Nanoemulsion of *Sideroxylon obtusifolium* as an Alternative to Combat Schistosomiasis

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Schistosomiasis is caused by the intestinal parasite *Schistosoma mansoni*. Individuals are affected by schistosomiasis when they are exposed to aquatic environments contaminated with *Schistosoma* cercariae that emerged from the infected intermediate host mollusk of the genus *Biomphalaria*. The WHO recommends using molluscicidal products to reduce the snail population and disease transmission. The WHO encourages the search for alternative substances in schistosomiasis control. Natural products are seen as a promising alternative because they are abundant in countries where schistosomiasis is endemic and have many different substances in their extracts, impairing cases of resistance. Therefore, the nanoemulsion effect of a butanol-soluble fraction of *Sideroxylon obtusifolium* leaves was evaluated against three study points in the biological cycle of the disease, that is, adults and young *Biomphalaria glabrata*, spawning by the host mollusk, and infectious larvae of the parasite. Extract-SOB (butanol fraction) and nano-SOB (nanoemulsion) demonstrated promising activity in adult *B. glabrata* population control with an LC50 of 125.4 mg/L, an LC90 of 178.1 mg/L, an LC50 of 75.2 mg/L, and an LC90 of 97 mg/L. Nano-SOB presented greater potency against young *B. glabrata*, with an LC90 of 72.1 mg/L and an LC50 of 58.3 mg/L. Still, relevant activity against *S. mansoni* cercariae was eliminated in 4 h (LC90: 34.6 mg/L).

Nano-SOB reduced viable spawning by approximately 30% at 178.1 and 97 mg/L. Referring to most substances in this extract, quercetin-3-rhamnosyl-(1-6)-galactoside and hyperoside may cause low environmental toxicity and human toxicity according to in silico analysis. Thus, nano-SOB is a promising agent to combat *B. glabrata* population growth and schistosomiasis transmission.

**Keywords:** nanoemulsion, molluscicide, *Biomphalaria*, *Sideroxylon obtusifolium*, plant extract

**INTRODUCTION**

Schistosomiasis is an acute parasitic and chronic disease caused by worm trematodes of the genus *Schistosoma* and is transmitted to many snail types. This disease affects 78 countries, mainly in tropical and subtropical regions. Schistosomiasis is the second largest infectious parasitic disease in the world after malaria (WHO, 2020a). In 2019, the estimates showed that at least 236.6 million...
people needed treatment for schistosomiasis. At least 105.4 million people in the world were treated for schistosomiasis (WHO, 2020b).

Schistosomiasis is caused by the intestinal parasite *Schistosoma mansoni*, which uses mollusks of the genus *Biomphalaria* as an intermediate host, favoring the transmission cycle (World Health Organization, 1983). The individuals were affected by schistosomiasis when they were exposed to aquatic environments contaminated with *Schistosoma* cercariae that emerged from the infected intermediate host. Therefore, in the installation of the evolutionary cycle of *S. mansoni*, the presence of susceptible *Biomphalaria* is necessary. Furthermore, *Biomphalaria glabrata* is the most efficient intermediate host of *S. mansoni* due to a high degree of susceptibility of this parasite (Rey, 2010). Therefore, without a compatible mollusk, transmission is restricted (WHO, 2019a,b).

In combination with other interventions, the WHO recommends using molluscidical products, with a strategy of focal or seasonal application in freshwater areas and their margins. Thus, the molluscidical application reduces the transmission rate by reducing the mollusk population (BRASIL, 2014; WHO, 2019a,b).

Niclosamide (Bayluscide®, Bayer, Leverkusen, Germany) is a commercially available molluscicide recommended by the WHO (World Health Organization, 1983; BRASIL, 2014) for a large-scale use in schistosomiasis control programs. However, this drug causes toxicity to other organisms, and mollusk populations exhibit resistance to niclosamide®. Thus, the search for new drugs and substances to control the vector is necessary. Therefore, the WHO encourages the search for alternative substances to control schistosomiasis (Inobaya et al., 2014). Natural products are seen as a promising alternative because they are abundant in countries where schistosomiasis is endemic and have many different substances in their extracts, making resistance difficult (Cantanhede et al., 2010; Ding-Feng, 2010).

