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RESEARCH ARTICLE

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Thioredoxin and glutathione systems differ in parasitic and free-living platyhelminths

Lucía Otero¹, Mariana Bonilla¹, Anna V Protasio², Cecilia Fernández¹, Vadim N Gladyshev³ and Gustavo Salinas*¹

Abstract

Background: The thioredoxin and/or glutathione pathways occur in all organisms. They provide electrons for deoxyribonucleotide synthesis, function as antioxidant defenses, in detoxification, Fe/S biogenesis and participate in a variety of cellular processes. In contrast to their mammalian hosts, platyhelminth (flatworm) parasites studied so far, lack conventional thioredoxin and glutathione systems. Instead, they possess a linked thioredoxin-glutathione system with the selenocysteine-containing enzyme thioredoxin glutathione reductase (TGR) as the single redox hub that controls the overall redox homeostasis. TGR has been recently validated as a drug target for schistosomiasis and new drug leads targeting TGR have recently been identified for these platyhelminth infections that affect more than 200 million people and for which a single drug is currently available. Little is known regarding the genomic structure of flatworm TGRs, the expression of TGR variants and whether the absence of conventional thioredoxin and glutathione systems is a signature of the entire platyhelminth phylum.

Results: We examine platyhelminth genomes and transcriptomes and find that all platyhelminth parasites (from classes Cestoda and Trematoda) conform to a biochemical scenario involving, exclusively, a selenium-dependent linked thioredoxin-glutathione system having TGR as a central redox hub. In contrast, the free-living platyhelminth *Schmidtea mediterranea* (Class Turbellaria) possesses conventional and linked thioredoxin and glutathione systems. We identify TGR variants in *Schistosoma* spp. derived from a single gene, and demonstrate their expression. We also provide experimental evidence that alternative initiation of transcription and alternative transcript processing contribute to the generation of TGR variants in platyhelminth parasites.

Conclusions: Our results indicate that thioredoxin and glutathione pathways differ in parasitic and free-living flatworms and that canonical enzymes were specifically lost in the parasitic lineage. Platyhelminth parasites possess a unique and simplified redox system for diverse essential processes, and thus TGR is an excellent drug target for platyhelminth infections. Inhibition of the central redox wire hub would lead to overall disruption of redox homeostasis and disable DNA synthesis.

Background

Platyhelminths (commonly known as flatworms) are a metazoan phylum that includes the neodermata lineage, composed exclusively of parasitic taxa, and the turbellaria lineage, mostly composed of free-living taxa [1]. Neodermatan flatworms comprise the classes Cestoda, Trematoda and Monogenea, of which the first two include the agents of serious human diseases. Notably, trematode infections caused by *Schistosoma* spp. affect more than 200 million people in Africa, South America and Asia [2].

Cestode infections are less prevalent, but include severe human diseases such as cysticercosis and hydatid disease, caused by *Taenia solium* and *Echinococcus* spp., respectively [3]. As yet, there is no vaccine that can prevent platyhelminth infections in humans, neither there are trials underway. Although chemotherapy is the mainstay of control, very few effective drugs are currently used to treat platyhelminth infections, being praziquantel the single drug readily available for schistosomiasis treatment [4].

In most organisms, including the mammalian hosts of platyhelminth parasites, cellular redox homeostasis, antioxidant defenses and supply of electrons for deoxyribonucleotide synthesis rely on two major and independent

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pathways: the glutathione (GSH) and the thioredoxin (Trx) systems [5], which have overlapping and differential targets, and function in a great variety of biological processes. These pathways operate through redox cascades that involve transfer of reducing equivalents from NADPH to targets through a series of dithiol-disulfide reactions or variations of this theme (e.g. when selenocysteine, Sec, replaces cysteine) [6]. The core enzymes of these pathways are glutathione reductase (GR) and thioredoxin reductase (TR), both of which are pyridine-nucleotide thiol-disulfide oxidoreductases that reduce the oxidized tripeptide glutathione (GSSG) and the oxidized disulfide reductase thioredoxin (Trx), respectively. In turn, GSH and Trx transfer electrons to downstream targets. Platyhelminth parasites, unlike their mammalian hosts, lack conventional GR and TR enzymes, and hence canonical thioredoxin and glutathione systems [7-9]. Instead, they possess a linked glutathione thioredoxin system that relies exclusively on the selenoenzyme thioredoxin glutathione reductase (TGR) for provision of reducing equivalents to both pathways. TGR achieves this broad substrate specificity by a fusion of an N-terminal glutaredoxin (Grx) domain to TR domains [10]. In the platyhelminth parasite *Echinococcus granulosus*, cytosolic and mitochondrial TGR variants derived from a single gene have been reported [11] and functional thioredoxin-glutathione systems have been recently described in both compartments [12], whereas only a cytosolic variant of TGR has been described in *S. mansoni* [13]. The differences in the thioredoxin and glutathione pathways between parasitic flatworms and their mammalian hosts, and the lack of redundancy of these redox pathways have prompted studies which have recently resulted in validation of TGR as a novel drug target for platyhelminths [14]. Indeed, new drug leads have been identified by quantitative high-throughput screenings using *Schistosoma mansoni* TGR as a target [15].

The fact that TGR is an essential enzyme that controls the overall redox homeostasis in these parasites warrants further studies on flatworm TGRs. In particular, little is known relating the genomic structure of flatworm TGRs, whether additional TGR variants are expressed, and how the variants are generated. More importantly, it is not known whether the presence of TGR and the absence of TR and GR genes is a signature of the platyhelminth lineage. In this work, we report that additional parasitic flatworms possess TGR and lack TR and GR. In contrast, the free-living platyhelminth *Schmidtea mediterranea* possesses TR, GR and TGR genes. In addition we investigate the existence, generation and expression of TGR variants in parasitic flatworms.

