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An example of molecular co-evolution: reactive oxygen species (ROS) and ROS scavenger levels in Schistosoma mansoni/Biomphalaria glabrata interactions

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Abstract

The co-evolution between hosts and parasites involves huge reciprocal selective pressures on both protagonists. However, relatively few reports have evaluated the impact of these reciprocal pressures on the molecular determinants at the core of the relevant interaction, such as the factors influencing parasitic virulence and host resistance. Here, we address this question in a host-parasite model that allows co-evolution to be monitored in the field: the interaction between the mollusk, Biomphalaria glabrata, and its trematode parasite, Schistosoma mansoni. Reactive oxygen species (ROS) produced by the hemocytes of B. glabrata are known to play a crucial role in killing S. mansoni. Therefore, the parasite must defend itself against oxidative damage caused by ROS using ROS scavengers in order to survive. In this context, ROS and ROS scavengers are involved in a co-evolutionary arms race, and their respective production levels by sympatric host and parasite could be expected to be closely related. Here, we test this hypothesis by comparing host oxidant and parasite antioxidant capabilities between two S. mansoni/B. glabrata populations that have co-evolved independently. As expected, our findings show a clear link between the oxidant and antioxidant levels, presumably resulting from sympatric co-evolution. We believe this work provides the first supporting evidence of the Red Queen Hypothesis of reciprocal evolution for functional traits at the field-level in a model involving a host and a eukaryotic parasite.

Keywords: Host-parasite co-evolution, Schistosoma mansoni, Biomphalaria glabrata, Reactive oxygen species (ROS), ROS scavengers
1. Introduction

Understanding the co-evolution of host-parasite interactions represents a challenge in evolutionary biology. Parasites cause substantial deleterious effects on their hosts, and therefore represent a major driving force in their evolution (Howard, 1991). Similarly, the host immune defenses represent the major selective pressure driving the evolution of parasites. For parasites to survive and develop in the host they must adapt to the host-defense system or they will die. This parallel co-evolution of host-parasite interactions can be viewed as an arms race in which both the host and the parasite develop mechanisms to circumvent the weapons developed by their opponent.

In this context of reciprocal co-evolution, illustrated by Van Valen (1974), under the Red Queen Hypothesis it is assumed that the parasitic genes responsible for infectivity will evolve alongside the host defense genes, resulting in adaptation of the interactions between local host and parasite populations (Dybdahl and Storfer, 2003). To date, however, only a few studies have sought to verify this prediction and convincing experiments have only been reported for models involving viruses, bacteria and unicellular eukaryotes (Lohse et al., 2006; Forde et al., 2008).

Demonstrating co-evolution in an animal host-parasite system is not straightforward and most prior discussions of such processes have been indirect, as in studies describing local adaptation when compatibility is higher between sympatric host-parasite combinations than between allopatric combinations (Gasnier et al., 2000; Gagneux et al., 2006; Munoz-Antoli et al.), or studies that have focused on only one trait of the interaction, such as host resistance (Green et al., 2000) or parasite infectivity (Little et al., 2006). Moreover it is important to take into account that non-co-evolutionary mechanisms could also explain correlations between the traits of interacting species and that the absence of correlated traits is not evidence for an absence of co-evolution (Nuismer et al., 2007, 2010; Yoder and Nuismer, 2010). However we assume that the direct examination of reciprocal selection in both the host and the parasite could provide supporting evidence of co-evolution.
Two relatively recent studies investigated this reciprocal response more thoroughly. In the first, reciprocal changes in resistance and infectivity were identified for co-evolving *Potamopyrgus* snail hosts and their trematode parasites; however, while these changes were identified using prevalence phenotypes, they were not supported by the studied functional markers (Koskella and Lively, 2007). The second report provided experimental support for the reciprocity of adaptation costs, rapid genetic changes and increased genetic diversity during the co-evolution of a multicellular host, the nematode *Caenorhabditis elegans*, and its pathogenic bacteria, the Gram-positive bacterium, *Bacillus thuringiensis* (Schulte et al., 2010). In both papers, co-evolution was studied using laboratory strains selected by experimental evolutionary approaches, and only the second paper focused on molecular changes induced by the co-evolutionary process. Both papers were based on an "over time" approach in which the evolution of phenotypes was monitored over the course of experimental laboratory generations. Other empirical studies have been based on a "point time" approach in which the pattern of co-variations between host-parasite populations or strains that co-evolved independently were assessed at a single experimental time point (Forde et al., 2004; Morgan et al., 2005).