Numerous plant species have been evaluated, some of which have been recognized for their molluscidical activity. However, there are no data in the literature on this activity with *Sideroxylon obtusifolium* (Humb. Ex Roem. & Schult.) T.D. Penn. (Sapotaceae).

*Sideroxylon obtusifolium*, also called “quixaba,” “sapotiaba,” and “ruptures gibbon” (Silva et al., 2012), is a medicinal tree species found in the Restinga and Caatinga areas. This species can reach up to 15 m. This plant is widely known and is used in popular medicine (anti-inflammatory and healing) and commercial herbal products (Albuquerque and Oliveira, 2007). In addition, *S. obtusifolium* possesses confirmed biological activities, such as antifungal (Pereira et al., 2016), antimicrobial, and anti-inflammatory (Aquino et al., 2016; Carnevale Neto et al., 2017) activities.

The species of the Sapotaceae family are characterized by the diversity of substances resulting from their secondary metabolisms, such as triterpenes, steroids, tannins, flavonoids, and polyphenols, in addition to alkaloids, carotenes, cyanogenic compounds, carbohydrates, and fatty acids (Beltrão, 2000; Oliveira et al., 2012, 2013). Some chemical classes of natural origin have been described to have biological activity, especially flavonoids. Therefore, developing stable carrier systems to increase aqueous dispersibility and bioavailability can reduce this inconvenience. In this context, nanotechnology presents a safe alternative capable of releasing hydrophobic active substances in an aqueous medium and allows stable and adequate systems to be obtained for therapeutic use. Among several options for drug nanocarriers, nanoemulsions (NEs) were chosen for this work (Donsì and Ferrar, 2016).

Most NEs are nanosized (20–200 nm range) oil-in-water droplet dispersions stabilized by surfactants (Sagalowicz and Leser, 2010; Jaiswal et al., 2015). The main advantages of these nanocarriers are enhanced solubility and bioavailability, improved pharmacological activity and stability, more sustained delivery, and protection from physical and chemical degradation (Campos et al., 2012).

Thus, we aimed to evaluate the effectiveness of a nanoemulsion produced with the extract-SOB of *S. obtusifolium* leaves, demonstrating this activity at three points in the biological cycle of the disease as follows: adults and young *B. glabrata*, spawning of the host mollusk, and infectious larvae of the parasite (*S. mansoni* cercariae).

**METHODOLOGY**

**Collection, Identification, and Extract Preparation**

The *S. obtusifolium* leaves were collected in Restinga de Jurubatiba National Park, located in the municipality of Carapebus in Rio de Janeiro, with the authorization for activities with scientific purpose number 13659-14 (SISBIO) and SISGEN authorization number A0D648D. Prof. Dr. Marcelo Guerra Santos identified and deposited the exsiccate (no. 2152) at the Herbarium of the Faculty of Teacher Education at the State University of Rio de Janeiro (RFFP).

**Extract Preparation**

The collected leaves (600 g) were dried in an oven at 40°C with forced ventilation for 48 h. Once dried, this vegetable material was crushed in a hammer mill and was subjected to extraction by maceration at room temperature using 96% GL ethanol as a solvent. This procedure was followed by filtration and concentration in a rotary evaporator, thus obtaining the crude extract (149 g). Then, liquid/liquid partitions were performed for the previous purification of this ethanolic extract of the tris way: 112 g of this ethanolic extract was resuspended in 90% ethanol and was partitioned with solvents of increasing polarity: hexane (14.6 g), dichloromethane (7.6 g), ethylacetate (17.6 g), and finally butanol to obtain 10.0 g of extract-SOB (Oliveira et al., 2012). The extract-SOB was subsequently diluted with dimethylsulfoxide (DMSO) at 50%. Dilutions were made with distilled water to realize the experiments.

**Obtaining the Nanoemulsion – Oil/Water**

Nanoemulsions were developed using an oil and a pair of surfactants previously selected, according to the methodology.
proposed by Azeem et al. (2009), with some adaptations by Falcão (2007) described as follows: purified water (60.0%), extract-SOB of S. obtusifolium (5.0%), transcutol® (22.5%) as the oil phase solvent and an aqueous solution of PEG-40 with hydrogenated castor oil (6.25%) as the chosen surfactant, and propylene glycol as the cosurfactant (6.25%).

