylglyoxal.

Introduction

Leishmaniasis constitutes a wide spectrum of diseases ranging from the simple self-limiting cutaneous form to the debilitating visceral form, which is often fatal if left untreated. The protozoan parasite Leishmania donovani is the major causative agent of visceral leishmaniasis. The current challenges in its chemotherapy include widespread resistance to pentavalent antimony; absence of safe and cost-effective antileishmanial agents and relapses in HIV Leishmania co-infected patients. Thus an urgent need exists to look for newer and more effective drug targets to treat leishmaniasis.

The glyoxalase system of Leishmania is one such pathway that may be exploited successfully for the development of anti-parasitic drugs. The glyoxalase system is a ubiquitous thiol-dependent detoxification pathway [1]. This system comprises of two enzymes, glyoxalase I (GLOI) (lactoylglutathione lyase, EC 4.4.1.5) and glyoxalase II (GLOII) (hydroxyacylglutathione hydrolase, EC 3.1.2.6). Glyoxalase I catalyses the formation of S-D lactoyl glutathione from the hemithioacetal formed nonenzymatically from methylglyoxal and glutathione. Glyoxalase II converts S-D lactoyl glutathione to lactate and free glutathione [2]. Recent reports demonstrated a unique glyoxalase system in pathogenic kinetoplastids, as a consequence of this protozoan possessing an unusual thiol metabolism [3,4,5]. Glyoxalase I and glyoxalase II in kinetoplastids showed high level of specificity to trypanothione hemithioacetal and little activity with glutathione hemithioacetal [3,4,5,6]. The unique nature of this trypanothione-dependent glyoxalase pathway represents a novel chemotherapeutic target for the kinetoplastids. A major function of the glyoxalase pathway is believed to be detoxification of α-ketoaldehydes, especially methylglyoxal (MG). MG is a cytotoxic metabolite produced primarily as a by-product of glycolysis through nonenzymatic phosphate elimination from the glycolytic pathway intermediates, dihydroxyacetone phosphate and glyceraldehyde 3-phosphate [2]. The glyoxalase enzymes have been reported to have a regulatory role in cell division including cell proliferation, tissue regeneration, and malignancy besides detoxification of methylglyoxal [7]. We had earlier cloned and characterized the genes encoding

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glyoxalase I (GLOI) (AY739896) and glyoxalase II (GLOII) (AY851655) from *Leishmania donovani* [5,6]. Glyoxalase I is a single copy gene and is placed at a single chromosomal band of ~2.2 Mb. Multiple sequence alignment and homology modeling clearly showed that there is a crucial difference in the active site of human and *Leishmania* glyoxalase I enzymes suggesting that the glyoxalase I may be a potential target for drug design [5].

In the present study we report that *L. donovani* promastigotes carrying a GLOI-green fluorescent protein (GLOI-GFP) chimeric construct display localization of GLOI in the kinetoplast of the parasite apart from the cytosol. To study the physiological role of GLOI in *Leishmania*, we have generated GLOI attenuated strains by targeted gene replacement. Attempts to disrupt both alleles of GLOI in *L. donovani* promastigotes resulted in change in levels of cell ploidy. These GLOI mutants were phenotypically characterized and showed reduced growth, lower GLOI enzymatic activity and had reduced ability to detoxify methylglyoxal in comparison to the wild-type parasites. Null mutants of GLOI were obtained after rescuing the heterozygous mutants with an episomal copy of GLOI. This data suggests that the GLOI is essential to *L. donovani*.

### Results

**Subcellular localization of Glyoxalase I in *L. donovani***

Amino acid sequence analysis of the LdGLOI protein in-silico using pSORT [8] and MitoProt II [9] web-based programs predicted that GLOI is localized to the mitochondria with a *P* = 0.8780 probability. However, no cleavage site was predicted for the mitochondrial targeting sequence using Target P web based program [10]. To determine the localization of GLOI in intracellular organelles, we investigated the localization of a GFP-tagged version of GLOI in *L. donovani* and compared with the wild type parasites, transfected with the vector alone (pGEM-7zfNeo-GFP) (Fig. 1A, panel 5 and 6). Confocal microscopy showed that the GLOI-GFP fusion protein was localized both in the cytosol and the *Leishmania* kinetoplast (Fig. 1A), as seen by the clear co-localization of the GLOI-GFP with the fluorescence associated to the MitoTracker Red CMX Ros that reveals the position of mitochondria within the promastigotes (Fig. 1A, panel 1, 3 and 4). Immuno-localization of GLOI using anti-GLOI antibodies and Alexa 546 labeled anti mouse IgG as secondary antibody in the *L.donovani* expressing GLOI as GFP translational fusion

![Figure 1. Localization of GLOI in *L. donovani*. A: Panel 1: confocal microscopy of wild type *L. donovani* transfected with pGEM7zf-neo5-GFP-GLOI expressing GLOI as a GFP translational fusion protein and stained with MitoTracker-Red CMX Ros dye, Panel 2: phase contrast image, Panel 3: wild type *L. donovani* expressing GLOI as a GFP translational fusion protein, Panel 4: merged micrograph, Panel 5: wild type *L. donovani* transfected with pGEM7zf-neo5-GFP vector alone expressing GFP, Panel 6: phase contrast image B: Panel 1: Immunofluorescence analysis by confocal micrograph of wild type *L. donovani* parasite transfected with pGEM7zf-neo5-GFP-GLOI using anti–GLOI antibody, Panel 2: phase contrast image, Panel 3: wild type *L. donovani* expressing GLOI as a GFP translational fusion protein, 4: merged micrograph C: Panel 1: Immunofluorescence analysis by confocal micrograph of wild type *L. donovani* parasite transfected with pGEM7zf-neo5-GFP-GLOI using anti–GLOI antibody, Panel 2: wild type *L. donovani* expressing GLOI as a GFP translational fusion protein, Panel 3: stained with DAPI, Panel 4: phase contrast image, Panel 5: merged micrograph. D: Western blotting using anti-GLOI antibody. Cytosolic and mitochondrial fractions were separated from late log phase promastigotes (1 × 10⁸) expressing GLOI as a GFP translational fusion protein. Lane 1: recombinant GLOI, Lane 2: cytosolic fraction, Lane 3: mitochondrial fraction. The blot was also probed with mitochondrial specific anti-F1 ATP synthase α-subunit. Results are representative data from three separate experiments. doi:10.1371/journal.pone.0006805.g001

