



Contents lists available at ScienceDirect

International Journal for Parasitology

journal homepage: www.elsevier.com/locate/ijpara

Independent origins of loss-of-function mutations conferring oxamniquine resistance in a Brazilian schistosome population [☆]

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ARTICLE INFO

Article history:

Received 1 December 2015

Received in revised form 31 March 2016

Accepted 31 March 2016

Available online xxxx

Keywords:

Schistosoma mansoni
Oxamniquine resistance
Sulfotransferase
Loss-of-function
Biochemical assay
Soft selective event

ABSTRACT

Molecular surveillance provides a powerful approach to monitoring the resistance status of parasite populations in the field and for understanding resistance evolution. Oxamniquine was used to treat Brazilian schistosomiasis patients (mid-1970s to mid-2000s) and several cases of parasite infections resistant to treatment were recorded. The gene underlying resistance (*SmSULT-OR*) encodes a sulfotransferase required for intracellular drug activation. Resistance has a recessive basis and occurs when both *SmSULT-OR* alleles encode for defective proteins. Here we examine *SmSULT-OR* sequence variation in a natural schistosome population in Brazil ~40 years after the first use of this drug. We sequenced *SmSULT-OR* from 189 individual miracidia (1–11 per patient) recovered from 49 patients, and tested proteins expressed from putative resistance alleles for their ability to activate oxamniquine. We found nine mutations (four non-synonymous single nucleotide polymorphisms, three non-coding single nucleotide polymorphisms and two indels). Both mutations (p.E142del and p.C35R) identified previously were recovered in this field population. We also found two additional mutations (a splice site variant and 1 bp coding insertion) predicted to encode non-functional truncated proteins. Two additional substitutions (p.G206V, p.N215Y) tested had no impact on oxamniquine activation. Three results are of particular interest: (i) we recovered the p.E142del mutation from the field: this same deletion is responsible for resistance in an oxamniquine selected laboratory parasite population; (ii) frequencies of resistance alleles are extremely low (0.27–0.8%), perhaps due to fitness costs associated with carriage of these alleles; (iii) that four independent resistant alleles were found is consistent with the idea that multiple mutations can generate loss-of-function alleles.

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1. Introduction

Surveys of drug resistance alleles using molecular markers provide a powerful approach to identify pathogen populations in which resistance is emerging, to map resistance spread, and for evidence-based resistance management. Molecular approaches

are now widely used for tracking resistance in malaria (Pearce et al., 2009; Ashley et al., 2014), HIV (Panichsillapakit et al., 2015) and bacterial diseases (Bhembe et al., 2014), and for managing insecticide resistance (Essandoh et al., 2013; Djègbè et al., 2014). Such methods are also now actively used for monitoring resistance in helminths of veterinary importance such as *Haemonchus contortus* (Rufener et al., 2009; Chaudhry et al., 2015; Redman et al., 2015), and have enormous potential for monitoring resistance in helminth parasites infecting humans, because existing phenotypic screening methods based on reductions in production of eggs or larval stages are insensitive for detection of low frequency resistance alleles and cannot detect recessive resistance alleles present in heterozygous worms. However, molecular

[☆] Note: Nucleotide sequence data reported in this paper are available in the GenBank[™], EMBL and DDBJ databases under the accession numbers KU951903–KU952091.

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<http://dx.doi.org/10.1016/j.ijpara.2016.03.006>

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surveillance is only possible when resistance genes have been identified, which is rarely the case for human helminth infections such as schistosomiasis.

Schistosomiasis, caused by three major species of blood flukes of the genus *Schistosoma* (Dye et al., 2013; Colley et al., 2014), is the second most important tropical parasitic disease after malaria (Steinmann et al., 2006), affecting an estimated 260 million people across Africa, Asia and South America, and killing over 200,000 people per year. Two drugs are available for treating schistosomiasis. Praziquantel (PZQ) is currently used as a monotherapy in expanding mass drug administration programs in Africa (Dye et al., 2013), making resistance evolution a major concern. A second drug, oxamniquine (OXA), is the focus of this paper. OXA acts specifically against *Schistosoma mansoni*, which is found in Africa, together with *Schistosoma haematobium*, and in South America, where it is the only species present. OXA was manufactured in Brazil by Pfizer (Cheetham, 1994) and widely used to treat *S. mansoni* infections from the mid-1970s to the mid-2000s (Coura and Amaral, 2004), but has now been replaced by PZQ as the first line drug.

Schistosomes resistant to OXA were first identified in Brazil in 1973 (Katz et al., 1973a,b), around the same time as the first clinical study of OXA treatment (Katz et al., 1973a,b). This rapid emergence of resistance was most likely due to the previous treatment of the same populations with the related drug hycanthon (Katz et al., 1968; Coura and Conceição, 2010), as both OXA and hycanthon have the same target (Jansma et al., 1977; Pica-Mattoccia et al., 1993) and the same mechanism of resistance (Pica-Mattoccia et al., 1992a). Later genetic studies showed that resistance is a recessive single locus trait (Cioli and Pica-Mattoccia, 1984; Pica-Mattoccia et al., 1993), most likely involving the absence of a sulfotransferase activity necessary for drug activation in resistant schistosomes (Pica-Mattoccia et al., 2006). The gene encoding the *S. mansoni* sulfotransferase involved in OXA resistance (*SmsULT-OR*) was recently identified by classical quantitative trait mapping in concert with crystallographic and functional analyses (Valentim et al., 2013). This work identified an amino acid deletion (p. E142del) in the laboratory-selected resistant parasite (HR), while an independent loss-of-function mutation (p.C35R) was identified in a field-collected resistant parasite line (MAP).