The O/W nanoemulsion was prepared by adding extract-SOB, PEG-40 hydrogenated castor oil and propylene glycol in Transcutol® and dissolving it, under manual stirring, during heating in an electric plate until reaching 70–80°C. After reaching the desired temperature, the aqueous phase was gently poured for 30 s. Then, both phases were mixed by the inversion method under mechanical agitation using a four-blade propeller with 400 turns/min speed, by remaining under constant stirring of 400 turns/min at an ambient temperature for 10 min. Finally, the outside of the container was subjected to a cold-water bath. The O/W nanoemulsion obtained in this way, through a low-energy emulsification technique, occurs in the form of nanometric droplets, whose average diameter, according to the steps above, is between 1 and 200 nm.

**Stability Study**
The stability of all emulsions was evaluated 1, 7, 15, 30, and 60 days after manipulation by macroscopic and microscopic analyses (color, visual aspect, phase separation, creaming, and sedimentation) (Falcão, 2007). All emulsions were maintained at room temperature (25 ± 2°C) in a screw-capped glass test tubes during this period.

**Molluscicidal Test**
The molluscicidal assay was performed according to Santos et al.’s (2017) methodology adapted from the WHO (1965). In this methodology, 24-well plates are used, where mollusks of the species B. glabrata (free of S. mansoni) are placed individually in the wells of the plate, forming groups of three mollusks per concentration to be tested. The tests were performed in triplicate on different days, totaling nine mollusks per concentration at the end of the tests. The usual test volume is 2,000 µL of test solution for 10–12 mm B. glabrata mollusks, which are satisfactorily stored during this period.

The exposure was executed on day 0, the viable embryos were counted, and the viability was evaluated at 24 and 48 h. The mortality parameters were color change and/or condensed appearance of the embryo.

**Carcicidal Test**
According to the 24-well plate methodology by Santos et al. (2017), in one well of a 24-well plate, 1 mL of the cercariae suspension was placed in contact with 1 mL of the nanoemulsion to obtain a volume of 2 mL per well, and 20 µL of the vital dye trypan blue 0.1% was added. The following concentrations were tested: 40, 60, 80, 100, 125, and 250 mg/L. Mortality was assessed every hour until the treatment was completed for 4 h using a stereoscope microscope with an 11.5 × magnification to observe the dead cercariae stained blue. The live cercariae did not present a color. The positive control was 1% niclosamide® (NCL), and the negative controls were chlorinated water and nanoemulsion without the active compound.

**Ovicidal Test**
Biomphalaria glabrata spawns are used with embryos from 24 to 72 h of life. The spawns were stored in the wells of a 24-well plate according to the methodology of Santos et al. (2017). We exposed 1 mL of nano-SOB, both in the concentration of CL90 obtained from adult B. glabrata tests. The positive control was 1% niclosamide® (NCL), and the negative controls were chlorinated water and nanoemulsion without the active compound. The exposure was executed on day 0, the viable embryos were counted, and the viability was evaluated at 24 and 48 h. The mortality parameters were color change and/or condensed appearance of the embryo.

**In silico Assays**
The prediction of the ecotoxicity profile of the major substances from the extract-SOB, namely, quercetin-3-rhamnosyl-(1-6)-galactoside and hyperoside, was performed by ADMET Predictor™ (version 9.5, Simulations Plus, Lancaster, CA, United States). The endpoints analyzed were bioconcentration, biodegradation, impact at three different trophic levels [Tetrahymena pyriformis, water flea (Daphnia), and fathead minnow], and effects due to interference with sex hormones (estrogen and androgen disruptors).

**Statistics**
For statistical verification, linear regression analysis and two-way ANOVA were used, with a value of $p < 0.0001$, applying GraphPad Prism software (version 5), showing $R^2 > 0.800$. The LC50 and LC90 values were calculated in Excel version “W” using the dispersion analysis formula.
RESULTS AND DISCUSSION

Nanoemulsion

Several emulsions were prepared with extract-SOB. The optimized process parameters found in the preliminary study were applied as the concentration of the extract-SOB and the surfactant couple (data not shown). Nano-SOB also demonstrated satisfactory characteristics for nanoemulsions, with an average particle diameter of 21.2 nm and stability after 60 days (Table 1). Therefore, it was possible to evaluate the polydispersity and microscopic aspects of the parameters. Due to their characteristic size, they possess stability against sedimentation, creaming, flocculation, or coalescence. This long-term physical stability of nanoemulsions makes them unique, and they are sometimes referred to as “approaching thermodynamic stability” (Solans et al., 2005).