**Glyoxalase I as Drug Target**

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protein, further confirmed the presence of GLOI in the kinetoplast and the cytosol and not in the nucleus (Fig. 1B panel 1, 3 and 4). Immunolocalization of GLOI using anti-GLOI antibodies and Alexa 546 labeled anti mouse IgG as secondary antibody was also done and counter stained with DAPI to confirm the position of the nucleus and the kinetoplast with in the parasites (Fig. 1C, panel 1, 2, 3 and 5). Western blot analysis with both the cytosolic and mitochondrial fraction (100 µg protein lysate) of the GLOI-GFP transfected parasites showed a band of ~44.5 kDa that corresponded to the size of the fusion protein and a band of 16 kDa that corresponded to the native L.donovani GLOI protein. These experiments confirmed the presence of GLOI with in the kinetoplast and the cytosol of the parasites (Fig. 1D). The mitochondrial fraction was further confirmed by reprobing the blot with mitochondrial specific anti-F1 ATP synthase α-subunit (Fig. 1D).

Construction and Molecular characterization of GLOI mutants

From our earlier studies it is known that GLOI is a single copy gene in L. donovani and is placed at a single chromosomal band of ~2.2 Mb [5]. Since Leishmania is considered to be diploid and no sexual crosses have been achieved, homologous recombination strategy has been used in an attempt to generate transgenic parasites lacking both alleles of GLOI. The wild type parasites were transfected individually with the hyg and neo replacement cassettes to create the GLOI heterozygotes; Ld GLOI :: HYG (+/+h) and Ld GLOI :: NEO (+/+n) respectively (Fig. 2). The Hyg and Neo transfectants were selected against hygromycin (200 µg/ml) and G-418 (40 µg/ml) respectively. The representative clone from the Ld GLOI :: HYG (+/+h) and Ld GLOI :: NEO (+/+n) single knock out mutants were screened for the correct integration of the hyg and neo replacement cassette respectively. The SalI digested genomic DNA from the wild-type L. donovani and representative L. donovani heterozygous clones Ld GLOI :: HYG (+/+h) and LdGLOI :: NEO (+/+n) were subjected to Southern blot analysis and results are presented in Fig. 3. The integrations of the selectable markers were confirmed by hybridization with hyg and neo specific probes. The hyg specific probe (1026-bp) resulted in no hybridizing bands with SalI-digested genomic DNA from wild type parasites and Ld GLOI::NEO (+/+n) clones, while a single band of ~6.8 kb,

![Figure 2](https://www.plosone.org/figure/2)

**Figure 2. Schematic representation of the wild type and the hyg, neo and phleo targeted Ld GLOI locus.** SalI restriction sites were used to analyze the recombination events. The restriction fragments obtained after SalI digestion are indicated by double headed arrows along with their respective size. Thick arrows indicate the ORF of genes as indicated. Bold arrows indicate the 5' and 3' flanking regions of GLOI ORF. Solid bars indicate the probes used for Southern blot analysis. Primers P 1 and P 2 were used to check the correct integration of hyg cassette and primers P 1 and P 3 were used to check the correct integration of neo cassette. Restriction sites (H- HindIII, B- BamHI, Bg - BglII, SalI) used for cloning 5' and 3' flank are also indicated. doi:10.1371/journal.pone.0006805.g002
corresponding to the hyg based replacement fragment was detected in L. donovani heterozygous clones, Ld GLOI :: HYG (+/h) (Fig. 3A). The neo specific probe (800-bp) resulted in no hybridizing band with SalI digested genomic DNA from the wild-type (+/+) parasites and representative Ld GLOI :: HYG (+/h) clones but a single band of ~6.7 kb was detected in the Ld GLOI :: NEO clone (+/n) (Fig. 3B). Southern blot analysis of SalI-digested genomic DNA from wild-type parasites (+/+), L. donovani heterozygotes: Ld GLOI :: HYG (+/h), Ld GLOI :: NEO (+/h/n), a double transfectant mutant clone Ld GLOI :: HYG (+/h/n) and a GLOI complementation mutant (+/h - GLOI*) was performed and blots were hybridized with the GLOI ORF probe (426 bp). An expected 2.2 kb band was recognized in the DNA from wild-type L. donovani (+/+) and L. donovani heterozygous mutants Ld GLOI :: HYG clone (+/h), Ld GLOI::NEO clone (+/n) and GLOI complementation mutant (+/h - GLOI*). An additional and expected band of 6.7 Kbp was observed in the (+/h)-GLOI* transfectant that corresponds to the presence of an episomal copy of GLOI in the Ld GLOI :: HYG (+/h) background. However, the double transfectant mutant clone, Ld GLOI :: HYG :: NEO (+/h/n) showed the presence of a GLOI specific band of 2.2 kb (Fig. 3C). A total of ten double transfectant mutant (+/h/n) clones were screened and a 2.2 kb specific GLOI band was present in all the clones (representative data from one clone is shown). Although GLOI gene disruption by neo and hyg took place, as clearly shown from the hybridization studies with the hyg (Fig. 3A) and neo probes (Fig. 3B), one GLOI allele remained intact since the 2.2 kb genomic SalI fragment was still present in the double targeted mutants (Fig. 3C, lane: +/h/n). The gene replacement of four of the (+/n) clones was further evaluated by polymerase chain reaction (PCR).

