Analysis of rhodopsin G protein-coupled receptor orthologs reveals semiochemical peptides for parasite (*Schistosoma mansoni*) and host (*Biomphalaria glabrata*) interplay

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Schistosomiasis is a medically significant disease caused by helminth parasites of the genus *Schistosoma*. The schistosome life cycle requires chemically mediated interactions with an intermediate (aquatic snail) and definitive (human) host. Blocking parasite development within the snail stage requires improved understanding of the interactions between the snail host and the *Schistosoma* water-borne free-living form (miracidium). Innovations in snail genomics and aquatic chemical communication provide an ideal opportunity to explore snail-parasite coevolution at the molecular level. Rhodopsin G protein-coupled receptors (GPCRs) are of particular interest in studying how trematode parasites navigate towards their snail hosts. The potential role of GPCRs in parasites makes them candidate targets for new antihelminthics that disrupt the intermediate host life-cycle stages, thus preventing subsequent human infections. A genomic-bioinformatic approach was used to identify GPCR orthologs between the snail *Biomphalaria glabrata* and miracidia of its obligate parasite *Schistosoma mansoni*. We show that 8 *S. mansoni* rhodopsin GPCRs expressed within the miracidial stage share overall amino acid similarity with 8 different *B. glabrata* rhodopsin GPCRs, particularly within transmembrane domains, suggesting conserved structural features. These GPCRs include an orphan peptide receptor as well as several with strong sequence homologies with rhabdomeric opsin receptors, a serotonin receptor, a sulfakinin (SK) receptor, an allatostatin-A (buccalin) receptor and an FMRFamide receptor. Buccalin and FMRFa peptides were identified in water conditioned by *B. glabrata*, and we show synthetic buccalin and FMRFa can stimulate significant rates of change of direction and turn-back responses in *S. mansoni* miracidia. Ortholog GPCRs were identified in *S. mansoni* miracidia and *B. glabrata*. These GPCRs may detect similar ligands, including snail-derived odorants that could facilitate miracidial host finding. These results lay the foundation for future research elucidating the mechanisms by which GPCRs mediate host finding which can lead to the potential development of novel anti-schistosome interventions.

**Abbreviations**

- GPCRs: G protein-coupled receptors
- HMMs: Hidden Markov Models
- MEGA: Molecular evolutionary genetics analysis
- PBS: Phosphate buffered saline

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Metazoan helminth blood flukes of the genus *Schistosoma* are the primary etiological agents of human schistosomiasis, a disease endemic in 74 countries that affects over 200 million people worldwide. Globally, up to 200,000 people die directly or indirectly due to schistosomiasis annually and an estimated 600 million people live in endemic areas. Schistosomes have a complex dioecious life cycle, involving asexually reproduced larvae in a molluscan intermediate host and sexually reproductive adult worms in the mammalian definitive host. In water, *Schistosoma mansoni* eggs hatch into free-living, mobile miracidia that must search and infect a compatible snail host, *Biomphalaria glabrata*. Following infection, a single miracidium can reproduce asexually via mother and daughter sporocysts, into several thousand cercariae that each when released may develop into an adult worm in the human host. The complex physiological and morphological changes associated with the *Schistosoma* life-cycle means that the anthropogenic control of transmission could be directed at several life cycle stages. Currently, the adult worms are usually targeted through treatment of infected humans with the drug praziquantel. Nevertheless, the disease remains a constant threat in developing countries and the World Health Organization has acknowledged that continued research on the snail infection stage is required. For example, alternative methods of interfering with the *Schistosoma* life-cycle could involve the application of anthelmintic drugs that target the host seeking behavior of miracidia or cercariae.