According to Oliveira et al. (2012), from extract-SOB, a total of four saponins and ten flavonol glycosides were isolated and identified. However, these substances are characterized by great structural diversity and need a stable carrier system to increase aqueous dispersibility and bioavailability, which was developed in this study.

The major substances in this extract are quercetin-3-rhamnosyl-(1-6)-galactoside (47.2 mg; tR 16.0 min), which was purified by preparative HPLC from Fr. 117–128 (0.300 g) with a gradient of (MeCN/0.1% aq. HCOOH 40:60–50:50 in 40 min), and hyperoside (619.5 mg; tR 17.0 min), which was purified by preparative HPLC from Fr. 141–160 (0.400 g) with a gradient of (MeCN/0.1% aq. HCOOH 40:60–50:50 in 40 min) according to the work of Oliveira et al. (2012), who used a fraction of the material used in the present study.

Hyperoside is a flavonol glycoside, also known as quercetin-3-O-D-galactoside, in various vegetables and fruits (Wu et al., 2007; Hosseinimehr et al., 2008). Identified in different plant products, it has become important due to its diverse potent pharmacological activities, including anti-inflammatory, antithrombotic, antidiabetic, antiviral, antifungal, hepatoprotective, and antioxidant protective effects (Raza et al., 2017).

Biological Activity Tests

According to documents from the WHO (World Health Organization, 1983; WHO, 2017, 2019a), a molluscicide originating from the plant is considered promising when this substance reaches mortality in 90% of the population at a concentration of 100 mg/L (LC$_{90}$) in 48 h.

We tested the butanol-soluble fraction derived from a crude ethanol extract of $S$. obtusifolium leaves (extract-SOB) against adult $B$. glabrata snails (10–12 mm in length). The results showed maximal activity of extract-SOB in the first 24 h with a maximal concentration of 200 mg/L, maintaining the result in 48 h. The treatment with extract-SOB promoted the LC$_{50}$ of 125.4 mg/L and the LC$_{90}$ of 178.1 mg/L (Figure 1A), in this case exceeding the WHO-recommended value (100 mg/L). On the other hand, niclosamide® (NCL) treatment at 2 mg/L caused 100% death after 24 h, and H$_2$O or nanoemulsion control did not cause toxic effects (Figure 1B).

Nano-SOB enhances the aqueous dispersibility and extracts yield as an optimization method. The molluscidal test of nano-SOB showed maximal activity at 125 mg/L after 24 and 48 h of analysis. The lethal concentrations obtained were

![FIGURE 1](image1.png) The molluscicidal test on adult Biomphalaria glabrata for 48 h using extract-SOB (A,B). The test was performed in triplicate on different days using 27 specimens/triplicate. These data are expressed as the mean ± SD. Using the linear regression analysis, the value of $R^2$ is determined as 1.00.
$LC_{50} = 75.2 \text{mg/L}$ and $LC_{90} = 97 \text{mg/L}$ when tested on adult $B. \text{glabrata}$ (Figure 2A). Niclosamide® (NCL) treatment at 2 mg/L caused 85% mortality after 24 h and 100% mortality after 48 h. The $H_2O$ or nanoemulsion control did not cause a toxic effect (Figure 2B).

Nano-SOB was evaluated against young $B. \text{glabrata}$ of two sizes (3–5 and 4–6 mm). The results were similar after 48 h of exposure, with a maximal activity at 100 mg/L. Interestingly, nano-SOB treatment was more potent in causing a toxic effect in both early stages when compared with adult snails. The $LC_{50}$ value was 72.1 mg/L and the $LC_{90}$ was 58.3 mg/L for $B. \text{glabrata}$ from 6 – 8 mm (Figure 3A) and 3 – 5 mm (Figure 3C). On the other hand, niclosamide® (NCL) treatment at 2 mg/L caused 100% mortality after 24 h, and $H_2O$ or nanoemulsion control did not cause toxic effects (Figures 3B,D).