The identification of the gene involved in OXA resistance now allows us to examine distribution of resistance alleles of *SmsULT-OR* in natural populations. Despite the fact that the Brazilian government has switched from OXA to PZQ during the last decade (Uttinger et al., 2003), the recessive nature of OXA resistance allows the persistence of these alleles, because alleles encoding non-functional enzyme are not counter-selected as long as they segregate with alleles encoding a functional enzyme. We collected miracidia larval stages from a village in Minas Gerais, Brazil, and sequenced the *SmsULT-OR* gene in these samples. We sought to answer several questions: How common are resistance alleles? How many times have OXA resistance alleles arisen? Are the OXA resistance alleles selected in the laboratory and identified using linkage mapping actually present in nature? More broadly, our goal is to better understand the evolution of drug resistance in schistosomes and to demonstrate the utility of molecular screening approaches in anticipation that the gene(s) underlying PZQ resistance will soon be identified.

2. Materials and methods

2.1. Ethics statement

Stool samples were collected in accordance to the procedures of the Research Ethics Committee of the Universidade Federal de São

Paulo, Brazil (process number CAAE: 15567313.8.0000.5091). The purpose of the study and the procedures to be followed were explained and written informed consent was obtained from all participants or their legal guardians prior to any collection.

2.2. Sampling of *S. mansoni* miracidia

We collected stools from school children from Ponto dos Volantes (Minas Gerais, Brazil, GPS coordinates: 16°45'3.301"S, 41°30'13.755"W) and shipped these at 4 °C by ground transportation overnight to the Universidade Federal de Minas Gerais in Belo Horizonte, Brazil. We processed samples as follows: several grams of stools were filtered through three layers of sieves (mesh size: 250–45 µm) to obtain schistosome eggs. Eggs were transferred from the third sieve grid to a Petri dish and exposed under artificial light for at least 1 h. All filtering steps and egg transfer were performed with locally available bottled mineral water.

Washed eggs were observed under a stereomicroscope. For each patient, 1–11 living miracidia were sampled individually in ~2 µL of water and spotted onto CloneSaver FTA cards (GE Healthcare Life Sciences, USA). Spotted samples can be easily located on the cards because the pink dye on the cards turns white after water contact. Full cards were allowed to dry for 1 h at room temperature on the bench before being stored in a plastic bag and finally shipped to San Antonio, Texas, USA.

2.3. Preparation of FTA samples for whole genome amplification (WGA)

For each sample, we removed a 2 mm diameter disc from the FTA card using a 2 mm Harris Micro-punch (GE Healthcare Life Sciences). The 2 mm disc corresponds to the entire spot containing the whole miracidium. Each punch was placed individually in a 1.5 mL sterile tube. Punches were washed three times with FTA Purification Reagent (GE Healthcare Life Sciences) then rinsed twice with TE⁻¹ buffer (10 mM Tris, 0.1 mM EDTA, pH 8). Washing and rinsing steps were performed by adding 200 µL of solution to each tube followed by 5 min of incubation on a nutating mixer (24 RPM) at room temperature and then discarding the solution while minimising contact between the pipette tip and the punch. Punches were finally dried in tubes for 10 min at 56 °C on a dry bath incubator.

2.4. WGA

We performed WGA on each punch using the illustra Genomi-Phi V2 DNA Amplification kit (GE Healthcare Life Sciences). Punches were transferred in 0.2 mL sterile tubes using a sterile tip. We performed reactions following the manufacturer's instructions, immersing each punch in 9 µL of sample buffer and keeping tubes on ice at all times after the denaturation step. After amplification, we quantified DNA using the Qubit dsDNA BR assay (Invitrogen, USA).

2.5. Sequencing of *SmsULT-OR* exons

The two exons of the *SmsULT-OR* gene were amplified and sequenced independently. Each PCR was performed using the TaKaRa Taq kit (Clontech, USA) and composed of 9.325 µL of sterile water, 1.5 µL of 10× buffer, 1.2 µL of dNTP (2.5 mM each), 0.9 µL of MgCl₂ (25 mM), 0.5 µL of each primer (10 µM; Table 1), 0.075 µL of Taq polymerase (5 U·µL⁻¹) and 1 µL of DNA template. Amplifications were done using a GeneAmp PCR system 9700 thermocycler (Applied Biosystems, USA) with the following program: 95 °C for 5 min; 95 °C for 30 s, 60 °C for 30 s, 72 °C for 45 s, for 35 cycles; then 72 °C for 10 min.

Table 1Primer sequences used for PCRs and sequencing of the two exons of the *Schistosoma mansoni* *SmsULT-OR* gene in this study.