PCR based approach was also used to analyze for the correct integration of the hyg and neo based replacement constructs. Integration specific primers P1 and P2 as mentioned in the material and methods and shown in Fig. 2 were used to determine hyg specific integration (Fig. 3E), whereas P1 and P3 was used to verify neo specific integration (Fig. 3F). Southern blotting of the PCR products of the wild type (+/+), Ld GLOI :: HYG clone (+/h),
**Ld GLOI**::**NEO** clone (+/n) and double transfectant mutant clone **Ld GLOI::HYG::NEO** (+/h/n) using **hyg** (Fig. 3E, panel b) and **neo** (Fig. 3F, panel b) specific probes further confirmed the expected integration. These observations confirmed the generation of single knockouts of **GLOI** by both **hyg** and **neo** based replacement cassettes. PCR studies further confirmed that in the double transfectant mutant clone **Ld GLOI::HYG::NEO** (+/h/n) disruption of **GLOI** gene by **neo** (Fig. 3F, panel b) and **hyg** (Fig. 3E, panel b) took place.

Since the double transfectant mutant clones **Ld GLOI::HYG::-NEO** (+/h/n) were found to contain 2.2-kb wild type **GLOI** allele as well as **hyg** and **neo** integrations, DNA content based analysis was performed to check the ploidy levels of the cells. This was determined by fluorescence activated cell sorting (FACS) analysis of fixed cells stained with propidium iodide (Fig. 4). In the wild type cells (+/+), two peaks corresponding to 2N (G1) and 4N (G2) DNA content were observed as expected (Fig. 4 panel A). Similar pattern was observed with the heterozygote mutants (+/h) and (+/n) and **GLOI** complementation mutant (+/h-**GLOI**+) (Fig. 4, panels B, C and D) thereby showing that these clones contain normal diploid DNA content similar to the wild type cells. However, analysis of the double transfected mutant clone (+/h/n) showed peaks that corresponded to 3N and ~6N DNA content (Fig. 4 panel E) (representative data from one of the clones is shown). This suggests a difference in the ploidy level in the cell population of the double transfectants when compared to the wild type cells. The G1 peak of the double transfected mutant clone indicates that these cells are aneuploid (Fig. 4, panel E).

In order to create null mutant of **GLOI**, a further attempt was made to express the gene from an episome prior to disrupting the second **GLOI** allele. The heterozygous **GLOI** mutant with an episomal copy of **GLOI** (+/h-**GLOI**+) was transfected with the linearized pX63-PHLEO-A**GLOI** construct. The transfectants were selected against hygromycin (200 μg/ml), G418 (40 μg/ml) and phleomycin (25 μg/ml). The null mutants thus obtained were designated as (−/−**GLOI**+). The **Sal**I digested genomic DNA

**Figure 4. DNA content analysis.** Wild type and **GLOI** mutant **L. donovani** cell lines were analysed for DNA content using flow cytometry. A: Wild type (+/+); B: heterozygote **LdGLOI**:**HYG** clone (+/h); C: heterozygote **LdGLOI**:**NEO** clone (+/n); D: **GLOI** complementation mutant (+/h-**GLOI**+); E: **L. donovani** double targeted transfectant, **LdGLOI**:**HYG**:**NEO** clone (+/h/n) were fixed and stained with propidium iodide and analysed using a Beckton Dickinson flow cytometer (FACScalibur). Representative peaks of G1 and G2 phase are shown as predicted by the CELL QUEST software. Results are representative data from three separate experiments.

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from nine clones was subjected to Southern blotting and the blot was hybridized with GLOI (426-bp) (Fig. 3D) or phiio (375-bp) specific probe (Fig. 3G). A representative data from two (−/−GLOI) clones is shown in Fig. 3D and 3G. The 2.2 kb GLOI fragment was present in the wild type L. donovani (+/+), heterozygous mutants, Ld GLOI :: HYG clone (+/h), LdGLOI::NEO clone (+/n) and GLOI complementation mutant (+/h – GLOI)
. The two mutant clones (−/−GLOI) lacked 2.2-kb wild type GLOI fragment. These two clones were found to contain an intensely hybridizing fragment of 6.7-kb which corresponded to the presence of an episomal copy of GLOI in the Ld GLOI :: HYG (+/h) background. These clones were also checked for the correct integration of the phiio replacement cassette by Southern blot analysis of Sall digested genomic DNA followed by hybridization with the phiio probe. Fig. 3G shows that both the clones (−/−GLOI) contain 6.1-kb fragment indicative of the correct integration of the phiio cassette.

A total of at least two clones of each of the mutant lines were picked and screened for phenotypic characterization. Representative data from one clone from each mutant line is presented below.