Nano-SOB exhibited molluscicidal activity against adult and young snails. Thus, we investigated the effect on cercariae released from infected $B. \text{glabrata}$. We treated cercariae for 4 h and analyzed viability each hour. Niclosamide® treatment at
2 mg/L caused 80% mortality after 4 h. Treatment for 1, 2, and 3 h produced toxicities of 30, 46, and 60%, respectively. H2O or nanoemulsion control caused a discrete toxic effect of 10% in 4 h. Thus, nano-SOB exhibited a dose-dependent effect (Figure 4). The nano-SOB concentration of 80 mg/L promoted a maximal activity of 80% after 1 h of exposure. The concentration of 40 mg/L reached this same maximal effect after 4 h of exposure. The LC50 value calculated for the nano-SOB treatment after 4 h was 20.09 mg/L and the LC90 value was 23.63 mg/L (Table 2).

Other studies, such as Vinaud et al. (2005), found maximal cercaricidal activity in 1 h at 20 to 200 mg/L concentrations. Another possibility of controlling the snail population is the control of spawning to influence the mollusk reproductive cycle. The nano-SOB LC50 concentration calculated for adult B. glabrata was determined after exposure to embryos for 24 and 72 h (Figure 5). Treatment with 2 mg/L NCL caused a complete reduction of viable spawning in 24 h. The nanoemulsion control did not cause toxicity. Nano-SOB treatment diminished the viable spawning number. However, the nano-SOB effect was poor when compared with that of NCL. Luna et al. (2005) found 100% mortality of spawning at 100 mg/L Caesalpinia echinata (Pau-Brasil). Nano-SOB, tested in the present study, reduces viable spawning within 48 h of exposure. We tested nano-SOB on another snail species, Melanoides sp., related to schistosomiasis transmission and environmental toxicity. The LC50 and LC90 values of nano-SOB were tested for 72 h (LC50: 75.2 mg/L and LC90: 97 mg/L). The NCL LC50 and LC90 values were obtained from Giovanelli et al. (2002). NCL LC90 promoted maximal Melanoides sp. toxicity after 48 h of exposure. The nanoformulation control did not cause a toxic effect in this snail (Figure 6). The LC50 and

### Table 2 | Kinetics of nano-SOB toxicity on Schistosoma mansoni cercariae.

| Time of exposition (h) | Nano-SOB LC50 (mg/L) | Nano-SOB LC90 (mg/L) |
|------------------------|----------------------|----------------------|
| 1                      | 57.49                | 80                   |
| 2                      | 55.11                | 77.79                |
| 3                      | 36.60                | 55.14                |
| 4                      | 20.09                | 23.73                |

FIGURE 4 | Nano-SOB activity against Schistosoma mansoni cercariae for 4 h (A,B). The test was carried out in triplicate on different days using a range of 80 cercariae per well during the testing of the samples. These data are expressed as the mean ± SD. Using the two-way ANOVA test, the value of R2 is determined as 0.8660, with a P-value of < 0.0001.
LC$_{90}$ values of nano-SOB exhibited a moderate toxic effect on this species compared with NCL. This result could indicate a possible selectivity of the nanoformulation according to the species studied and indicate a possible environmental tolerance compared with NCL.

Evaluation of the environmental effects caused by commercial chemicals may prevent accidental toxic effects on aquatic species. Therefore, the chemical industry, pharmaceutical companies, and government agencies have established standard experimental protocols to test chemicals for their toxic potential (Zhu et al., 2008).

Although experimental protocols for toxicity testing have been developed for many years, computational chemical toxicology is a viable approach to reduce the number of efforts and the cost of experimental toxicity assessment. In this regard, we used the ADMET Predictor (Simulation Plus) to predict the environmental effects caused by the majority of substances from the extract-SOB. Thus, we evaluated the ecotoxicity of (1) quercetin-3-rhamnosyl-(1-6)-galactoside and (2) hyperoside (Scheme 1).