	Primer type	Primer sequence (5'–3' orientation)	Expected amplicon size (bp)	Usage
Exon 1	Outer forward primer	GCGAGATTCAAACCCAGGAT	822	PCR
	Outer reverse primer	GCCGTGATATTACTATCAATCCC		PCR
	Nested forward primer	GGGTAAGGAAGAGGGTTGG	545	PCR
	Nested reverse primer	TAAGAACAGACATATTAGACGAGT		PCR and sequencing
	Sequencing forward primer	TATATATGAAATATTATAACATTAC	–	Sequencing
Exon 2	Forward primer	ACTTCAACCAATCCCAAATCC	672	PCR and sequencing
	Reverse primer	AGTCCATTCAATCAATGTTTCAA		PCR and sequencing

Exon 1 required a nested PCR in order to obtain a specific product. Products from the first PCR were cleaned up by adding 4 µL of ExoSAP-IT (Affymetrix USB products, USA). Tubes were then incubated at 37 °C for 30 min and at 80 °C for 15 min using a thermocycler. Cleaned PCR products were then used as templates for the second PCR following the above protocol.

Sequencing reactions were performed using a BigDye® Terminator v3.1 cycle sequencing kit (Applied Biosystems) on final PCR products. PCR products were cleaned up using ExoSAP-IT as described above. Sequencing reactions were performed using 2.59 µL of sterile water, 1 µL of 5× running buffer, 0.25 µL of BigDye Terminator ready reaction mix (Applied Biosystems), 0.16 µL of forward or reverse primer used in the final PCR step, and 1 µL of final PCR product. Sequencing fragments were generated using a GeneAmp PCR system 9700 thermocycler (Applied Biosystems) with the following program: 96 °C for 1 min; 96 °C for 10 s, 50 °C for 5 s, 60 °C for 4 min, for 25 cycles. Sequencing reactions were cleaned up using a BigDye XTerminator® purification kit (Applied Biosystems). In each reaction, 20.45 µL of SAM™ solution and 4.55 µL of XTerminator™ solution were added. Reactions were then vortexed for 30 min and run on a 3730xl DNA Analyzer (Applied Biosystems).

Sequencing files were first screened using FinchTV (v1.4.0; Geospiza Inc.) to identify failed sequencing reactions. In the case of failure, sequencing reactions were performed a second time.

2.6. Variant identification and functional impact evaluation

We scored variants using PolyPhred software (v6.18) (Nickerson et al., 1997) which relies on Phred (v0.020425.c), Phrap (v0.990319), and Consed (v29.0) software, analysing each exon independently. We identified single nucleotide polymorphisms using a minimum phred quality score (-q) of 40, a minimum genotype score (-score) of 90, and a reference sequence that includes the *SmsULT-OR* gene and surrounding regions (position 1519500 to 1525200 of chromosome 6 of *S. mansoni* reference genome v5.0, ftp://ftp.sanger.ac.uk/pub/pathogens/Schistosoma/mansoni/genome/Assembly-v5/sma_v5.0.chr.fa.gz). Variant sites were labelled as non-reference alleles if they differed from the reference sequence. We identified insertion/deletion (indel) polymorphisms using a minimum phred quality score (-q) of 40 and a minimum genotype score (-score) of 80. Polymorphisms were visually validated using Consed. All the sequences were submitted to GenBank (GenBank accession no KU951903–KU952091).

Nucleic sequences showing mutations were translated in silico into protein sequences using the Translate tool from ExpASY portal (Artimo et al., 2012).

We evaluated the potential functional impact of identified polymorphisms on RNA features and on protein structure in silico. Modifications of RNA motifs and sites were analysed using the RegRNA2.0 website (Chang et al., 2013) for all available features with *Drosophila melanogaster* as the reference species when required. Modifications in protein structure were assessed using the mutagenesis function of PyMol software (v1.7.2.0; Schrödinger,

LLC) using the structure of *SmSULT-OR* determined previously (PDB code 4MUB, Valentim et al., 2013).

2.7. Population genetics analysis

We evaluated population structure by testing for Hardy–Weinberg equilibrium, measuring fixation indices (F_{st} and F_{is}) of parasites using Genepop (Rousset, 2008) with default options and considering schistosomes from a given patient as belonging to the same population. To identify positive selection, we calculated the number of synonymous (s) and non-synonymous (n) sites (Nei and Gojobori, 1986) and compared these values to synonymous (S) and non-synonymous (N) changes. We also performed a McDonald–Kreitman (MKT) test using the MKT server (<http://mkt.uab.es>; Egea et al., 2008). The *SmSULT-OR* homologue sequence in *Schistosoma rodhaini* was used as an outgroup and was obtained using tblastx from the NCBI server (NCBI Resource Coordinators, 2015) on the *S. rodhaini* genome assembly (GenBank accession no. GCA_000951475.1) using the *SmSULT-OR* sequence (GenBank accession no. KF733459.1) as a query.