Schistosome miracidia have restricted vision and limited time (12–16 h) to find an appropriate host snail. Host identification involves a range of behavioral responses that promote host localization, thereby increasing the likelihood of successful infection. Sharing the same living environment with the intermediate host makes the miracidial stage an ideal point to interrupt the parasite’s life cycle and represents a target window for assessing the molecular components required for host finding. Two main behavioural responses occur comprising an initial dispersal and directional phase influenced by photoreceptor mediation, and a secondary seeking and circling phase (chemokinesis) that is olfactory-mediated. To date, little information is available concerning the underlying molecular mechanisms that dictate olfactory-mediated host detection by schistosome miracidia although a peptide was recently discovered from *B. glabrata* that induces behavioural changes in miracidia.

G protein-coupled receptors (GPCRs) are well recognized as chemosensory receptors in eumetazoans, and are a promising research focus for understanding parasite host-finding. GPCRs and downstream biochemical pathways are often used as selective pharmacological targets for parasite lifecycle interruption and, and thus may be effective targets for miracidial manipulation. At the molecular level, there are several reports on *S. mansoni* receptor biology and signal transduction pathways such as the discovery of an array of genome-encoded sensory-type proteins. These include GPCRs that may interact with chemical ligands, a concept supported by proteomic and functional expression analyses that identified GPCRs in the miracidia and adult tegumental matrix of schistosomes. Scrutiny of the *S. mansoni* genome revealed numerous rhodopsin-type GPCR sequences using a combination of three bioinformatic algorithms, including the Phobius, HMMerSearch and Pfam scan. Those expressed in the miracidia included two opsin receptors, which may underpin miracidial photokinetic behavior.

Emerging information shows that the close association of the snail and its obligate schistosome parasite has helped in shaping their respective genomes. The genetic variability of the snail host, rather than the human host, may be a more significant factor in influencing the variability of life history traits in schistosomes. For example, it has been noted that the presence of homologous mucin proteins between the snail and different strains of *S. mansoni* may be key elements underlying snail host-parasite compatibility. This is consistent with the discovery of significant homology of the non-coding 5’ and 3’ regions of non-long terminal repeat retrotransposon nimbus sequences; these class I transposable elements copy and paste themselves into different genomic location, in the host and parasite, suggesting possible horizontal transfer of host sequences into the parasite.

In this study, we identified *S. mansoni* miracidia rhodopsin-like GPCRs that share significant sequence identity with rhodopsin-like GPCRs in *B. glabrata*. This new knowledge guided peptide behaviour bioassays on *S. mansoni* miracidia which demonstrated that FMRFa and buccalin peptides can elicit behaviours consistent with host finding.

**Materials and methods**

**Ethics approval and consent to participate.** The conduct and procedures involving animal experimentation were approved by the Animal Ethics Committee of the QIMR Berghofer Medical Research Institute (project number P242). This study was performed in accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The study was carried out in compliance with the ARRIVE guidelines.

**S. mansoni miracidia collection and transcriptome preparation.** Livers were obtained from ARC Swiss mice infected with *S. mansoni* (Puerto Rican strain) under conditions specified by the Australian Depart-
ment of Agriculture, Fisheries and Forestry (DAFF). Mice were euthanized with CO2 gas and their livers were perfused with chilled PBS. Eggs of S. mansoni were collected during perfusion of mice. Four infected mouse livers were sliced with scalpel blades and blended to a smooth consistency in 50 mL phosphate buffered saline (PBS). The egg pellets were washed using PBS twice on a second Percoll column (2.5 ml Percoll + 7.5 ml 0.25 M sucrose in a 15 mL tube). Purified eggs were transferred into a 200 ml hatching measuring cylinder wrapped completely in light-blocking black tape with the exclusion of the top 4 cm from the lip, thereby producing a light-gradient. The hatching cylinder was topped with pH neutral MilliQ water until ~ 1.5 cm above the tape-covered area and exposed to bright light at 27 °C. Eggs were incubated for 6 h post-hatch, and the top 10 ml of miracidium-containing water was collected for miracidia isolation. Hatched miracidia were isolated by centrifugation at 8,000 × g for 1 min at 4 °C, and washed twice with water. For RNA isolation, miracidia were collected at 6 h post-hatch and kept at -80 °C. Total RNA was isolated from S. mansoni miracidia (6 h post-hatch in triplicate) using TRIzol reagent following the manufacturer's user guide (Invitrogen, USA), the RNA quantity and quality were assessed by UV spectrophotometry (NanoDrop ND-1000) and the RNA was sent to Novogene (Hong-Kong) for next-generation Illumina 2500 platform RNA-seq. Raw sequence data was deposit into the GenBank NCBI under accession number SRR17224866. De novo transcriptome assembly of the S. mansoni miracidia raw sequence data was performed using Trinity, as previously described35,36 and contigs were translated into protein sequences using Transdecoder35. Gene expression levels of the S. mansoni miracidia were calculated by mapping raw sequence data against the S. mansoni reference genome derived from WormBase Parasite (https://parasite.wormbase.org/Schistosoma_mansoni_pjreia36577/Info/Index/) using CLC Genomic Workbench with default parameters37.