The bioconcentration factor (BCF) is the ratio of the chemical concentration in biota to that in water at a steady state (Hamelink and Spacie, 1977). Environmentally, the BCF describes the accumulation of pollutants partitioning from the aqueous phase into an organic phase (typically fish) and does not include uptake due to diet. The BCF has no units, as shown in equation (a):

$$\text{BCF} = \frac{\text{Conc.inorganisms}}{\text{Conc.in the environment}}.$$ (a)

Some reference values have been defined for BCF risk assessment. A substance with a BCF greater than 2,000 will be regarded as bioaccumulative for EU REACH. A substance with a BCF greater than 5,000 will be regarded as very bioaccumulative. In the United States, a substance is considered not bioaccumulated for a BCF value less than 1,000, bioaccumulated for a BCF value from 1,000 to 5,000, and very bioaccumulated for a BCF value greater than 5,000.

From the reference values cited, all investigated compounds predicted had a low bioaccumulative risk. Interestingly, none of the tested compounds presented an alarming bioaccumulative risk (greater than 5,000) (Table 3).
TABLE 3 | Results of the compound ecotoxicity.

| Compounds                          | BCF   | BOD    | Aquatic toxicity       | Endocrine receptor binding |
|------------------------------------|-------|--------|------------------------|----------------------------|
|                                    |       |        | Th_pyr_pIGC\(_{50}\) | Daphnia\(_{LC50}\) | Minnow\(_{LC50}\) | Andro\(_{Filter}\) | Estro\(_{Filter}\) |
|------------------------------------|-------|--------|------------------------|----------------------------|
| Quercetin-3-rhamnosyl-(1-6)-galactoside | 0.422 | No     | -1.292                 | 32.942                     | 1365.521               | Toxic               | Toxic               |
| Hyperoside                         | 1.015 | No     | -0.557                 | 54.841                     | 350.178                 | Toxic               | Non-toxic           |

The endpoints evaluated for each compound are defined as follows: BCF, bi-concentration factor; BOD, biodegradation – categorizing the compounds as positive (readily biodegradable) if % biodegradation is greater than or equal to 60% and as negative otherwise; Th_pyr_pIGC\(_{50}\), concentration (mM) of toxicant needed to inhibit growth by 50% (IGC\(_{50}\)) of Tetrahymena pyriformis; Daphnia\(_{LC50}\), concentration (mg/L) of compound required to kill 50% of a D. magna population; Minnow\(_{LC50}\), concentration (mg/L) of a compound that kills 50% of a population of minnows; Andro\(_{Filter}\) and Estro\(_{Filter}\) assess a compound’s likelihood of binding to the androgen and estrogen receptors, respectively.

Such findings agree with the compound physical–chemical characteristics. The compounds that presented a higher risk of being bioaccumulative were those with lower water solubility.

Biodegradation (BOD) of chemicals in the environment is becoming increasingly important. For the biodegradation assessment, the compound of interest is mixed with sludge from several geographical locations; then, oxygen consumption is measured for the specified period. Biological oxygen demand (BOD) is calculated from equation (b):

\[
BOD = \frac{[O_2 \text{ uptake by test substance}] - [O_2 \text{ uptake by blank}]}{\text{amount of test substance}} \tag{b}
\]

The percentage biodegradation is obtained from equation (c):

\[
\% \text{biodegradation} = \frac{BOD}{\text{ThOD}} \times 100\% \tag{c}
\]

ThOD represents the amount of oxygen consumed in converting all the compounds to carbon dioxide (CO\(_2\)), i.e., the theoretical oxygen demand. The ADMET predictor categorizes the compounds as positive (readily biodegradable) if % biodegradation is greater than or equal to 60% and as negative otherwise. Thus, the model returns a “yes” if the compound is readily biodegradable or “No” if not.

All the compounds tested are predicted to be non-biodegradable. Therefore, attention must be paid to the amount of waste generated in the environment and its potential harm (Table 3).

Aquatic toxicity can also be measured as its potential to inhibit the growth of living aquatic beings. Therefore, the ADMET predictor provides three models to investigate this characteristic at different trophic levels [Tetrahymena pyriformis, water flea (Daphnia), and fathead minnow].

The model that evaluates T. pyriformis growth inhibition predicts the concentration (mM) of toxicant needed to inhibit growth by 50% (IGC\(_{50}\)) in the protozoan after approximately 40 h of exposure (eight to nine cell cycles in the control group) at 27°C.