2.8. Recombinant *SmsULT-OR* protein production

The *SmSULT-OR* sequence from the reference genome (Smp_089320; GenBank accession no. HE601629.1) was used to create a codon optimised synthetic gene (GenScript, USA) which was subcloned into the pAG8H vector derived from pKM260 (Melcher, 2000). We introduced mutations using a Phusion site-directed mutagenesis protocol (ThermoFisher Scientific, USA). Transformed *Escherichia coli* strain BL21 pLysS (Promega) were grown at 37 °C until the absorbance ($\lambda = 600$ nm) reached 0.7. We then decreased the temperature to 18 °C and induced expression by the addition of isopropyl- β -D-thiogalactoside (IPTG) at a final concentration of 1 mM. The cells were washed and resuspended in 50 mL of 50 mM Tris pH 8.0, 500 mM NaCl (column buffer) containing 250 µL of Sigma protease inhibitor cocktail per litre of culture and lysed by sonication on ice. The clarified supernatant was loaded onto a Ni²⁺-NTA affinity chromatography column (GE Healthcare), washed with five volumes of column buffer, and eluted using a 10–500 mM imidazole gradient. We pooled fractions identified as the Smp_089320 via SDS–PAGE and added His-tagged Tobacco etch virus (TEV) protease at a Smp_0893230:protease ratio of 15:1. The resulting solution was dialyzed overnight at 4 °C against 50 mM Tris pH 8.0, 100 mM NaCl, 2 mM dithiothreitol (DTT). We passed the dialysate over the Ni²⁺-NTA column again to remove the His tag and the His-tagged TEV protease, while the cleaved target protein flowed through. The sample was loaded onto a GE-pre-packed Q anion exchange column and eluted with a 0.1–1.0 M NaCl gradient. We pooled fractions containing Smp_089320 as identified by SDS–PAGE and dialyzed overnight at 4 °C against 25 mM Tris pH 8.5, 50 mM NaCl, 2 mM reducing agent tris-(carboxyethyl)-phosphine (TCEP) to prevent formation of intermolecular disulphide bonds. The product was ~98% pure as estimated by SDS–PAGE. We concentrated the purified

Smp_089320 protein to $10 \text{ mg}\cdot\text{mL}^{-1}$ using the calculated extinction coefficient $\varepsilon = 39,880 \text{ mol}^{-1}\cdot\text{cm}^{-1}$.

2.9. OXA activation assay

This assay utilises the fact that OXA is a prodrug that is enzymatically converted into a highly reactive molecule that covalently binds intracellular components such as DNA (Pica-Mattoccia et al., 1989). Because the original assay employs worm extracts which require time for preparation and introduce unpredictable variations in SmsULT-OR concentration (Pica-Mattoccia et al., 1992a; Valentim et al., 2013), we developed an improved in vitro assay that uses purified recombinant SmsULT-OR enzyme. This in vitro assay measures the enzymatic sulphonation of tritiated OXA molecules which then bind genomic DNA (gDNA). The resultant radioactive DNA-OXA complexes can then be quantified by a scintillation counter. Enzymes produced by non-functional *SmsULT-OR* alleles cannot sulphonate OXA, so no DNA-OXA complexes are formed.

Recombinant proteins were expressed and purified commercially by GenScript (p.C35R, p.P67L, p.E142del; Valentim et al., 2013) or by the Hart laboratory (University of Texas Health Science Center, USA) (p.G206V and p.N215Y). To determine which recombinant protein has the ability to activate OXA, 1 nM from each recombinant protein was added to 90 μL of a protease inhibitor cocktail (PIC) consisting of 0.1 M HEPES pH 7.4, 0.1 mM leupeptin, 2 μM E-64, 2 μM pepstatin A, 0.1 U of aprotinin, and 10 $\text{ng}\cdot\mu\text{L}^{-1}$ sheared *S. mansoni* gDNA as a final target. For each reaction 100 μCi of ^3H -OXA (Pica-Mattoccia et al., 1989) was solubilised in 2 μL of DMSO and added to 10 μL of a mixture containing the enzyme cofactors ATP and MgCl_2 at 50 mM each, and 3'-phosphadenosine-5'-phosphosulfate (PAPS) at 1 mM. The radiolabelled OXA and co-factor mix was then added to the PIC mix containing the recombinant protein. The resulting reaction was incubated for 2.5 h at 37 °C and stopped by adding three volumes of 0.1% SDS in 0.1 M sodium bicarbonate. The reaction was then extracted three times with two volumes of dichloromethane and the aqueous phase was counted in a liquid scintillation spectrometer (Beckman LS 6500 Scintillation Counter, USA) for 10 min. We also measured a blank solution (water) and the background scintillation. Blank and background values were subtracted from sample values. We performed three independent reactions for each recombinant protein.

2.10. Statistical analysis

Statistical analyses were done using R (v3.1.3) (R Core Team, 2015). For synonymous and non-synonymous changes, data were compared using a Fisher's exact test. For the OXA activation assay, the data were compared with a Welch *t*-test after testing for normality (Shapiro test, $P > 0.05$).

3. Results

3.1. Brazilian samples

We collected 232 FTA preserved miracidium samples from 51 patients (range: 1–12, mean \pm S.D.: 4.55 ± 3.65). We successfully amplified DNA from 204 samples (87.93%) from 50 patients (range: 1–11, mean \pm S.D.: 4.08 ± 3.26). Among the amplified samples, 189 from 49 patients (range: 1–11, mean \pm S.D.: 3.86 ± 3.18) contained schistosome DNA (92.65% of amplified samples, 81.47% of total samples). Among the 189 samples, we sequenced exon 1 from 183 samples and exon 2 from 188 samples. All samples had at least one exon sequenced (Supplementary Table S1).