GLOI heterozygous mutants are growth defective

In order to verify phenotypic alteration in these gene replacement mutants, the growth rate of each of the cell lines was determined for a period of 120 h at an interval of 24 h. Fig. 5 shows the growth of the wild-type L. donovani, Ld GLOI :: HYG (+/h), Ld GLOI :: NEO (+/n) and complementation clone (+/h)-GLOI and double targeted transfectant clone Ld GLOI :: HYG :: NEO (+/h/n). The promastigotes of the GLOI heterozygous mutants (+/h) and (+/n) showed a slower growth rate when compared to that of the wild-type cells (+/+). The GLOI heterozygous mutants (+/h) clone when complemented with an episomal copy of GLOI showed an increase in the growth rate as compared to the heterozygous mutant, thereby revealing that the presence of GLOI as an episomal copy facilitates in overcoming the defect in the growth that probably has arisen because of the deletion of one of the wild type GLOI allele. Double targeted transfectant clone (+/h/n) had growth pattern similar to that of the heterozygous GLOI mutant strains.

GLOI heterozygous mutants exhibit reduced methylglyoxal (MG) detoxification

The effect of methylglyoxal concentration on the growth rate of the L. donovani wild type and the mutant parasites is shown in Fig. 5. The effective concentration was 0.41±0.04 mM towards MG. This date indicates that complementation of GLOI in heterozygous mutant clone Ld GLOI :: HYG (+/h) decreases the sensitivity towards methylglyoxal.

Reduction in glyoxalase I activity in GLOI heterozygous mutant

The specific activity of GLOI in heterozygous mutants, Ld GLOI :: HYG (+/h) and Ld GLOI :: NEO (+/n), was 173±31 nmol min−1 mg−1 protein and 116.5±33 nmol min−1 mg−1 protein respectively. The specific activity in the double targeted transfectant clone (+/h/n) was 109.5±13 nmol min−1 mg−1 protein. However, a higher specific activity of GLOI in the wild-type L. donovani (486±19.8 nmol min−1 mg−1 protein) was observed when compared to the respective mutants. Thus, Glyoxalase I activity in L. donovani mutants showed an approximately two fold reduction as compared to the wild-type L. donovani. The heterozygous mutant Ld GLOI :: HYG (+/h) when complemented with an episomal copy of GLOI had a specific activity of 345±28 nmol min−1 mg−1 protein (Table 2).

The rescued null mutant clones (−/−GLOI) had normal diploid content. The growth pattern, IC50 towards MG and specific activity of GLOI null mutant clones (−/−GLOI) was similar to that of the GLOI complementation mutant (+/h – GLOI) (data not shown).

Discussion

The glyoxalase system, consisting of the enzymes glyoxalase I (GLOI) and glyoxalase II (GLOII), is an integral component of cellular metabolism in mammalian systems. Although the glyoxalase pathway was reported as early as 1913 [11,12] and is

| Strain | IC50 ± S.D. (mM) |
|--------|------------------|
| Wild type (+/+) | 1.0±0.14 |
| Ld GLOI :: hyg (+h) | 0.45±0.03* |
| Ld GLOI :: neo (+/n) | 0.3±0.02* |
| Ld GLOI :: hyg neo (+/n) | 0.41±0.04* |
| Ld GLOI :: hyg GLO I (+/h – GLOI) | 0.8±0.07 |

IC50 represents the concentration of methylglyoxal, which inhibited the growth of the wild type and the mutant parasites by 50%. Each value is a mean±S.D. of three independent experiments.

*Significantly different from the wild type (P<0.001).

Table 1. Effect of methylglyoxal on the cell viability of the Wild type and the mutant parasites.
widely distributed in living organisms, its full biological function has never been elucidated. A major function of the glyoxalase pathway is believed to be detoxification of α-ketoaldehydes, especially methylglyoxal (MG). MG is a cytotoxic metabolite produced primarily as a by-product of glycolysis [13]. Another role of GLOI and methylglyoxal is their involvement in the regulation of cellular growth [14]. In tumor tissues high activities of glyoxalase I have also been reported [15]. Inhibitors of glyoxalase I have been reported to be selectively toxic to proliferating cells, this could be due to increased accumulation of methylglyoxal that could lead to inhibition of DNA synthesis [16,17]. Glyoxalase I inhibitors have been reported to have antimalarial [18] and antitrypanosomal activities [19]. The glyoxalase system of the pathogenic kinetoplastids has been recently reported to be unique, as a consequence of these protozoa possessing an unusual thiol metabolism. In these organisms, instead of glutathione, the major low molecular mass thiol is trypanothione [N,N 8-bis (glutathionyl) spermidine] [4,5]. It has been recently reported that the GLOI system in L. major and L. donovani uses trypanothione as the substitute for glutathione [4,5]. The metal cofactor is zinc in eukaryotes and nickel in E. coli [20,21] and L. major [4]. Thus both the substrate and cofactor of Leishmania glyoxalase are different from the mammalian glyoxalases. The difference in cofactor dependence is reflected in differences between the active sites of the human and Leishmania enzymes, suggesting that the latter may be a target for antimicrobial therapy [22,23].

Localization studies using L. donovani GLOI–GFP translational fusion shows that GLO I is localized not only in the cytosol of the parasites but also in the kinetoplast. Analysis of L. donovani GLOI sequences with MitoProt II [9] suggested targeting of proteins to the mitochondria. The GLO I enzyme is likely to be exported to the mitochondria with a P=0.8780. Analysis of the L. donovani GLOI sequences with PSORT also suggested it to be a mitochondrial protein. This observation is supported from the sequence analysis of T. cruzi and L. major GLOI with MitoProt II having a P=0.909 and 0.8704 respectively [24].