Results

Analysis of thioredoxin and glutathione systems in free-living and parasitic platyhelminths

We carried out an exhaustive *in silico* analysis of available genome and transcriptome data from platyhelminth organisms to examine the presence of TGR, TR and GR sequences. A tblastn search of the *E. multilocularis* genome using *E. granulosus* TGR sequence as protein query revealed that *E. multilocularis* genome possesses a single TGR gene and lacks genes encoding conventional TR or GR, consistent with previous experimental evidence from *E. granulosus*. *E. multilocularis* TGR is a selenoprotein: its gene contains an in-frame TGA codon as well as a SECIS element. *Echinococcus* TGR orthologs are virtually identical; they possess 95% identity at the nucleotide level and 98% identity at the amino acid level (Figure 1). On the other hand tblastn searches of the *S. mediterranea* genome revealed that, in contrast to the parasitic flatworms *E. multilocularis* (class Cestoda) and *Schistosoma* spp. (class Trematoda), the free-living platyhelminth *S. mediterranea* (class Turbellaria) possesses a TR gene, a GR gene and a TGR gene. The coding sequences of *S. mediterranea* TR, GR and TGR were predicted based on EST sequences when available; final adjustments of intron-exon boundaries were performed based on a multiple alignment of TRs, GRs and TGRs from different species. The deduced amino acid sequences are shown in Figure 1. All *S. mediterranea* pyridine-nucleotide thiol-disulfide oxidoreductases identified contain the canonical CX₄C redox center. Both TR and TGR genes from *S. mediterranea* contain an in-frame TGA codon and a SECIS element, and thus encode selenoproteins, with a GCUG (U denotes selenocysteine) C-terminal redox center. *S. mediterranea* TGR possesses a dithiol Grx domain, containing the CPFC redox active center, similar to *S. japonicum* TGR and TGRs from other organisms. The TR gene from *S. mediterranea* has higher identity to mammalian mitochondrial TRs. Indeed, an exon encoding a putative mitochondrial leader peptide was detected upstream of the TR coding sequence. In the absence of additional platyhelminth genomes sequenced, we searched databases for expressed TRs, GRs and TGRs of other flatworms. A tblastn search identified a cDNA encoding a Sec-containing full-length TGR from the platyhelminth parasite *Fasciola hepatica* (class Trematoda), with high similarity to *Schistosoma* TGRs and containing the canonical CPYC redox center at the Grx domain, the CX₄C redox center of pyridine-nucleotide thiol-disulfide oxidoreductases, and a Sec-containing C-terminal redox motif. Additionally, a tblastn search revealed ESTs encoding a fragment of a TGR from the platyhelminth parasite *Taenia solium* (class Cestoda). This TGR is highly similar to *Echinococcus* TGRs. Figure