3.2. SmsULT-OR variants in a Brazilian schistosome population

We scored nine mutations: four non-synonymous single nucleotide polymorphisms (SNPs), one insertion and one deletion in the coding region, and three non-coding SNPs (Table 2, Supplementary Table S1). The number of mutations in each exon (three in exon 1 and three in exon 2) did not differ regarding the length of the exon (327 bp and 447 bp, respectively) (Fisher's exact test, $P = 0.7$).

Among the seven SNPs, five (71.43%) were transitions and two (28.57%) were transversions. Four of the SNPs were located in the exonic region, two in exon 1 and two in exon 2. Three were present at very low frequency (0.0027–0.0053) while one was present at very high frequency (0.95). One SNP was identified at the first position of the intron at a low frequency (0.0056). The two remaining SNPs were found in the 3' untranslated region (UTR), one at high frequency (0.96), and one at low frequency (0.03).

One insertion was identified in exon 1 while one deletion was identified in exon 2, both at very low frequency (0.0027 and 0.0080, respectively).

Table 2

Mutations scored in the exons, intron, and 3' untranslated regions (UTR) of the *Schistosoma mansoni* *SmsULT-OR* gene. For each mutation, the corresponding nucleotide found in *Schistosoma rodhaini*, the number of homozygous and heterozygous samples carrying the non-reference allele, the number of samples sequenced (sample size), the allele frequency of the non-reference allele, the corresponding amino acid mutation, and the functional impact are shown.

	Nucleic mutation	<i>Schistosoma rodhaini</i> state	No. of homozygous samples for non-reference allele	No. of heterozygous samples for non-reference allele	Sample size	Frequency of the non-reference allele	Amino acid mutation	Functional impact
Exon 1	c.103T>C	T	0	1	183	0.0027	p.C35R	Misfolded protein (Valentim et al., 2013)
	c.200C>T	C	169	12	183	0.9563	p.P67L	No effect (Valentim et al., 2013)
	c.214_215insA	–	0	1	183	0.0027	p.T72NfsX5	Truncated protein with no active site
Intron	g.328G>A	G	1	0	179	0.0056	p.V110IfsX3	Splicing site disrupted leading to truncated protein with no active site
Exon 2	c.424_426delGAA	–	1	1	188	0.0080	p.E142del	Impaired oxamniquine binding (Valentim et al., 2013)
	c.617G>T	G	1	0	188	0.0053	p.G206V	No effect (Fig. 2)
	c.643A>T	A	0	1	188	0.0027	p.N215Y	No effect (Fig. 2)
3' UTR	g.4720C>T	C	177	9	188	0.9654	–	–
	g.4741T>C	T	2	8	188	0.0319	–	–

The code used for nucleic mutations indicates the sequence type (c = coding, g = gene), the position, and the mutation type (X>Y = substitution of X by Y, insN = insertion of N, delN = deletion of N). The code used for protein mutations indicates the sequence type (p = protein), the reference amino acid, the position, and finally the alternative amino acid, and when frame shift (fs) occurs, the position of the stop codon (X) after the mutation. For details about the nomenclature, see Ogino et al. (2007).

Low frequency mutations were found in heterozygous or in homozygous states among the samples. Observing rare SNPs present as homozygotes was surprising, suggesting population structure. The test for Hardy–Weinberg equilibrium showed a global deficit in heterozygosity ($P < 0.0001$). This is likely due to a deficit of heterozygous genotypes within host ($F_{is} = 0.3251$) rather than due to population differentiation between infections ($F_{st} = 0.0153$). Null alleles are not likely to explain the deficit. While we had small numbers of samples for which WGA failed, *SmSULT-OR* was successfully amplified from all samples for which WGA was successful.

Due to the absence of the synonymous mutations which precludes dN/dS calculation, we compared synonymous (S) and non-synonymous (N) changes with synonymous (s) and non-synonymous (n) sites to evaluate evidence for selection on *SmSULT-OR*. This comparison reveals no differences (Fisher's exact test, $P = 1$) indicating that there was no evidence for selection. We also tested directional evolution by performing a McDonald–Kreitman test using the homologue sequence of *SmSULT-OR* from *S. rodhaini* as an outgroup. We found no evidence for directional evolution ($\chi^2 = 1.466$, $P = 0.225$).

3.3. Functional impacts of mutations

3.3.1. Non-coding variants

We evaluated the functional impacts of the polymorphisms identified based on RNA features such as binding sites or splicing or regulatory motifs (Table 2). The mutation g.649G>A was predicted to modify the unique splice donor site at the end of exon 1. The disruption of the splicing site leads to translation of the beginning of the intron which ends four codons later due to the

introduction of a stop codon. This results in a truncated protein with no active site (Fig. 1, Supplementary Movie S1). Mutations present on the 3' UTR were not predicted to be in any regulatory sites.