**Gene identification and functional annotation.** The pipeline for identification of ortholog GPCRs shared between B. glabrata and S. mansoni is shown in Fig. 1. For B. glabrata, data on GPCRs and their expression levels in different tissues were retrieved from a previous study6. With S. mansoni, transcriptome-derived protein sequences were searched for Pfam-based profiles and TM domains to identify receptors that belong to the rhodopsin GPCR family. Specifically, this included two bioinformatic tools to predict TM domains for all proteins, including TMHMM (http://www.cbs.dtu.dk/services/TMHMM-2.0/) and Phobius (http://phobius.sbc.su.se/). As TM domains are convenient markers for GPCRs, we only focused on those sequences with 7-TM domains. Next, we applied a Pfam-based profile search using HMMerSearch (http://www.hmmer.wustl.edu/). Proteins containing putative rhodopsin-type GPCR domains were systematically identified by profile hidden Markov model searches using the HMMer package (http://www.hmmer.wustl.edu/) and the PFAM model PF00001 (7tm_1). Putative GPCRs identified in S. mansoni miracidia were used to query (using tBLASTp) the B. glabrata GPCR database. Those sequences with E-values < 1.0E-20 were retrieved and screened for the presence of recurrent transmembrane motifs using TMHMM (http://www.cbs.dtu.dk/services/TMHMM/). Those containing 7 transmembrane (TM) domains were selected for further analysis. Multiple sequence alignments were created with Molecular Evolutionary Genetics Analysis (MEGA) software version 6.049 with the MUSCLE49 algorithm. Phylogenetic trees were constructed using the neighbor-joining method with 1000 bootstrap replicates for node support. Gene ontology mapping and functional annotation were applied by using OmicsBox (BioBam)40. The final phylogenetic tree and heatmap were modified with iTOL v541.

**Miracidia behaviour in response to test solutions.** Test solutions. Synthetic FMRFa (FMRF-NH₂), buccalin (RLDRFGFAGQL-NH₂) and SK (NYGDYGIGGGRFGR) were provided by China Peptides (Shanghai, China) (purity ≥95%). Serotonin (5-hydroxytryptamine/5-HT) was provided by Sigma (Burlington, United States).
**Identification of candidate peptide ligands from B. glabrata-conditioned water extract.** To identify whether candidate peptide ligands were present in B. glabrata-conditioned water, two approaches were taken. First, mass spectrometry data derived from prior analysis of B. glabrata-conditioned water was searched using target precursor proteins.