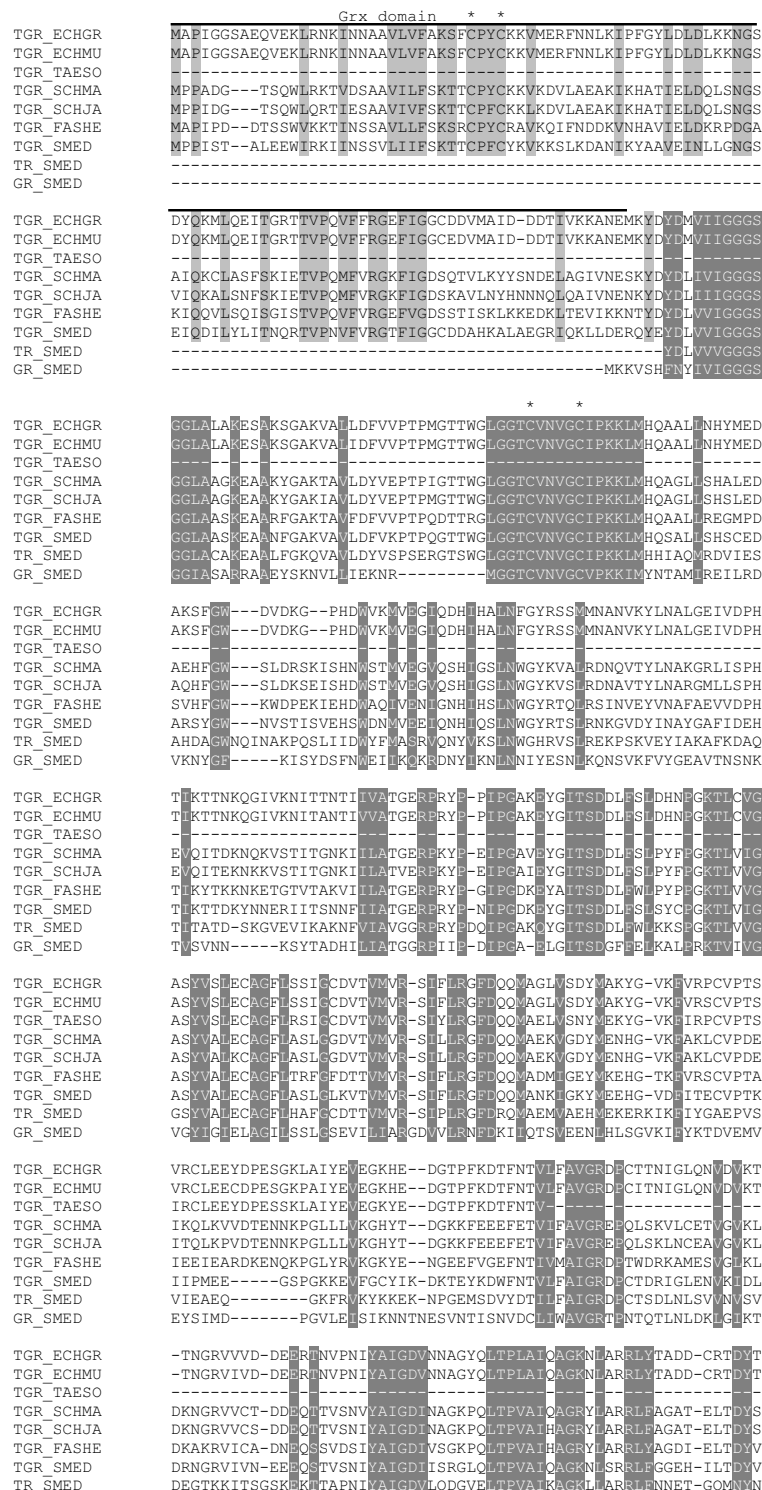


Figure 1 Amino acid sequence alignment of TR, GR and TGRs of platyhelminths. Sec is indicated by U. The position of the redox active residues in the sequences is indicated by a star. Conserved residues in all proteins are highlighted in dark grey, conserved residues in the Grx domain of TGRs are highlighted in light grey. Location of the Grx domain is indicated above the sequence. ORFs for TGR, TR and GR from *S. mediterranea* (SCHME) genome (assembly 31) were predicted in the contigs 000676, 000203 and 001663, respectively. ORF for *E. multilocularis* (ECHMU) TGR was assembled from contigs 0007357 and 0007358. Full-length TGR sequences of *S. mansoni* (SCHMA), *S. japonicum* (SCHJA), *E. granulosus* (ECHGR), and *F. hepatica* (FASHE) were retrieved from Genebank (gb|AAK85233.1|AF395822.1, gb|AAW25951.1, emb|CAM96615.1, and gb|AAN63052.1 respectively). *T. solium* (TAESO) TGR partial sequences were retrieved from the EST repository at Genebank (gb|EL757065.1 and gb|EL743442.1). The putative mitochondrial leader peptide of *S. mediterranea* TGR is not included in the sequence, neither leader peptide variants of *Schistosoma* and *Echinococcus* TGRs. Sequences were aligned with Clustal W2 [29], with final manual adjustment after inspection.

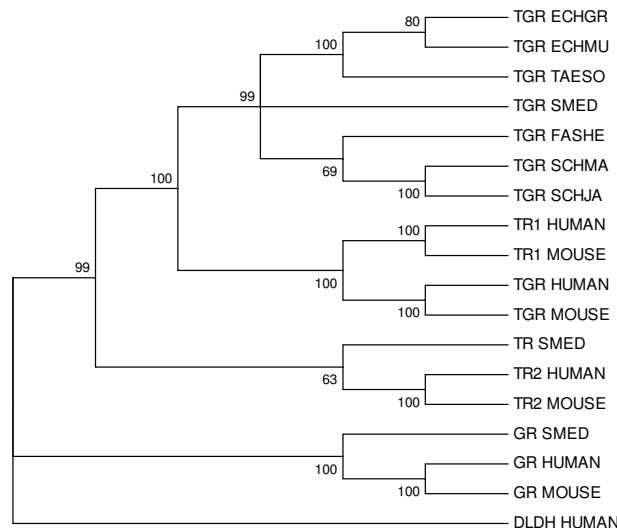


Figure 3 Phylogenetic relationships of GRs, TRs and TGRs of platyhelminths and mammals. TRs, GRs and TGRs from platyhelminths and mammals were aligned with Clustal W2 [29]. Human dihydroliipoamide dehydrogenase (DLDH) was used as outgroup. A Neighbor-Joining tree was constructed using MEGA4 [30] with pairwise deletion and default parameters. A condensed tree is shown, and bootstrap values of reliable nodes (above 50) are indicated. The polytomy displayed at the TGR node denotes that the evolutionary relationships within the node can not be resolved with at least 50% of bootstrap support. In other words, nodes with less than 50% of bootstrap support were collapsed and are displayed as polytomies. The results indicate that the TR and GR genes present in the planarian lineage were lost in the neodermata lineage. Very similar topology and statistical support were obtained using different phylogenetic reconstruction methods (i.e. Maximum Parsimony, UPMGA and Minimum Evolution). ECHGR: *E. granulosus*, ECHMU: *E. multilocularis*, TAESO: *T. Solium*, SCHMA: *S. mansoni*, SCHJA: *S. japonicum*, SMED: *S. mediterranea*, FASHE: *F. hepatica*.

sequence; they occurred in the coding sequence corresponding to the C-terminal half of TGR. Thus, the gene structure appeared to have more constraints at the Grx domain than at the interface domain. In turn, *S. mediterranea* TR and GR genes displayed a completely different exon/intron structure than platyhelminth TGR genes.

Finally, since only a handful of genes from *Echinococcus* have been sequenced, we analyzed in detail various aspects of *Echinococcus* TGR genes. We examined exon-intron boundaries to identify common features and differences (Additional file 1 shows all the exon-intron boundaries of TGR gene). All but one intron contained the canonical GU-AG donor-acceptor sites, typical of eukaryotic nuclear pre-mRNA, being the donor site of intron 15 the exception to the consensus. In addition, we searched for conserved bases or motifs around the splice sites and found that C or A followed by a purine (A or G) preceded the 5' GU splice site in most cases; and that U or A was present at the -3 base with respect to the 3'AG splice site (Additional file 1). The T+A content in TGR introns was 59% (contrasting 50.7% in exons), indicating a neutral mutational bias towards T+A, as previously noted for *Echinococcus* genes [16]. The small size of all *Echinococcus* TGR introns is noteworthy: while *Echinococcus* TGR genes span 6 kb, *S. mediterranea* and *Schistosoma* spp genes are significantly longer. These data agree with the previous observations that *Echinococcus* genes possess small introns [16]. Taken as a whole, our results are

also in agreement with previous findings that both *Echinococcus* species are remarkably similar with regard to sequence information.