3.3.2. Coding variants

All SNPs identified in the coding sequence of the gene were non-synonymous, and derived mutations relative to the outgroup *S. rodhaini* (Table 2). c.103T>C induced a substitution of cysteine to arginine (p.C35R) leading to a misfolded protein as shown previously (Valentim et al., 2013) (Fig. 1). c.200C>T induced a substitution of proline to leucine (p.P67L) and was previously found in the OXA sensitive strain (Valentim et al., 2013) (Fig. 1); this mutation therefore does not reduce enzyme activity. c.617G>T induced a substitution of glycine to valine (p.G206V) which occurs close the binding site of the PAPS co-factor and was previously observed in field-collected OXA resistant strain (Valentim et al., 2013) but was never tested. We postulated that this mutation may have a potential detrimental effect on co-factor binding and finally enzyme activity (Fig. 1, Supplementary Movie S1). The mutation c.643A>T induced a substitution of asparagine to tyrosine (p.N215Y) which occurs on a helix connected to a loop involved in co-factor binding. This mutation is predicted to have little impact on protein structure. We therefore postulated that this will not change protein function (Fig. 1, Supplementary Movie S1).

3.3.3. Indels

Both indels have detrimental effects on the enzyme (Table 2). The insertion c.214_215insA induces a frame shift leading to an early stop codon six codons after the mutation. This frame shift is predicted to produce a truncated enzyme with no active site.

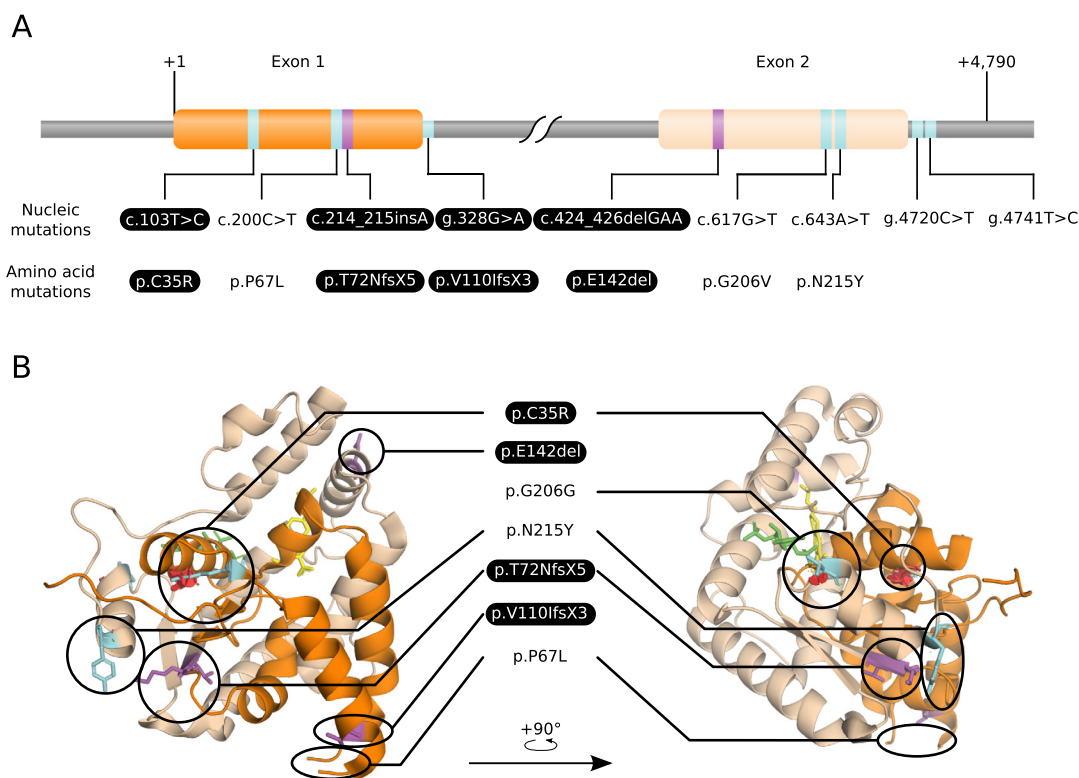


Fig. 1. Mapping of the mutations on the gene sequence and structure of *Schistosoma mansoni* SmSUTL-OR sulfotransferase. Exon 1 and exon 2 are represented in orange and beige, respectively. Single nucleotide polymorphisms and insertion/deletion events are represented in cyan and magenta, respectively. Loss-of-function mutations are highlighted in black. (A) Linear representation of the *SmSULT-OR* gene showing the relative position of the mutations and their translation in amino acid sequences. (B) Positions of mutations on the SmSULT-OR protein. Oxamniquine is represented in yellow, 3'-phosphoadenosine-5'-phosphosulfate (PAPS) co-factor is represented in green, and spatial distortions are represented by red discs. For a more detailed view of the mutations on the structure and their functional impact, see Supplementary Movie S1.

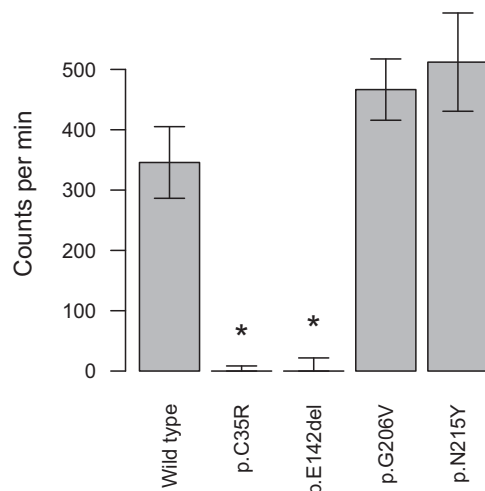


Fig. 2. Enzymatic activity of recombinant *Schistosoma mansoni* SmSULT-OR sulfo-transferase expressed from different allelic variants. This in vitro oxamniquine activation assay quantifies DNA-oxamniquine complexes by scintillation (counts per min) (see Section 2.9). Bars show the mean of three replicates, while error bars are S.E.M. Enzyme carrying loss-of-function mutations, such as p.C35R or p.E142del, showed no oxamniquine activation, while two newly identified alleles (p.G206V and p.N215Y) did not impair oxamniquine activation. * $P < 0.05$.