The biological function of a gene in an organism can be assessed by generating a knockout line and then characterizing the phenotype of the mutant. The functions of several genes of Leishmania have been established using gene targeting procedures [25,26,27,28]. Our earlier studies have shown that the GLOI is a single copy gene and the expression is highest in late log phase promastigotes [5]. Homologous recombination strategy has been used to obtain single knock out mutants Ld GloI :: HYG (+/h) and Ld GLOI :: NEO mutants (+/n). Both these mutants exhibited slow growth rate and decrease in the specific activity of the glyoxalase I enzyme when compared to the wild-type. The decrease in enzymatic activity probably causes accumulation of MG derived adducts and induces oxidative stress in the mutants. Loss of one GLOI allele by gene disruption was compensated by genetic complementation with an episomal copy of GLOI. Our attempts to generate GLOI null mutants by step wise gene replacement failed despite the successful disruption of the two alleles by gene targeting. Earlier reports show that in L. major, generation of dihydrofolate reductase-thymidylate synthase null mutant [29] resulted in an increase in the chromosome number in order to keep one dfr-ts allele intact. Similarly disruption of the trypanothione reductase (TR) gene of Leishmania led to partial trisomy for the TR locus [30]. It has been proposed that such alteration in the genome could be an indicator of the essential nature of the gene. In order to address this problem in the present study, the DNA content of the mutant lines was determined by FACS analysis and compared to that of the wild type. A normal diploid DNA content was observed in the wild type, heterozygous mutants with hyg replacement cassette (+/h), heterozygous mutant with neo replacement cassette (+/n), and GLOI complementation mutant (+/h/-GLOI +). However, analysis of the DNA content of the double targeted transfectant (+/ h/n) indicated that the second round of gene replacement resulted in aneuyploidy. This change in ploidy has occurred during the introduction of the second replacement construct. The mechanism by which such changes in ploidy occur during gene replacement is not known, but this phenomenon is an indication that the gene being targeted for replacement is essential.

In order to test the hypothesis that the GLOI is an essential gene, an attempt was made to express the gene from an episome by previous transfection with a plasmid carrying the target gene for episomal expression and genetic complementation before the second round of gene replacement. We were successful in replacing both GLOI loci with two selection markers only in the presence of an episomal copy of GLOI in the heterozygous mutant. This shows that the Leishmania donovani GLOI gene is essential for the survival.

Difference in response to MG of the wild-type and the single allele knockout mutant further shows that mutants probably have a significantly reduced rate of metabolism of MG compared to the wild-type parasites. The heterozygous mutant Ld GloI :: HYG (+/h) when complemented with an episomal copy of GLOI was able to overcome the effect caused by the deletion of one of the allele of GLOI. It has been implicated that glyoxalase I serves as an indicator of cell growth status in numerous cells, as its activity is low in resting cells and high during active cell growth and division [31,32,7]. The balance between methlyglyoxal (the naturally occurring substrate of glyoxalase I) and glyoxalase I play a decisive role in the regulation of cell growth [33]. In the present study, it is possible that the single knock out mutants were unable to detoxify the cytotoxic effects of endogenous methlyglyoxal and its accumulation induced cell growth retardation [34]. Similar observations have been made in the GLOI null mutants of Escherichia coli [35]. Inactivation of the GLOI gene sensitizes cells to MG. The phenotypic characteristics of the null mutants in the presence of an episomal copy of GLOI (+/-GLOI +) when studied were found to be similar to that of GLOI complementation mutants (data not shown). This data suggests that glyoxalase I may play an essential role in survival of this pathogenic organism and the major therapeutic perspective emerging from this study is that Leishmania GLOI may be further validated as an interesting target to develop specific chemotherapeutic inhibitors against the parasite.

Table 2. Comparison of the Glyoxalase I activity of the Wild type and the mutant strains of glyoxalase I.

| Strain                     | Specific activity ± S.D (nmoles/min/mg) |
|---------------------------|-----------------------------------------|
| Wild type (+/+)           | 486±19.8                                 |
| Ld GLOI :: hyg (+/h)      | 173±31*                                  |
| Ld GLOI :: neo (+/n)      | 116.5±33*                                |
| Ld GloI hyg :: neo (+/h/n)| 109.5±13*                                |
| Ld GLOI :: hyg GLOI +    | 345±28                                   |

The Glyoxalase I activity of the wild type and mutant parasites was determined. Results are mean±S.D. values of three independent experiments. *Significantly different from the wild type (P<0.001).

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Materials and Methods

Materials

Trypanothione disulphide was purchased from Bachem (Switzerland). [\(\alpha\)-\(P\)]-dCTP (3000 Ci/mmol) was acquired from Amersham Biosciences, U.K. All restriction enzymes and DNA modifying enzymes were obtained from MBI Fermentas (Germany). The pX63-NEO and pX63-HYG, vectors that encompass the neomycin phosphotransferase (neo) and hygromycin phosphotransferase genes (hyg) respectively were kindly provided by Dr. Stephen M. Beverley (Washington University, St. Louis, MO). Construct pGEM-7zf(+)Neo and pGEM-7zf(+)Hyg containing a 426-bp GLOI specific probe. Hygromycin B, Phleomycin and Geneticin (G418) were obtained from Sigma Aldrich Corp. (St. Louis, MO). Alexa-546 labeled IgG was obtained from Molecular Probes, Eugene, OR. The other materials used in this study were of the highest purity and were commercially available.