The T. pyriformis model showed that 20% of the most toxic compounds presented pIGC\(_{50}\) values less than approximately -1.0. Therefore, quercetin and hyperoside presented a higher potential to inhibit T. pyriformis growth. Water fleas (Daphnia magna) occupy the next rung up on the aquatic food chain from Tetrahymena. Daphnia\(_{LC50}\) predicts the concentration of compounds required to kill 50% of a D. magna population within 48 h. The data for this study were obtained from the EPA’s ECOTOX database.

The developed D. magna model showed that 20% of the most toxic compounds presented an LC\(_{50}\) of less than approximately 3.0. Therefore, the compounds, quercetin and hyperoside, did not present potential aquatic toxicity.
The third trophic level investigated was the effect of the compound on *Pimephales promelas*. The ADMET predictor model was developed based on the database created by the Mid-Continent Ecology Division of the U.S. Environmental Protection Agency testing industrial compounds for lethal effects on the fathead minnow (*P. promelas*). The experimental protocol was used to determine the concentration of a compound that killed 50% of a population of minnows after an exposure time of 96 h.

The *P. promelas* model showed that approximately 20% of the most toxic compounds presented an IGC<sub>50</sub> less than 2.0. Therefore, quercetin and hyperoside did not present potential aquatic toxicity.

The potential environmental damage caused by exposure to endocrine receptor binding disruptors has gone unappreciated until relatively recently. The compounds compete for binding with estrogenic or androgenic sex hormones, blocking the timely transmission of standard hormonal signals or producing them in a few cases.

ADMET predictor models of endocrine receptor binding are used to assess a compound's likelihood of binding to the estrogen/androgen receptor. Estro_Filter and Andro_Filter are straightforward predictions of whether the molecule will have an appreciable affinity for the estrogen or androgen receptor at all. A negative (non-toxic) classification indicates that the compound is unlikely to cause endocrine disruption by binding to the receptor. In contrast, a positive (toxic) indicates that the compound is likely to bind detectably to the receptor.

Regarding the estrogen receptor, hyperoside is predicted to have a non-toxic effect compared with quercetin. On the other hand, quercetin and hyperoside have the potential to interact with the androgen receptor. Therefore, these compounds risk adverse health outcomes, including cancer, reproductive impairment, cognitive deficits, and obesity.

**CONCLUSION**

Nano-SOB ameliorated the extract-SOB mOLLUSCICIDAL activity on the population of *B. glabrata*. This biotechnological strategy rescued extract-SOB activity with an LC<sub>50</sub> value in the range recommended by the WHO. Nano-SOB also acted on young *B. glabrata* mollusks with similar potency to adult *B. glabrata*. Additionally, nano-SOB exhibited notable activity against the cercariae of *S. mansoni*, where, after 4 h, all the cercariae were eliminated. Furthermore, extract-SOB and nano-SOB were not effective in reducing viable spawning. Concerning ambient toxicity, nano-SOB was moderately toxic to *Melanoides sp*. Thus, nano-SOB proved to be an excellent alternative in controlling and fighting against schistosomiasis, acting in several disease transmission cycle phases. Additionally, quercetin-3-rhamnosyl-(1-6)-galactoside and hyperoside are biodegradable and pose a low risk of environmental accumulation and aquatic toxicity. However, these substances pose a risk for human manipulation.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

**AUTHOR CONTRIBUTIONS**

APO, DQF, and LR performed the experiments based on chemical isolation of the essential oil and nanoemulsion. LSR performed the experiments based on biological assays. NLVR and CRR performed the experiments based on *in silico* assays. MGS identified and deposited the exsiccate. RXF, JAAS, and LR wrote and supervised the manuscript. All authors contributed to the article and approved the submitted version.

**FUNDING**

This work was supported by CNPq (National Council of Research of Brazil) (RXF holds a grant with Fellowship Process Number 308755/2018-9), FAPERJ (Research Support Foundation of the State of Rio de Janeiro) [JCNE (Young Scientist from Our State) with fellowship process number: E-26/203.246/2017], and the Emergent Group of Research from Rio de Janeiro (E-26/211.025/2019) for financial support, and CAPES (Coordination for the Improvement of Higher Education Personnel) provided support through scholarships.

**ACKNOWLEDGMENTS**

We thank CNPq, CAPES, and FAPERJ for their financial support.

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