Identification of TGR variants in platyhelminths

We have previously demonstrated the existence of mitochondrial and cytosolic TGR variants derived from a single gene in *E. granulosus*. In order to identify TGR variants expressed in other flatworms, we performed a tblastn search at the NCBI server (EST others option). Only a single TGR variant was identified in the case of *S. mediterranea*. In the case of *S. japonicum*, ESTs encoding two additional TGR variants to the already reported cytosolic TGR were identified, whereas a single additional variant was identified in *S. mansoni*. Similarly to what has been described in *E. granulosus*, the *Schistosoma* cDNAs encode variants differing in their N-termini (Figure 4). One of the *S. japonicum* variants (TGR_SCHJA_v1) would encode an N-terminal mitochondrial signal peptide. The other *S. japonicum* variant (TGR_SCHJA_v2) and the *S. mansoni* variant identified (TGR_SCHMA_v2) encode a leader peptide related to, but shorter than, the mitochondrial one. It is not possible, *in silico*, to ascribe a topological signal to this variant. The genomic sequence and exon-intron structure of *Schistosoma* TGR genes strongly support the existence of these variants. Indeed, the information for the identified *Schistosoma* leader peptides is found in an exon upstream of the one encod-

Table 1: Exon and intron structure of flatworm TR, GR and TGRs

Exon/Intron	Em_TGR	Eg_TGR	Sma_TGR	Sja_TGR	Sme_TGR	Sme_TR	Sme_GR
E1	69	69	93	93			
l1	281	284	1507	971			
E2	120	120	108	108	105	55	132
l2	40	40	34	35	45	48	43
E3	61	63	61	61	63	57	116
l3	41	41	33	30	3066	48	46
E4	87	85	87	87	81	164	70
l4	118	127	36	36	301	397	44
E5	122	122	125	125	132	54	133
l5	701	705	2298	2614	2389	52	4105
E6	142	142	142	142	144	139	55
l6	162	164	214	115	41	9	4262
E7	48	48	54	54	48	54	73
l7	74	74	2166	179	3006	533	71
E8	143	143	143	143	141	74	246
l8	300	305	840	643	49	44	8817
E9	116	116	116	116	342	20	378
l9	121	119	4231	848	998	75	60
E10	226	226	226	226	165	92	216
l10	885	912	819	1240	59	1704	
E11	105	105	105	105	159	178	

Table 1: Exon and intron structure of flatworm TR, GR and TGRs (Continued)

I11	211	223	1865	700	58	46	
E12	74	74	74	74	339	89	
I12	68	68	2924	2180	1382	18	
E13	154	154	157	157	338	45	
I13	198	202	1788	2054		54	
E14	108	108	108	112		195	
I14	81	81	1618	1191		2858	
E15	93	93	93	89		63	
I15	502	678	2491	293		58	
E16	138	138	138	138		104	
I16	305	309	170	908		44	
E17	484	485	480	423		378	
Σtot	6378	6623	25344	16290	13451	7749	18867
Σexons	2290	2291	2310	2253	2057	1761	1419

The length of exons and introns is shown for *E. multilocularis* TGR (EmTGR), *E. granulosus* TGR (EgTGR), *S. mansoni* TGR (SmaTGR), *S. japonicum* TGR (SjaTGR), *S. mediterranea* TGR (SmeTGR), *S. mediterranea* TR (SmeTR) and *S. mediterranea* GR (SmeGR). Numbering corresponds to *Echinococcus* TGR.

ing the N-terminal sequence of the Grx domain. TGR_SCHJA_v1 (putatively encoding a mitochondrial TGR variant) and TGR_SCHJA_v2 (with a shorter leader peptide) are derived from alternative splicing of exon I and exon II at a canonical GU donor site present in intron I and at a leaky GU donor site present in exon I, respectively. If the transcript is spliced at this latter GU donor site, it would give rise to a shorter exon I and, consequently to a shorter leader peptide (Figure 4c). The same gene structure is observed in *S. mansoni* TGR suggesting that, in addition to TGR_SCHMA_v2, a mitochondrial variant of TGR (TGR_SCHMA_v1) also exists in this species (Figure 4c). The sequences of the full-length cDNA variants encoding *Schistosoma* cytosolic TGRs, and the structure of the genes suggest that the cytosolic variants (TGR_SCHJA_v3 and TGR_SCHMA_v3) are derived from alternative initiation of transcription, from a puta-

tive promoter at intron I, similar to what has been hypothesized for the *E. granulosus* variants. Figure 4d shows a model for the generation of *Schistosoma* TGR variants, which takes into account all this information.

In order to confirm the presence of the newly identified variant in *S. mansoni* and assess the occurrence of the putative mitochondrial variant (identified in *S. japonicum* ESTs, but not in *S. mansoni*), we performed PCRs from different *Schistosoma* samples (schistosomula, adult worm and cercariae) with forward primers specific for each of the leader peptide variants and a common TGR reverse primer; in addition we used, as a control, a forward primer corresponding to the 5'end of the cytosolic variant (Figure 5a). PCR products of the expected sizes were obtained in the three reactions in all materials, confirming the expression of both leader peptide variants, TGR_SCHMA_v1 and TGR_SCHMA_v2. Figure 5b