Cell Culture

*L. donovani* promastigotes were grown at 22°C in M199 medium (Sigma St. Louis, MO) supplemented with 100 units/ml penicillin (Sigma, St. Louis, MO), 100 μg/ml streptomycin (Sigma, St. Louis, MO) and 10% fetal bovine serum (FBS, Hyclone, U.K.). All mutants of GLOI gene used in this study were derived from the wild-type *L. donovani* strain AG83 (MHOM/IN/1983/AG83) designated GLOI (+/+) for the purpose of genetic manipulations described here where GLOI denoted the Glyoxalase I locus and + specifies the wild-type allele. Promastigotes of mutant parasites containing functional copies of the neomycin or phleomycin resistant gene or hygromycin phosphotransferase gene were routinely maintained in the same medium supplemented with either 40 μg/ml Geneticin (G418) or 25 μg/ml phleomycin or 200 μg/ml hygromycin respectively. For characterizing the mutant parasites phenotypically, cells were subcultured without the selection marker prior to the experiments.

Growth studies

Growth rate experiments were conducted by inoculating stationary phase parasites at a density of 1 × 10⁷ cells/ml in modified M199 medium with 10% FBS in 25 cm² flasks without respective selection drug and cultured at 22°C. Growth rates of each of the cultures were determined at 24 h intervals by Neubauer haemocytometer. Growth studies with each individual cell line were done at least three times and similar results were obtained consistently.

MTT assay

25 μl of 1 × 10⁶ parasites/ml of mid log phase promastigotes were cultured in 96 well tissue culture plates and incubated with 25 μl of varying concentrations of methylglyoxal at 22°C. After 72 h, 20 μl of 5 mg/ml MTT [3-(4, 5-dimethylthiazol-2-yl)-2, 5-diphenyltetrazolium bromide] (Sigma) dissolved in PBS (phosphate buffered saline with 1% of glucose, pH, 7.4) was added. Plates were incubated for 6 h at 37°C. Reaction was stopped by the addition of 50 μl of 50% isopropanol and 10% SDS by gentle shaking at 37°C for 30 min to 1 h. O.D. was measured spectrophotometrically at 570 nm [36]. Results are representative of three independent experiments.

DNA extraction, Southern Blotting and Hybridization

Genomic DNA was extracted from *Leishmania* using a standard method [37]. 5 μg of DNA was digested with appropriate restriction enzymes, separated by 0.5% agarose gel electrophoresis, stained with ethidium bromide, transferred onto nylon membranes (Amersham Pharmacia Biotech) and subjected to Southern blot analysis. Blots were hybridized with denaturated [\(\alpha\)-\(P\)]-dCTP labelled DNA probe at 10⁷ c.p.m/ml, which was labelled by random priming (NEB Blot Kit, New England Biolabs, Inc.) Membranes were washed, air dried and exposed to an imaging plate. The image was developed by Phosphoimager (Fuji FLA – 5000) using ImageQuant software (Amersham Biosciences).

Construction of GLOI-GFP fusion construct (pGEM-7zfNeo-GLOI-GFP)

The GLOI gene was amplified from the genomic DNA of *L. donovani*, using the primers 5’-CCCAAGCTTATGGCCGTCGCTGGATGATG-3’ and 5’-CCCAAGCTTGGCAGTCCGGCCTGAGTCCTT-3’ containing a flanking HindIII site (Underlined). The conditions for PCR were as follows: 94°C for 10 min, then 35 cycles of 94°C for 45 sec, 57°C for 45 sec and 72°C for 1 min. Final extension was carried out for 10 min at 72°C. The GLOI PCR product was digested with HindIII and cloned in to the vector pGEM-7zfNeo (containing neomycin phosphotransferase as the selection marker) with GFP as the carboxy terminal of the GLOI gene product. The clone was checked for correct orientation of the insert by restriction enzyme digestion and confirmed by sequencing. 20 μg of the plasmid, pGEM-7zfNeo-GLOI-GFP were transfected into the wild type *L. donovani*. Methods of electroporation and plating of *L. donovani* have been described previously [37]. The transfected cells were maintained in 40 μg/ml of G418. These transfected parasites were used for localization studies. A control transfection with vector alone, pGEM-7zfNeo-GFP was transfected in to the wild type *L. donovani* parasites for comparison.

Localization of GLOI

To detect the site of localization of GLOI in *L. donovani*, pGEM-7zfNeo-GLOI-GFP transfected promastigotes were used. Fluorescent imaging of the stabilized culture was performed using the confocal laser scanning microscope (Zeiss LSM 510 META) equipped with a 63× objective, at an excitation wavelength of 488 nm. Briefly 10⁷ promastigotes/ml were pelleted and stained with Mitotracker Red –CMX-Ros (1 nM) (Molecular Probes, Eugene, OR) in a fresh media for 10–15 min to locate the kinetoplast with in the cell. The cells were then washed with phosphate-buffered saline (PBS) containing 1% fetal bovine serum (FBS) and resuspended in the same PBS solution with identical final cell concentration. The promastigotes were then immobilized on poly (L) lysine coated cover-slips. The cover-slips were incubated for 1 h at 37°C to allow parasite adherence. The cells were then treated with methanol/acetone (1:1) for 15 min at −20°C followed by washing with PBS for 5 min. For observing the fluorescence of the parasites stained with Mitotracker Red – CMX-Ros the parasites were visualized under confocal laser scanning microscope at an excitation wavelength of 568 nm.

Immunofluorescence Microscopy

10⁷ promastigotes/ml of the were pelleted and washed twice with phosphate-buffered saline (PBS) containing 1% fetal bovine
serum (FBS) and resuspended in the same PBS solution with identical final cell concentration. The parasites were then immobilized and fixed as mentioned above and were permeabilized with PBS containing 0.5% Triton X-100 for 5 min, followed by washing with PBS. Blocking was done using PBS with 1% BSA. The cells were then incubated in a humidity chamber for 1 h at room temperature with mouse polyclonal anti-GLOI antibody (1:100 dilution). After washing with PBS, cells were incubated with secondary anti-mouse Alexa-546 labeled IgG (Molecular Probes, Eugene, OR) diluted 1:2000 for 1 h. The cells were washed with PBS and incubated with DAPI (0.1 μg/ml) for 15 min at room temperature. The cover-slips were mounted on the slides and the parasites were visualized under the confocal laser scanning microscope at an excitation wavelength of 405 nm for immunofluorescence. The parasites stained with DAPI were observed at an excitation wavelength of 405 nm.