The deletion c.424_426delGAA is known from previous functional analyses to disrupt OXA binding (Valentim et al., 2013).

3.4. OXA activation assay

We produced two recombinant SmSULT-OR enzymes carrying the mutations p.G206V and p.N215Y in order to experimentally test whether these two mutations impact OXA activation, resulting in OXA resistance. We also tested known resistance alleles (p.C35R and p.E142del) as controls. The newly identified alleles are able to activate OXA as well as the reference allele (wild type) which does not carry any of these mutations (Welch t -test, $t < -1.55$, $P > 0.18$) while the known resistance alleles showed no activation as expected (Welch t -test, $t > 5.58$, $P < 0.021$) (Fig. 2). Therefore the two mutations tested (p.G206V and p.N215Y) did not disrupt co-factor binding or otherwise interfere with enzymatic activity.

4. Discussion

SmSULT-OR sequences from 189 miracidia collected from 49 Brazilian patients revealed nine mutations, including both mutations previously implicated in OXA resistance (Valentim et al., 2013). We found the p.C35R mutation, previously identified in the MAP strain. MAP was sampled from a patient living in a city of São Paulo state, Brazil (Pica-Mattoccia et al., 1992b), the neighbouring state of that from which our samples were collected (Supplementary Fig. S1). This suggests that the allele has been segregating in Brazilian parasite populations for more than 25 years. The second variant identified in the field, p.E142del, was previously found in the HR laboratory strain, which was initially sampled from a Puerto-Rican patient and subsequently selected with hycanthone in the laboratory (Cioli and Pica-Mattoccia, 1984) (Supplementary Fig. S1). This mutation could have arisen spontaneously in the laboratory. However, given that laboratory schistosome lines are maintained as outbred populations, the simplest explanation is that this mutation was segregating within the parasite population originally established in the laboratory. That we located this same deletion in two miracidia from Brazil and a parasite line collected from Puerto-Rico suggests that this allele may be widespread in schistosome populations from the New World.

It is not clear from our data whether the p.E142del seen in HR and Brazilian field samples arose independently or has a single origin. The HR p.E142del is found together with another mutation (p.L256W) (Valentim et al., 2013), which is absent from the p.E142del allele found in the Brazilian miracidia, providing some evidence for independent origins. Similarly, MAP differs from the Brazilian miracidia carrying the p.C35R mutation; while both also carry p.P67L, MAP carries an additional p.G206V mutation. Additional flanking SNP data will be required to critically test whether the p.E142del and p.C35R OXA resistance mutations have arisen a single time, or have multiple independent origins.

Laboratory selection is commonly used to explore the genetics of pathogen resistance, but a concern with this approach is that the mutations selected in the laboratory may poorly represent those occurring in nature. That we identified the same mutations in laboratory and field selected parasites is extremely encouraging, because ongoing work to identify mutations involved in PZQ resistance also utilises laboratory selected parasites (Couto et al., 2011). We note that laboratory selection experiments with *Plasmodium* also tend to identify the same genes and often the same mutations that are observed in the field (Anderson et al., 2011), further validating this approach.

Besides the two known alleles, two of these new mutations, c.214_215insA and g.328G>A, are predicted to confer OXA resistance; both introduce premature insertion of a stop codon producing a protein without an active site. These mutations therefore add two additional loss-of-function mutations that were probably selected during the OXA treatment. That four independent mutations are found in a single sampled parasite population is remarkable. Multiple origins have previously been observed in the evolution of resistance to benzimidazole drugs in the gastrointestinal nematode *Haemonchus* (Redman et al., 2015). The number of origins of resistance alleles is expected to depend on the size of the parasite population and the rate at which mutation generates resistant alleles (Messer and Petrov, 2013). In the case of *H. contortus*, the enormous size of parasite populations is likely to be the main driver, as only several specific mutations within β -tubulin can confer resistance. In the case of OXA resistance, a high mutation rate may be expected as the main driver, because multiple different mutations within SmSULT-OR can generate non-functional proteins. It is also possible that OXA resistant alleles were present within Brazilian *S. mansoni* populations prior to hycanthone or OXA treatment. Such standing variation may even have been present prior to the introduction of *S. mansoni* into South America. Analysis of SmSULT-OR in African populations, where OXA was not used extensively, will help answer this question.