Second, antibody-mediated dot blot analysis was performed using B. glabrata-conditioned water extracts. B. glabrata were washed with pH neutral MilliQ water and placed in beakers containing 20 ml water for 2 h at room temperature (RT). Snails were removed, conditioned water was collected from 20 snails (in different aquaria), and 20 ml methanol was added to each beaker and mixed thoroughly. The conditioned water was filtered through PVDF Millex-HV syringe filter units (0.45 µm) to remove particles and microbes. The filtrate was snap frozen and lyophilized. For negative controls, water not previously exposed to snails was similarly processed. When required for dot blot assay, samples were rehydrated with 200 µl MilliQ water and centrifuged at 12,000 rpm for 5 min. The supernatant was collected and diluted 1:1 in MilliQ water. Quantitation was performed using a NanoDrop for dot bot assay, samples were rehydrated with 200 µl MilliQ water and centrifuged at 12,000 rpm for 5 min.

**Availability of data and materials.** The S. mansoni miracidial transcriptomic raw sequence data was deposited into the GenBank NCBI under accession number SRR17224866 (https://www.ncbi.nlm.nih.gov/sra/?term=SRR17224866). The S. mansoni reference genome is available from WormBase Parasite (https://parasite.wormbase.org/Schistosoma_mansoni_prjea36577/Info/Index/). The original B. glabrata protein dataset is available from the VectorBase(https://vectorbase.org/vectorbase/app/downloads/Current_Release/BglabrataBB02 fasta/data/).

**Results**

**Identification of ortholog GPCRs shared between B. glabrata and S. mansoni miracidia.** In total, 96 proteins with 7-TM domains were extracted from the S. mansoni transcriptome-derived protein models based on TMHMM prediction. Phobius prediction led to the identification of 139 proteins with 7-TM domains. PLAM profiling of both predictions led to the classification of 87 proteins (E-value < 0.0004) as rhodopsin-type receptors. BLASTp analysis of these GPCRs against all B. glabrata rhodopsin GPCRs showed significant matches (E-value < 1.0E−20) for 8 GPCRs (Table 1), with between 26 to 48% amino acid identity (File S2a). BLASTp searches using target precursor proteins.

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validated by miracidial behaviour changes in the presence of snail FMRFamide and AST-A/buccalin peptides. These include GPCRs with similarity to neuropeptide GPCRs that bind FMRFa, AST-A/buccalin and sulfakinin within the snail mucus, while in silico analysis from snail-conditioned-water proteins predicted biomolecule(s) has not been clearly defined. One study implicated “miracidia-attracting glycoproteins” present within the water. The latter expectation was of snail-derived biomolecules influence S. mansoni miracidia. The 5-HT is a well-established eumetazoan neurotransmitter, while and B. glabrata neuropeptides FMRFa, buccalin [recognised as AST-A homolog in molluscs] and SK have previously been identified in B. glabrata, so were tested for bioactivity on miracidia. The trajectories of miracidial movement before and after addition of the FMRFa and buccalin peptides (2 μl at 100 μM) are compared in Fig. 3A,B. Representative movies can be viewed in Movies S1 and S2. Before addition, miracidia generally swam across the FOV in a direct, linear path (Fig. 3A,B—Before). Following application, miracidia showed localized movement within the FOV, as well as increased circular swimming (Fig. 3A,B—After). Upon application of buccalin or FMRFa peptides, miracidia within the FOV swam for longer, except for buccalin at 10 μM (P-value = 0.2964) (Fig. 3C). The change in acceleration were significant after the addition of buccalin or FMRFa (100 μM and 10 μM) (Fig. 3D). Peptides stimulated a swimming pattern concentrated around the location of the peptides. In contrast, following application of 5-HT (5 nM; Movie S3), the time within the FOV was insignificant, yet the change of acceleration was significant (File S3). Peptides (100 μM) resulted in miracidia staying longer in the region where the peptide was added, but there was no significant change in average velocity, as indicated by twoway ANOVA analysis (File S4). There was also no significant change in the time within FOV, nor in average velocity, upon addition of the SK peptide at 3 different concentrations (100 μM, 10 μM and 1 μM; Movie S4). As negative controls, MilliQ water was tested on S. mansoni miracidia and no behavior changes was observed.