Cloning procedures and molecular constructs for the replacement of GLOI alleles by homologous replacement

Plasmid constructs pX63-HYG, pX63-NEO and pX63-PHLEO were used for the targeted replacement of GLOI allele in the wild-type L. donovani strain AG83. To construct drug resistance cassettes to replace GLOI alleles, 5′-5′ flanking region and 1011-bp of 3′flanking regions of GLOI genes were sub-cloned into the appropriate sites of the pX63-NEO, pX63-HYG and pX63-PHLEO plasmids. The 5′-5′ flanking region was amplified from the GLOI genomic DNA by PCR using a sense primer 5′GCCAGGATCTACGGAATAGCGCAGAC-3′ and an antisense primer 5′CCCGCTCGAAGAGGGAGGCAGA-3′, containing HindIII and SalI sites (underlined) corresponding to a region ~217 nucleotides upstream from GLOI ORF. The 1011-bp 3′ flanking region was also amplified from the GLOI genomic DNA by PCR using a sense primer, 5′CCCGATCTCCCGTCTGCGTCGCTGGAC-3′ and the antisense primer, 5′GAAGATCTGGACGATTCCAGTTCTC-3′ containing BsmII and BglII sites (underlined).

The 3′ flank PCR product (~1011-bp) was then sub-cloned into the BsmII- BglII site of pX63-NEO, pX63-HYG and pX63-PHLEO. The resulting plasmids were termed pX63-NEO-3′F and pX63-HYG-3′F, pX63-PHLEO-3′F respectively. The 5′ flank PCR product (~542-bp) was then sub-cloned into the HindIII site of pX63-NEO-3′F, pX63-HYG-3′F and pX63-PHLEO-3′F to yield the three replacement constructs, pX63-NEO-ΔGLOI, pX63-HYG-ΔGLOI and pX63-PHLEO-ΔGLOI respectively. The correct orientation of 5′ flank region and 3′ flank region within the three drug resistance cassettes was confirmed by automated sequencing and restriction mapping.

In order to generate heterozygous mutants of GLOI, five micrograms of the linearized DNA of the targeting constructs pX63-NEO-ΔGLOI or pX63-HYG-ΔGLOI (digested with HindIII and BglII) was transfected into mid log phase L. donovani Ag83 wild-type promastigotes by using standard electroporation conditions for transfection as reported earlier [37]. Transfection with linearised pX63-NEO-ΔGLOI or pX63-HYG-ΔGLOI into the wild type L. donovani was done to create the heterozygote mutants, L. donovani GLOI::NEO (+/n) and L. donovani GLOI::HYG (+/h) respectively. Transfected L. donovani strains were maintained in liquid medium for at least 24 h before plating on a drug-containing semi-solid medium. After selection and stabilization of the liquid culture, mutant parasites were spread on the plates containing 1% Bacto-Agar (DIFCO) in M199 media (with 10% FBS) with either G418 (40 μg/ml) or hygromycin (200 μg/ml) as the case may be, to select for single colonies. The genotype of the L. donovani GLOI::NEO (+/n) and L. donovani GLOI::HYG (+/h) heterozygotes was confirmed by Southern blot analysis. A heterozygous clone of L. donovani GLOI::HYG (+/h) was then subjected to a second round of transfection with linearised pX63-NEO-ΔGLOI to obtain null line ΔGLOI. These double targeted transfecants, L. donovani GLOI::HYG::NEO (+/h/n) were selected on solid plates containing 1% Bacto-Agar (DIFCO) in M199 media (with 10% FBS) with both G418 (40 μg/ml) and hygromycin (200 μg/ml). A total of 10 clones of double targeted transfected line (ΔGLOI/+), were picked and screened by Southern blotting using GLOI ORF as a probe.

Selected clones from L. donovani GLOI::HYG::NEO (+/h/n) mutants were also confirmed for the presence of the hygromycin integration cassette using the hyg specific internal forward primer, P2, 5′-GACTGTCGACGTTACACA-3′ corresponding to a region 765-bp downstream of the hyg start codon and a reverse primer, P1, 5′-GCACGGATGCAACCAGAC-3′ corresponding to a region 138-bp downstream from the 3′flanking region of GLOI ORF (Fig. 2). Selected clones from L. donovani GLOI::NEO (+/n) mutants were checked for the presence of the neomycin integration cassette using the neo specific internal forward primer, P3, 5′-GGACCGGCTATCAGGACAT-3′ corresponding to a region 131-bp downstream of the neo start codon and a reverse primer, P1, 5′-GCACGGATGCAACCAGAC-3′ corresponding to a region 138-bp downstream from the 3′flanking region of GLOI ORF (Fig. 2). The gene replacement of four of the double transfectant clones L. donovani GLOI::HYG::NEO (+/h/n) was further evaluated by polymerase chain reaction using primers P1 and P3 and also P2 and P1 (Fig. 2).