Work on OXA resistance is simplified because we have an effective in vitro functional assay for screening allelic variants. The assay we used is an improvement on those previously described (Pica-Mattoccia et al., 1992a; Valentim et al., 2013), because worm homogenates are replaced by recombinant SmSULT-OR proteins. We identified two additional exon 2 substitutions (p.G206V, p.N215Y): Structural analyses suggested that one of these mutations (p.G206V) may disrupt OXA activation, by interfering with binding of the co-factor (PAPS), while the other (p.N215Y) is likely to have minimal impact on function. We were able to directly test these predictions by performing OXA activation assays. These assays demonstrated that neither mutation prevents OXA activation, allowing us to reject our prediction for p.G206V and confirm our prediction for p.N215Y. Hence, while structural studies are useful for formulating hypotheses about the consequences of mutations, functional assays are essential for critical testing of these hypotheses.

The two last mutations were found in the 3' UTR of the cDNA. In silico analysis, using predictions based on *Drosophila*, did not reveal regulatory sites in these regions, suggesting that these mutations

do not impact function. However, *Drosophila* may be a poor model to use. When more information on *SmSULT-OR* regulatory regions become available, new analyses of these two mutations may reveal potential effects on mRNA stability or translation rate.

The allele frequency of all four OXA resistance alleles combined is 3.8%, and only two parasites of 183 sampled (1%) are homozygous and therefore expected to be phenotypically resistant. The low frequency of the OXA resistance alleles can be explained by two non-exclusive hypotheses. First, resistance alleles may have remained at low levels even when OXA was the first line drug in Brazil. Second, fitness costs associated with these alleles may have driven reductions in allele frequency after the abandonment of OXA treatment. However OXA resistance alleles still persist within the populations in a heterozygous state, the cost being present only when worms are homozygous for the defective alleles.

Whether there is a cost associated with OXA resistance is questionable because miracidia homozygous for resistance alleles were found in the field in the present study. A previous study showed a reduction in infectivity and egg production in OXA resistant parasites relative to the OXA sensitive populations from which they were isolated (Cioli et al., 1992). However, this study suffers from a methodological limitation, because resistant and sensitive laboratory populations had different genetic backgrounds, complicating interpretation. Comparison of isogenic wildtype and genetically manipulated resistant parasites, or analysis of fitness in the progeny of a genetic cross between OXA resistant and sensitive parasites would allow measurement of associated fitness costs, while minimising confounding background effects.

New drugs are urgently needed for schistosome control because treatment currently relies on widespread monotherapy with PZQ. OXA is effective only against *S. mansoni*, but new OXA derivatives are under active development, with the aim of making compounds that are active against all three major schistosome species infecting humans (Taylor et al., 2015). If such derivatives are to be deployed clinically, understanding the capacity for resistance evolution in schistosome populations is of critical importance. For example, surveys of sequence variation in *SmSULT-OR* or the *S. haematobium* homologue would be an important prerequisite for field deployment of an OXA derivative active against both these species in Africa.

Our results have both positive and negative implications for field deployment of OXA derivatives. That multiple resistance alleles are present in a single parasite population suggests that resistance alleles may evolve and spread rapidly. On the positive side, existing resistance alleles are currently at extremely low frequency. It will be essential that OXA derivatives are deployed with appropriate partner drugs to minimise the rate of resistance evolution. We note that resistance to the antimalarial atovaquone was observed in the first clinical trial of this drug (Looareesuwan et al., 1996) and evolved de novo in different treated patients (Musset et al., 2007). Yet this drug is widely and effectively used with proguanil as a combination drug under the trade name Malarone.

Acknowledgements

This study was supported by NIH grants [R01-AI097576 (T.J.C. A.), 1R01AI115691 and P50 AI 098507 (P.T.L./P.J.H.)], World Health Organization [HQNTD1206356 (P.T.L.)], the UTHSCSA Presidents Collaborative Research Fund (P.T.L./P.J.H.) and the Robert A. Welch Foundation [AQ-1399 (P.J.H.)]. The molecular work at TBRI was conducted in facilities constructed with support from Research Facilities Improvement Program Grant (C06 RR013556) from the National Center for Research Resources (NIH). The AT&T Genomics Computing Center supercomputing facilities were supported by the AT&T Foundation and the National Center for Research

Resources Grant (S10 RR029392). The X-ray Crystallography Core Laboratory is supported by the University of Texas Health Science Center, San Antonio, Office of the Vice President for Research and by the San Antonio Cancer Institute Grant (P30 CA054174). W.L. was supported by a Cowles fellowship from Texas Biomedical Research Institute. N.E. was supported by the City of San Antonio Summer Ambassador Program. R.R.A. received support from CNPq (168260/2014-0). G.O. received support from CNPq (309312/2012-4), CAPES (070/13, REDE 21/2015) and FAPEMIG (RED-00014-14, PPM-00439-10). We thank Guilherme Oliveira, Dr. Fernanda Ludolf Ribeiro, Maycon Bruno and Nilvande Ferreira for technical assistance during stool processing and miracidia sampling, Dr. Andrea Gazzinelli and Dr. Leonardo Matoso for stool sampling at Ponto dos Volantes, Dr. Ricardo Toshio Fujiwara at Universidade Federal de Minas Gerais and Dr. Rodrigo Corrêa-Oliveira at Centro de Pesquisas René Rachou for providing laboratory space for stool processing and miracidia sampling, and Joanne Curran for use of a capillary sequencer.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.ijpara.2016.03.006>.

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