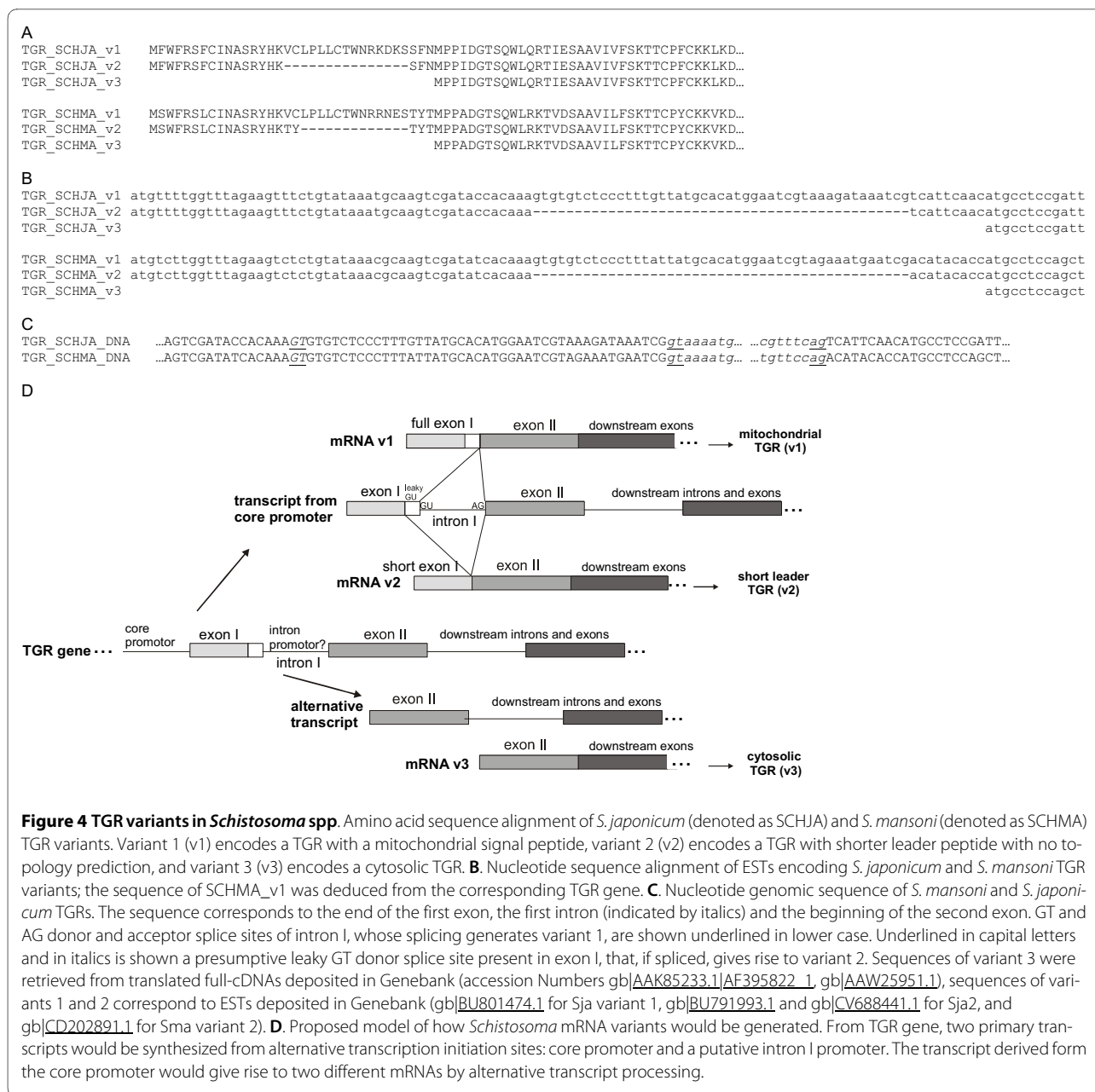


Figure 4 TGR variants in *Schistosoma* spp. Amino acid sequence alignment of *S. japonicum* (denoted as SCHJA) and *S. mansoni* (denoted as SCHMA) TGR variants. Variant 1 (v1) encodes a TGR with a mitochondrial signal peptide, variant 2 (v2) encodes a TGR with shorter leader peptide with no topology prediction, and variant 3 (v3) encodes a cytosolic TGR. **B.** Nucleotide sequence alignment of ESTs encoding *S. japonicum* and *S. mansoni* TGR variants; the sequence of SCHMA_v1 was deduced from the corresponding TGR gene. **C.** Nucleotide genomic sequence of *S. mansoni* and *S. japonicum* TGRs. The sequence corresponds to the end of the first exon, the first intron (indicated by italics) and the beginning of the second exon. GT and AG donor and acceptor splice sites of intron I, whose splicing generates variant 1, are shown underlined in lower case. Underlined in capital letters and in italics is shown a presumptive leaky GT donor splice site present in exon I, that, if spliced, gives rise to variant 2. Sequences of variant 3 were retrieved from translated full-cDNAs deposited in Genebank (accession Numbers gb|AAK85233.1|AF395822_1, gb|AAW25951.1), sequences of variants 1 and 2 correspond to ESTs deposited in Genebank (gb|BU801474.1 for Sja variant 1, gb|BU791993.1 and gb|CV688441.1 for Sja2, and gb|CD202891.1 for Sma variant 2). **D.** Proposed model of how *Schistosoma* mRNA variants would be generated. From TGR gene, two primary transcripts would be synthesized from alternative transcription initiation sites: core promoter and a putative intron I promoter. The transcript derived from the core promoter would give rise to two different mRNAs by alternative transcript processing.

shows the result of the PCR reactions using cercariae as a cDNA template.

Finally, we investigated whether there are additional variants in *E. granulosus* or whether the already known variants are developmentally regulated. To this end, we performed RT-PCRs using the splice leader exon of *E. granulosus* and TGR reverse specific primers on different *E. granulosus* materials (larval worms, adult worms and germinal layer of hydatid cyst). To allow discrimination of variants that may subtly differ in size, the PCR products obtained were analyzed on polyacrylamide gels. The results suggest that no additional variants to the already described are expressed in *E. granulosus* and that the

mitochondrial and cytosolic variants are expressed in all stages analyzed (data not shown).

Analysis of transcription from alternative promoter site

We previously postulated that the mRNA variants derived from a single TGR gene may arise from two alternative transcription initiation sites located upstream of the first exon (core promoter) and upstream of the second exon (intron promoter), because: i) both variants contained the trans-splice leader exon and ii) a putative TATA box was present within the intron sequence [11]. The comparison of the core promoter and the putative intron promoter nucleotide sequences of *Echinococcus*