Detection of buccalin and FMRFa peptides in B. glabrata-conditioned water. The B. glabrata buccalin precursor contains numerous buccalin-like peptides (Fig. 4A). The most conserved region in the buccalins is a C-terminal FXGGIG, which following post-translational processing, becomes an amidated peptide. Dot blots performed on conditioned-water extracts from B. glabrata showed the presence of a buccalin-like peptide at extract dilutions from 10 μg to 0.1 μg (Fig. 4B). The B. glabrata FMRFa precursor contains numerous FMRFa related peptides (FaRPs), including FMRFa, FLRFa and FIFRa (Fig. 4C). Analysis of proteomic mass spectrometry data derived from naïve B. glabrata snail-conditioned water, identified 4 peptide matches for the FMRFa precursor, although not specifically within any FaRPs. Dot blots performed on conditioned-water extracts from B. glabrata showed the presence of peptide(s) with similarity to FaRPs at extract dilutions from 10 μg to 0.1 μg (Fig. 4D).

Discussion

S. mansoni miracidia respond to snail-derived biomolecules, although the precise identity of the active biomolecule(s) has not been clearly defined. One study implicated “miracidia-attracting glycoproteins” present within the snail mucus, while in silico analysis from B. glabrata snail conditioned-water proteins predicted interactions of uncharacterized S. mansoni proteins with B. glabrata proteins. Peptides have also been implicated, whereby a snail-derived novel peptide (named P12) stimulated changes in the behaviour of the S. mansoni miracidia.

In this study, to narrow down biomolecules potentially involved in the parasite and host interplay, we utilised gene resources from both the B. glabrata and S. mansoni to identify ortholog GPCRs that are likely used by each organism to detect similar ligands. We reported that S. mansoni miracidia GPCRs share significant identity with B. glabrata GPCR orthologs with putative ligands presented an opportunity to investigate how snail-derived biomolecules within the FOV , as well as increased circular swimming (Fig. 3A,B—After). Upon application of buccalin or FMRFa (100 μM and 10 μM) (Fig. 3D). Peptides stimulated a swimming pattern concentrated around the location of the peptides.

| S. mansoni ID | B. glabrata ID | E-value | Overall identity (%) | TM number (% identity) | Best match GPCR |
|---------------|---------------|---------|----------------------|------------------------|-----------------|
| Smp_140620    | BGLB028445    | 2.78E-50| 26.8                 | 1 (45.5), 2 (64.0), 3 (58.3), 4 (40.0), 5 (39.1), 6 (45.8), 7 (65.2) | FMRFa           |
| Smp_131980    | BGLB002561    | 2.88E-42| 30.5                 | 1 (40.0), 2 (40.0), 3 (76.2), 4 (50.0), 5 (68.4), 6 (70.8), 7 (69.6) | Orexin type 2   |
| Smp_173010    | BGLB013427    | 2.67E-23| 48.5                 | 1 (45.6), 2 (60.0), 3 (45.8), 4 (52.0), 5 (39.1), 6 (50.0), 7 (63.2) | Sulfakinin      |
| Smp_007070    | BGLB003586    | 2.69E-60| 31.2                 | 1 (29.2), 2 (41.7), 3 (42.9), 4 (27.8), 5 (36.8), 6 (56.5), 7 (60.9) | Neuropeptide F  |
| Smp_126730    | BGLB013877    | 9.83E-37| 34.5                 | 1 (27.3), 2 (26.1), 3 (61.9), 4 (66.7), 5 (34.8), 6 (60.9), 7 (60.9) | Serotonin       |
| Smp_180030    | BGLB032600    | 4.18E-20| 24.9                 | 1 (52.2), 2 (68.2), 3 (74.9), 4 (52.6), 5 (38.1), 6 (66.7), 7 (85.0) | Opsin           |
| Smp_203500    | BGLB004467    | 2.08E-44| 30.1                 | 1 (47.8), 2 (34.8), 3 (37.0), 4 (34.8), 5 (47.6), 6 (62.5), 7 (58.3) | Allstatatin A/buccalin |
| Smp_049330    | BGLB001538    | 1.44E-59| 36.8                 | 1 (42.8), 2 (54.1), 3 (32.8), 4 (39.4), 5 (41.6), 6 (64.5), 7 (54.7) | Rhodopsin       |