Genetic complementation of the GLOI heterozygote mutant

GLOI gene was amplified using the primers 5′CCCAAAGCT- TATGGCCGTCTGCGTCGATG-3′ and 5′CCCAAATGCAGGGCAGATC CCTGCTGCTGTCCAGTTT-3′ using a flanking HindIII site (Underlined). The conditions for PCR were as follows: 94°C for 10 min, then 35 cycles of 94°C for 45 sec, 57°C for 45 sec and 72°C for 1 min. Final extension was carried out for 10 min at 72°C. The GLOI PCR product was digested with HindIII and cloned in the vector pGEM-7zf-LdGLOI (containing neomycin phosphotransferase as the selection marker) to create pGEM-7zf-LdGLOI containing the neomycin phosphotransferase gene. The representative heterozygous mutant, L. donovani GLOI::HYG (+/h) were transfected with pGEM-7zf-LdGLOI construct and transfected parasites were then selected in G418 (40 μg/ml) and hygromycin (200 μg/ml) in M199 media containing 10% FBS. Transfectant lines harboring epimorphic encompassing the wild-type and single deletion construct (linearized pX63-HYG-ΔGLOI) were designated as+/h GLOI.

Creating double mutants after episomal complementation of the GLOI heterozygote mutant

The transfecant cell line harboring the epimorphic encompassing the wild-type and single deletion construct (linearized pX63-HYG-ΔGLOI designated as+/h GLOI) was transfected with linearized pX63-PHLEO-ΔGLOI construct in an an attempt to create double mutants. These transfecants were selected on semi-solid plates containing 1% Bacto-Agar (DIFCO) in M199 media (with 10% FBS) with G418 (40 μg/ml) and phleomycin (25 μg/ml) and hygromycin (200 μg/ml). A total of 9 clones were picked and screened by Southern blotting using GLOI probe. In order to check for the correct integration of PHLEO based replacement cassette phleo gene probe was used.

Glyoxalase I activity

1×10⁸ promastigotes each from wild type L. donovani, L. donovani mutant cells were harvested in the late log phase by centrifugation...
at 1500 × g, at 4°C for 15 min, washed with PBSG, pH 7.4. The activity of glyoxalase I was assayed spectrophotometrically at room temperature by measuring the initial rate of formation of S-D-lactoyl trypanothione at 240 nm as described by Racker with slight modification [30]. Trypanothione disulfide (1 mM) (Bio-Rad) was reduced with 3 mM DTT at 60°C for 20 min before the assay. The resulting reduced trypanothione was used for GLOI assay. The assay mixture contained, in a final volume of 0.5 ml; 100 mM MOPS buffer, pH 7.2; 400 μM methylglyoxal (Sigma); 300 μM reduced trypanothione and 20 μM NiCl₂ [4]. The assay mixture was incubated for 10 min followed by the addition of Leishmania cell lysate. Trypanothione hemithioacetal concentration was calculated by using the published Kd value of 3 mM for the methylglyoxal – glutathione equilibrium [39]. The value for the ΔA240 nm was taken as 2.86 M⁻¹ cm⁻³ for the isomerization of trypanothione hemithioacetal of methylglyoxal [40]. All assays were performed in triplicate.

Western blot analysis

Late log phase promastigotes (1 × 10⁶) from GLOI-GFP transformants were harvested and the resultant cell pellet was resuspended in lysis buffer (20 mM MOPS, pH 7.2; 1 mM DTT; 2 mM phenylmethylsulfonylfluoride (PMSF); 0.5 μg/ml each of leupeptin and aprotinin). The cell pellet was lyzed by freeze thaw and cell supernatants were prepared by centrifugation at 20,000 × g. The supernatant was collected and was used as cytosolic fraction [41].

For crude mitochondrial fraction mid log phase promastigotes were harvested and washed with PBSG. The cell pellet was resuspended in 5 mM Tris, pH 7.4 and kept at room temperature for 10 min. Suspension was homogenized using Potter Elvehjem Homogenizer and centrifuged at 3,500 rpm for 15 min to remove cellular debris. Supernatant was taken in eppendorf and centrifuged for 20 min at 12,000 rpm. Pellet was re-suspended in phosphate buffer (5 mM Na-PO₄), pH 7.4 at a protein concentration of 10 mg/ml [42]. All centrifugation steps and other operations were performed at 4°C. 100 μg of protein from each fraction was fractionated by SDS-polyacrylamide gel electrophoresis, blotted on to PVDF membrane using electrophoretic transfer cell (Bio-Rad) and probed with GLOI specific polyclonal antibody raised in mice (diluted 1:500) [5]. The blots were developed with 3', 5'-diaminobenzidine (Sigma). The mitochondrial and cytosolic fractions were also probed with mitochondrial specific anti –F1 ATP synthase α- subunit (anti-mouse polyclonal serum; diluted 1:250) [43].

DNA Content Analysis

L. donovani promastigotes (1 × 10⁷ cells/ml) were harvested by centrifugation at 2000 × g for 5 min at 4°C. Cells were washed once in phosphate-buffered saline (PBS; 0.01 M phosphate buffer, 2.7 M KCl, 0.137 M NaCl, pH 7.4). The cells were then fixed by incubation in 70% methanol/30% PBS for 1 h at 4°C. Prior to analysis, fixed cells were harvested by centrifugation at 1000 × g, for 10 min at 4°C, washed in PBS and then re-suspended in 1 ml PBS with RNase A (10 μg/ml) and incubated at 37°C for 20 min. Following this, propidium iodide (50 μg/ml) was added. Samples were then analyzed using a flow cytometer (FACS, Beckton Dickinson). For each sample at least 10,000 cells were analyzed.

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Author Contributions

Conceived and designed the experiments: RM. Performed the experiments: SCC. Analyzed the data: SCC RM. Contributed reagents/materials/analysis tools: RM. Wrote the paper: RM.

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