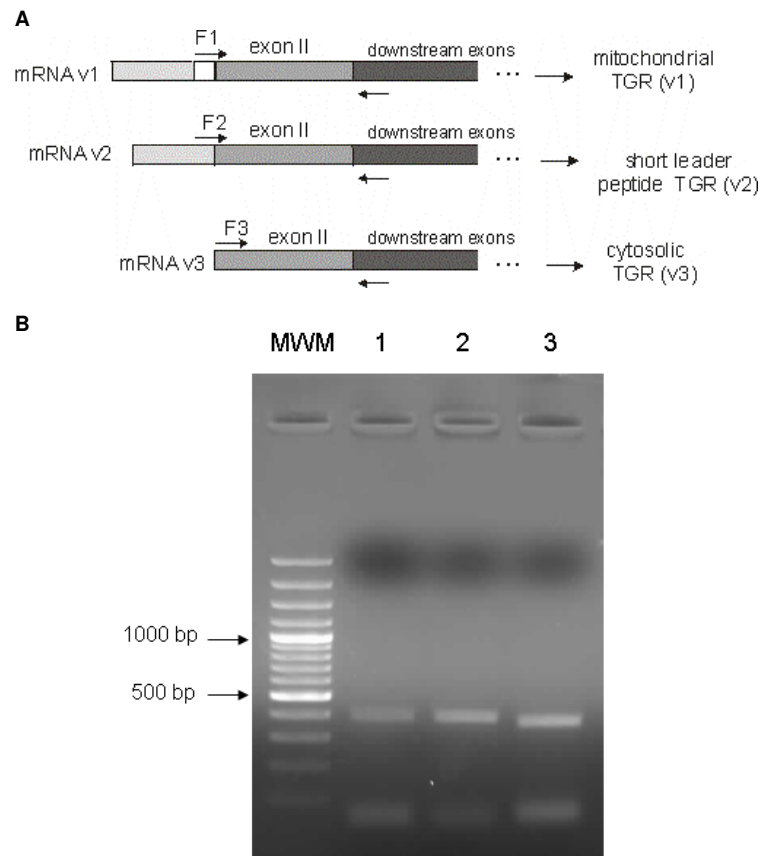


Figure 5 Expression of *S. mansoni* variants detected by RT-PCR. **A.** Schematic representation of the three mRNA variants of *S. mansoni* TGR, the variant 1 and variant 2 specific primers (F1 and F2) and the primer corresponding to the 5'-end of the cytosolic variant (F3) used in RT-PCRs in combination with a reverse primer derived from exon 3 sequence. **B.** Electrophoresis of PCR products from *S. mansoni* cercariae: a band of the expected size was observed in the PCRs with F1, F2 and F3 primers and the reverse primer (lanes 2 to 4, respectively). MWM lane corresponds to 100 bp plus ladder (Fermentas).

and *Schistosoma* TGRs did not evidence significant similarity, although there were some conserved regions at both presumptive promoter regions (data not shown). We tested the promoter activity of the putative intron promoter of *E. granulosus* TGR in a heterologous system (CHO-K1 cells) using a reporter vector (pAcGFP1-1™, Clontech), but this approach was not successful. Thus, in order to obtain evidence of alternative transcription initiation in *E. granulosus*, we performed RT-PCR with forward primers mapping the first intron and a reverse primer from a downstream exon. This strategy has been previously used [17] and its rationale relies on the fact that trans-splicing is, kinetically, a much slower process than cis-splicing, and thus it is possible to find a population of transcripts not completely processed; these transcripts will have been already cis-spliced, but have not been yet trans-spliced. The results indicate that alternative transcription initiation takes place from intron 1, downstream of the TATA box, since a PCR product was obtained with the forward primer 50 bp downstream of

the putative TATA sequence present in the intron, but not with primers immediately downstream or upstream from the TATA sequence. Since TGR mRNAs from *Schistosoma* TGRs are not trans-spliced [13] this strategy could not be used to assess alternative initiation of transcription from intron I.

Discussion

Previous studies on thioredoxin and glutathione systems in parasitic flatworms indicated that these systems are different to those of their vertebrate hosts [7-9,11]. The main difference is the absence of conventional GR and TR in these parasites, and the substitution of both enzymes by TGR as a sole redox "wire". Our analysis of platyhelminth genomes and transcriptomes indicates that this is the case for all flatworm parasites for which information is available. The *E. multilocularis* genome revealed that this cestode also lacks conventional TR and GR, and its redox thiol-disulfide homeostasis relies exclusively on a single Sec-containing TGR, in agreement with previous bio-

chemical and molecular studies of *E. granulosus*. A cDNA encoding TGR was also identified from *T. solium*, the other medically important cestode. A similar scenario was observed in the case of trematodes: lack of conventional GR and TR and presence of TGR in the genomes of *Schistosoma* spp. [18,19], and a cDNA encoding TGR from *E. hepatica*, an additional member of the class. Thus the dependence on the single oxidoreductase is common to the neodermata lineage (a platyhelminth subphylum that contains the classes Cestoda and Trematoda, but not the free-living class Turbellaria). Our results further revealed that the absence of TR and GR is not characteristic of the entire platyhelminth phylum. The turbellarian *S. mediterranea* possesses TR, GR and TGR, indicating that flatworm parasites have lost TR and GR genes. Elimination of GR and TR from their genomes suggests streamlining of redox regulation.

Another interesting finding in our study is the identification of novel variants of *S. mansoni* and *S. japonicum* TGRs. Previously, cytosolic and mitochondrial variants derived from a single gene were described in *E. granulosus* [11]. In the case of *Schistosoma*, we could detect *in silico* and confirm experimentally the existence of two variants in addition to the previously reported cytosolic form. One of the novel forms encodes a mitochondrial TGR; and this is in agreement with the occurrence of cytosolic and mitochondrial thioredoxins in *Schistosoma* [18]. The second newly identified TGR variant encodes a shorter leader peptide. The functional significance of this leader peptide (*e.g.* topological localization information or a regulatory role) must await functional studies. In many species, distinct genes encode for mitochondrial and cytosolic TRs. Nevertheless, complex expression patterns and rare mRNA variants have also been observed for all TR genes in mammals [20-26], and the functional significance of this complexity is not fully understood. In this context, it is important to note that the three variants of *S. mansoni* TGR and the cytosolic and mitochondrial variants of *E. granulosus* are expressed in all the life cycle stages examined (cercariae, schistosomule and adult worms in the case of *S. mansoni*; larval worms, hydatid cyst wall and adult worms in the case of *E. granulosus*). The existence of TGR variants reveals a relatively complex expression of the gene in these organisms that may be relevant when considering TGR as a drug target; in particular if inhibition is to be achieved at different sub-cellular compartments.

The study of the mechanism by which TGR variants may be generated in parasitic flatworms was addressed by two different strategies. The initial promoter reporter strategy was not successful. However, in the case of *E. granulosus* it was possible to map the initiation of transcription from the intron promoter, downstream from the presumptive TATA box. Thus, our results suggest

that two promoters (core and intron promoters) give rise to two different transcripts. The absence of the splice leader exon in *Schistosoma* TGR mRNAs precluded the use of this strategy. Nevertheless, the information is consistent with the model depicted in Figure 4d. One of the transcripts would be derived from the core promoter and would be alternatively spliced, generating the two variants with related but different N-terminal signal peptides, and an additional variant would be generated from alternative initiation of transcription.