Table 1. Comparative sequence identity between Biomphalaria glabrata and Schistosoma mansoni GPCR homologs.
The AST-A and its receptor have been characterised in various insects where they are involved in multiple functions such as inhibition of juvenile hormone biosynthesis and reduction of food intake, AST-A-like neuropeptides have been identified in gastropods and bivalve molluscs, including *Lottia gigantea*, *Theba pisana*, *Aplysia californica* and *Crassostrea gigas*. Buccalin, named following its first identification in the accessory radula closer muscle of *A. californica*, has been implicated in various activities in molluscs such as the inhibition of muscle contraction, regulation of feeding and spawning. Also in gastropods and bivalves, the AST-A/buccalin receptor was identified through in silico analysis of publicly available genomic datasets including that of *B. glabrata*. In our study, we identified an AST-A/buccalin receptor ortholog in *S. mansoni*, although there are no reports that *S. mansoni* has a buccalin-like peptide. In fact, a comprehensive neuropeptide investigation of 10 platyhelminth species showed that only the free-living turbellarian *Macrostomum lignano* has a buccalin-like peptide (*npp-9* gene; GAYSGFLG). We identified a buccalin-like peptide in the *B. glabrata* conditioned water. Despite the presence of neuropeptides in mucus secretions having not been well investigated, we previously identified neuropeptides (including buccalin) within the salivary gland mucus of the land snail *T. pisana*. The FMRFa was first discovered in the hard clam *Mercenaria mercenaria* and it is thought to have a pleitropic role in molluscan physiology. Extensive studies performed on the freshwater snail *Helisoma* showed that FMRFa and related peptides are densely concentrated not only in the nervous system but also within the salivary glands.

Figure 2. Characterization of GPCR orthologs shared between *B. glabrata* and *S. mansoni* miracidia. (A) Phylogenetic tree analysis of sharedGPCRs, where each *B. glabrata* GPCR clusters with an ortholog receptor from *S. mansoni*. Bootstrap values support the confidence levels of clades. (B) Heatmap showing expression of shared GPCR- (in TPM) encoding genes in *S. mansoni* miracidia at 6 h post-hatch and different *B. glabrata* tissues. The columns represent *S. mansoni* miracidia biological replicates (1–3) and their mean, as well as *Biomphalaria* (glabrata) tissues.
glands. An FMRFa receptor has been identified in the heart and nervous tissue of the land snail *Helix* and the optic lobe membrane of the squid *Loligo pelei*. The *S. mansoni* genome contains a gene encoding a FLP precursor (*npp-13* gene) that may be processed to release two RFamide peptides (HFMPQRFa and YTRFVPQRFa). A synthetic FLP (GNFFRFa) derived from non-schistosome platyhelminth precursors stimulates contraction of *S. mansoni* muscle fibres in vitro. An FLP receptor has also been reported in the turbellarian flatworm *Girardia tigrina* based on sequence similarity and a receptor calcium mobilization assay. The homolog receptor in *S. mansoni* miracidia was investigated in the current study due to its similarity with the *B. glabrata* FMRFa receptor.

Figure 3. Miracidial behaviour assay using buccalin and FMRFa peptides. (A) Representative trajectories of miracidial movement before and after the addition of the buccalin solution (100 µM) to the center of the recording area. Each colour represents one indistinguishable miracidium individual. See Movie S1 for representative assay video. (B) Representative trajectories of miracidia movement before and after the addition of the FMRFa solution (100 µM) to the center of the area. Each colour represents one indistinguishable miracidium individual. See Movie S2 for representative assay video. (C) Graph of time duration of miracidia remaining in the videoing zone, and (D) mean acceleration values, before and after the addition of buccalin and FMRFa.
Our behaviour assays also indicated that snail-derived FMRFa can be detected by *S. mansoni* miracidia due to their staying significantly longer in FOV and the increased acceleration of miracidia, supported by the observed presence of FMRFa precursor peptides in *B. glabrata* conditioned water. However, as *S. mansoni* has the potential to generate endogenous FLPs, we cannot preclude the possibility that the applied FMRFa may stimulate endogenous effects, leading to the observed miracidial behaviour changes.

The monoamine 5-HT plays a critical role in neural transmission and has been very well documented throughout eumetazoans, as has the conservation of 5-HT GPCRs. In adult *S. mansoni*, 5-HT stimulates motor activity\(^6^9\), while in the miracidia, an immunofluorescent approach localized 5-HT to within sensory nerves\(^7^0\). The 5-HT GPCR was identified within our interspecies GPCR ortholog analysis yet we found that 5-HT at 5 mM did not modify miracidial behaviour, while the significant change in acceleration could be attributed to its high concentration.

Sulfakinin is a sulfated neuropeptide best known for its function as a satiety (food intake) factor\(^7^1\). In silico data-mining showed that molluscan SK has the C-terminal RF(W)amide sequence common to insect sulfakinins, as well as the DY motif shared by both insect SKs and vertebrate cholecystokinin (CCK)\(^7^2\). Since vertebrate CCKs and insect SKs reveal similar biological function relating to digestive enzyme secretion, satiety and smooth muscle contraction\(^7^3\), it is possible that their molluscan counterparts have retained similar basic biological activities. In contrast, there is no obvious SK in *S. mansoni*, suggesting that the parasite may only recognize the *B. glabrata* SK, either as a secreted semiochemical, or once it penetrates the snail as a guidance peptide to navigate to the hepatopancreas where it proliferates\(^7^4\). Our behavior assays demonstrated that SK did not alter miracidial behavior (neither the velocity nor duration present under FOV were affected), and therefore it is more likely to act as an internal stimulus in *S. mansoni*.

FMRFa and buccalin peptides may contribute to a cocktail of biomolecules that could be used as an effective, species-specific attractant. Our serial dilution assays suggested sustained bioactivity for both buccalin and FMRFa peptides at a concentration of at least 1 µM. We also report 1 orphan peptide GPCR ortholog within *B. glabrata* and *S. mansoni* miracidia, which is consistent with the possibility that uncharacterized species-specific peptides could help attract miracidia to the appropriate snail host due to its presence in many tissues of *B. glabrata* and its high expression level in *S. mansoni* miracidia.

**Conclusions**

To minimise transmission and reduce schistosomiasis prevalence, interference with the snail-miracidium interaction is a promising plan of biocontrol. We characterised ortholog GPCRs shared between *B. glabrata* and *S. mansoni* miracidia, important biomolecules commonly used for chemosensory communication. The *B. glabrata* buccalin and FMRFa GPCRs represented good targets for bioassay, the results from which indicated that buccalin and FMRFa stimulated miracidial behaviour changes, despite the fact that homologs of buccalin-like peptides are not present in *S. mansoni*. These GPCRs could present novel targets for the development of anti-helminthic compounds to be applied to lakes and specifically interfere with Schistosoma detection of snail host. For greater species-specificity, we suggest that deorphanizing the ortholog orphan peptide GPCR will be most advantageous. These findings further help our understanding of chemosensory interaction between parasites and their hosts, particularly within aquatic environments.
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Author contributions
Conceived and designed the study and supervised the project: S.F.C. and T.W. Undertook the study and data analysis: P.P., D.L., C.F., M.Z., and M.D. Contributed to analysis using various tools: D.L., M.Z., and T.W. Wrote the paper: P.P., D.L., R.W., D.P.M., and S.F.C. All authors read and approved the final version of the manuscript.
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