Conclusions

Our study found that the biochemical scenario in parasitic flatworms differs from the one observed in free-living flatworms. A unique and simplified redox system, termed linked thioredoxin-glutathione system, is present in all platyhelminth parasites examined. In contrast, conventional and linked thioredoxin and glutathione systems were found in *S. mediterranea*, a free-living platyhelminth. The data also show that canonical GR and TR were specifically lost in the parasitic platyhelminths. Overall, our results reinforce previous studies indicating that TGR is an excellent drug target for flatworm infections [14,15]; indeed, maintenance of redox homeostasis and basic cellular processes such as synthesis of deoxyribonucleotides depend on a single selenoenzyme in different classes of flatworm parasites. Furthermore, the phylogenetic analysis of TRs, GRs and TGRs indicates that flatworm TGRs form a clade with no clear ortholog to either mammalian TRs or TGR; these differences may be relevant for selective inhibition. The fact that all flatworm TGRs contain selenocysteine at the active site constitutes a unique opportunity to target this particularly reactive nucleophilic residue. Alternatively, flatworm parasite redox homeostasis could be disrupted targeting selenium incorporation, for instance by selectively inhibiting platyhelminth selenophosphate synthetase or SECIS-binding protein.

Our work also demonstrates the expression of TGR variants in the human parasites *Schistosoma* spp. Finally, we provide evidence that alternative initiation of transcription and transcript processing contribute to the generation of TGR variants in platyhelminth parasites, revealing plasticity of gene expression in these organisms.

Methods

Identification of TR, GR and TGR genomic, cDNA and coding sequences in platyhelminth databases

In order to obtain information on platyhelminth TR, GR and TGR sequences, the *S. mansoni*, *S. japonicum*, *E. multilocularis* and *S. mediterranea* genomic databases (http://www.sanger.ac.uk/Projects/S_mansoni, <http://lifecenter.sgst.cn/schistosoma/en/schistosomeBlast.do>, <http://www.sanger.ac.uk/Projects/Echinococcus>, and

<http://smedgd.neuro.utah.edu>, respectively) were searched with tblastn, using the sequences of *S. mansoni* and *E. granulosus* TGRs and human GR, TRs and TGR as queries. The coding sequence of *E. multilocularis* TGR was determined based on the cDNA and amino acid sequences of *E. granulosus* TGR previously described in [11]. To map the coding sequence of *S. mediterranea* TGR, TR and GR, the amino acid sequences of *E. granulosus* and *S. mansoni* TGRs, human mitochondrial TR and human GR, were respectively used, as well as available ESTs encoding *S. mediterranea* TGR, TR and GR, retrieved from <http://smedgd.neuro.utah.edu/>. The final adjustments of intron-exon boundaries were performed based on a multiple alignment of TRs, GRs and TGRs from different species. Finally, in order to obtain information on expressed TR, GR and TGR from other platyhelminth organisms, tblastn (EST others option) and blastp searches were performed at the NCBI server with flatworm and mammalian pyridine-nucleotide thiol-disulfide oxidoreductases as queries.

Sequence and phylogenetic analyses

The selenocysteine insertion sequence (SECIS) elements present in flatworm TR and TGR genes were identified using the SECISearch program, version 2-19 <http://genome.unl.edu/SECISearch.html> [27] using the default energy cutoffs in all cases, and the canonical pattern for all but *Schistosoma* TGRs; in these latter instances the non-canonical pattern revealed the SECIS element. Topology prediction of the polypeptides was carried out using SignalP 3.0 [23] and ipSORT [28]. Phylogenetic analyses of flatworm and mammalian GRs, TRs and TGRs were performed on amino acid sequences aligned with ClustalW2 [29]. Neighbor-joining, maximum parsimony, minimum evolution and UPMGA methods were used to reconstruct phylogenies, with MEGA4 [30], and the following parameters: pairwise deletion, Poisson correction and uniform rates among sites. In all cases bootstrap test was performed on 500 replicates. The analyses were performed including and excluding the Grx domain of TGR.

In silico and experimental identification of platyhelminth TGR variants

To identify platyhelminth TGR variants, tblastn searches were performed on the blast servers of NCBI (option EST others) and *S. mansoni*, *E. multilocularis* and *S. mediterranea* genomic databases using the TGR sequences as queries. Experimental evidence of newly identified *S. mansoni* variants was analyzed using forward primers that allow discrimination of the variants (for details see Figure 5) used in combination with a TGR gene-specific reverse primer in PCR reactions from cDNAs of different *S. mansoni* materials. Total RNA was isolated from cer-

cariae, 3 hours schistosomula and adult worms using a modified TRIZOL (Invitrogen)/RNeasy (Qiagen) procedure [31]. Schistosomula were prepared by mechanical transformation of cercariae as previously described [32]. cDNA was generated from 1 µg of total RNA using Superscript II (Invitrogen) according to manufacturer's specifications. The *S. mansoni* life cycle is maintained at the Schistosoma Research Group at the Pathology Department, University of Cambridge, UK. In the case of *E. granulosus*, experimental identification of additional TGR variants was examined on different parasite materials (larval worms, adult worms and hydatid cyst wall) by RT-PCR, using *E. granulosus* 5' splice-leader exon as forward primer and different gene specific reverse primers derived from the previously published TGR cDNA sequence [11]. In all cases RNA was obtained from trizol-treated *E. granulosus* samples and subsequently used as template for reverse transcription reactions and PCRs as previously described [33], using ThermoScript (Invitrogen) reverse transcriptase and Pfu (Fermentas) DNA polymerase, respectively. In all cases, PCR products were analyzed by electrophoresis.

Amplification of *Echinococcus granulosus* TGR gene and its promoter from genomic DNA

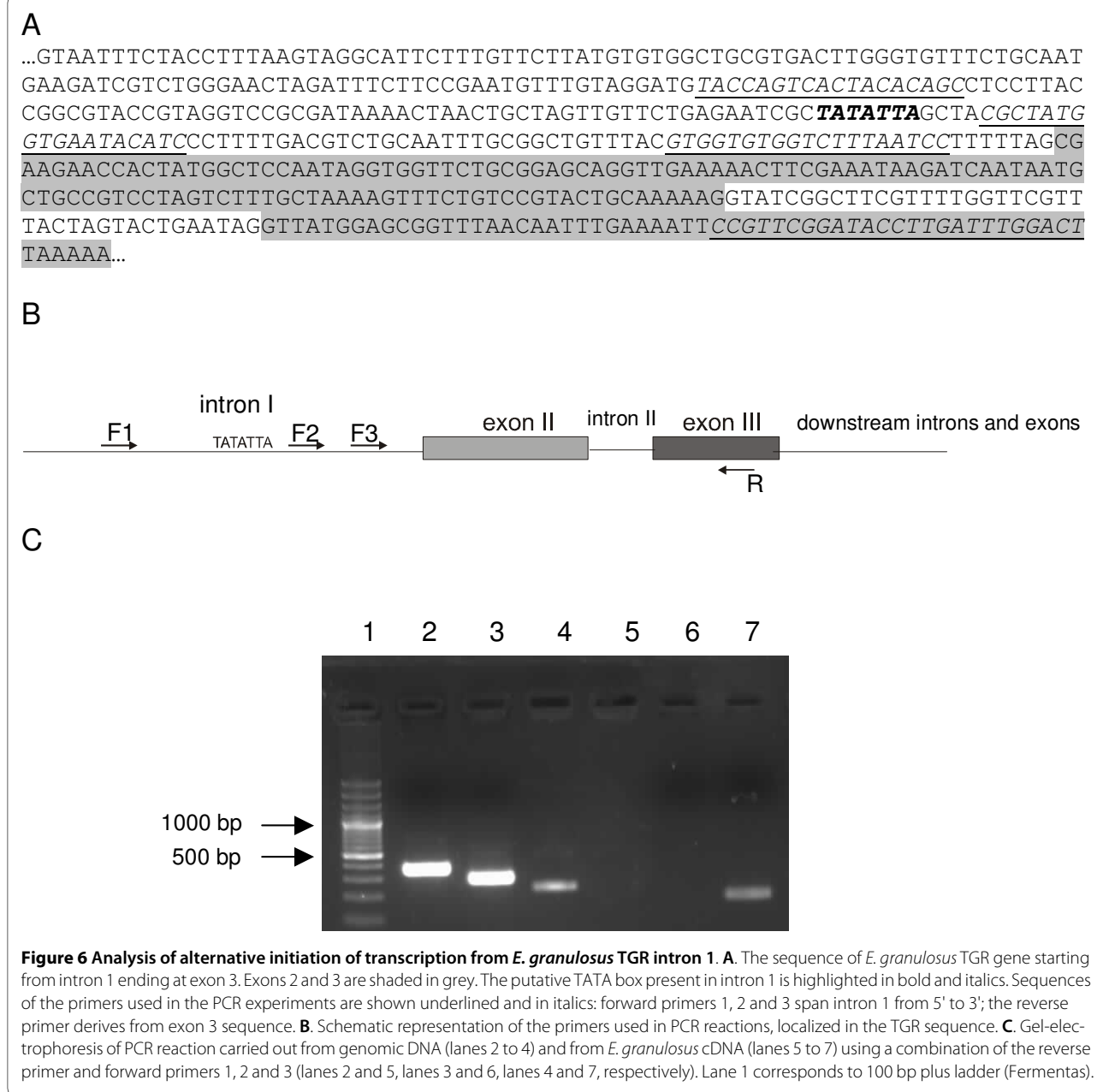
Two fragments corresponding to the Grx domain and the TR domains of the TGR gene were amplified by PCR from *E. granulosus* DNA, using forward and reverse primers derived from *E. granulosus* TGR cDNA sequence. The PCR products were cloned in pGEM-T easy™ (Promega) and subsequently sequenced by primer walking. The core promoter of TGR gene (1.5 kbp sequence upstream of TGR coding sequencer) was amplified using a forward primer derived from *E. multilocularis* homologous region, and a reverse primer derived from the 5'-end of *E. granulosus* TGR cDNA sequence; this PCR product was cloned and sequenced as described above.

Analysis of alternative initiation of transcription from *E. granulosus* TGR gene

In order to investigate and map alternative transcription initiation, we also performed RT-PCRs from *E. granulosus* hydatid cyst germinal layer using forward primers spanning the first intron of *E. granulosus* TGR (upstream and downstream of the presumed TATA box present in intron 1) and a reverse primer corresponding to exon 3 (Figure 6). Control PCRs were carried out from genomic DNA.

Additional material

Additional file 1 *Echinococcus granulosus* and *Echinococcus multilocularis* TGR exon-intron boundaries. Exon-intron boundary sequences of TGRs and canonical donor and acceptor splice sites.



Abbreviations

GSH: Glutathione; GR: glutathione reductase; GSSG: oxidized glutathione; SECIS: selenocysteine insertion sequence; Trx: thioredoxin; TGR: thioredoxin glutathione reductase; TR: thioredoxin reductase; EST: expressed sequence tag.

Authors' contributions

LO carried out most of the work described: cloned *E. granulosus* gene and promoter, performed the experiments using a heterologous system, analyzed the existence of TGR variants in *S. mansoni* and *E. granulosus* cDNAs by PCR, searched the genomes of *E. multilocularis*, *Schistosoma* spp. and *S. mediterranea* and analyzed the gene structure of the genes of interest. AVP prepared mRNA and cDNA from different *S. mansoni* samples. CF and MB prepared cDNA and genomic DNA from *E. granulosus* samples and provided technical help. GS performed *in silico* data mining of ESTs and genomes, phylogenetics analysis, and performed the experiments to test alternative transcription from intron promoter by PCR. LO and GS primarily conceived and designed the study, with insights from CF and VNG. Analyzed all the data: LO, VNG, CF and GS. GS wrote

the manuscript, LO, VNG and CF helped to draft the manuscript. All authors read, revised and approved the final manuscript.

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