Here, we investigated the reciprocal evolution of molecular mechanisms directly at the core of the host-parasite interaction in a natural system of co-evolution, by comparing host and parasite populations that have co-evolved independently. As a model, we used the interaction between the trematode, *Schistosoma mansoni* (responsible for human intestinal schistosomiasis), and its mollusk intermediate host, *Biomphalaria glabrata*. This interaction is a model of choice for the study of potential co-evolutionary dynamics (Webster and Davies, 2001; Webster et al., 2004; Beltran and Boissier, 2008; Beltran et al., 2008; Bouchut et al., 2008; Roger et al., 2008a; Roger et al., 2008b; Roger et al., 2008c; Steinauer, 2009).

During its intramollusksal stage, the parasite must cope with the snail’s immune system. One of the main immune effectors in mollusks are the reactive oxygen species (ROS) produced by hemocytes (the circulating immune cells of snails) (Hahn et al., 2000; de Jong-Brink et al., 2001;
Hahn et al., 2001b; Mourao et al., 2009b). Previous studies conducted by Hahn and co-workers demonstrated that hydrogen peroxide ($\text{H}_2\text{O}_2$) plays a crucial role in the killing of $S. \text{mansoni}$ sporocysts (Hahn et al., 2001a, b). Furthermore, hemocytes from $S. \text{mansoni}$-resistant snails were shown to generate significantly more $\text{H}_2\text{O}_2$ than susceptible snails, perhaps due at least in part to the former having constitutively elevated levels of the mRNA encoding the copper/zinc superoxide dismutase (Cu-ZN SOD) (Goodall et al., 2004; Bender et al., 2005; Bender et al., 2007). To resist ROS-mediated attacks, the $S. \text{mansoni}$ larvae produce ROS-detoxifying enzymes (Vermeire et al., 2006; Guillou et al., 2007; Vermeire and Yoshino, 2007; Roger et al., 2008c; Wu et al., 2009), several of which appear to be secreted by sporocysts (Guillou et al., 2007; Wu et al., 2009). Supporting this, a recent report showed that antioxidant enzymes produced by $S. \text{mansoni}$ sporocysts are directly involved in protecting the pathogen against immune cell-mediated oxidative stress (Mourao et al., 2009b).

In this context, the snail-produced ROS and the parasite-produced ROS scavengers are involved in a co-evolutionary arms race, and we can hypothesize that their production levels will be closely related. Here, we tested this hypothesis by comparing host oxidant and parasite antioxidant abilities for two $S. \text{mansoni}/B. \text{glabrata}$ strains that have evolved independently, originated from different geographic endemic zones and which are found to display significant differences in compatibility.

### 2. Materials and methods

#### 2.1 Ethics statement

Our laboratory has received the permit # A66040 for experiments on animals from both the French Ministry of agriculture and Fishing and the French Ministry of National Education, Research and Technology. Housing, breeding and animal care of the mice followed the ethical requirements of French government. The experimenter possesses the official certificate for animal experimentation.
2.2. Biological materials

Two strains of *S. mansoni* were used in this study: a Brazilian strain (*Sm*BRE) and a Guadeloupean strain (*Sm*GH2). Each strain was maintained: (i) in their sympatric strain of *B. glabrata* (*Bg*BRE and *Bg*GUA, respectively); and (ii) in hamsters (*Mesocricetus auratus*), as described previously (Théron et al., 1997).

Miracidia from *Sm*BRE and *Sm*GH2 were hatched from eggs axenically recovered from 60-day-infected hamster livers, according to the previously described procedure (Roger et al., 2008c). Briefly, livers were collected and kept overnight at 4°C in sterile saline solution (NaCl 150 mM) containing an antibiotic/antimycotic mixture (penicillin 100 units/ml, streptomycin 0.1 mg/ml, amphotericin B 0.25 μg/ml; Sigma). The livers were then homogenized and the eggs were filtered out, washed, and transferred to spring water. The miracidia were allowed to hatch out under illumination.

2.3. Schistosome-snail compatibility: snail exposure, infection rates and intensities

The compatibilities of the tested snail-schistosome combinations were evaluated by monitoring the infection rates (% of snails infected) and the intensity of infection (number of mother sporocysts (SpI) developed) among snails individually challenged with different numbers of miracidia. As the miracidial dose increased, a larger fraction of the phenotypic diversity in the parasitic isolate was sampled; thus, dose-response curves are much more informative than single-dose challenges when examining the dynamics of compatibility between two host-parasite combinations (Théron et al., 2008).

For each experiment, snails (7-9 mm in diameter) were exposed individually to a fixed number of miracidia in approximately 10 ml of water for 8 h. Following exposure to miracidia, the
snails were replaced in their original containers until their infection status (presence of SpI) was assessed. For the detection of SpI, the snails were fixed 15 days post-exposure, following previously described methods (Gerard et al., 1995; Moné et al., 2010b). In brief, each snail was relaxed in pond water containing an excess of crystalline menthol for 6 h, the body was removed and fixed in modified Raillet-Henry’s solution, exhaustive dissection of the head-foot zone was performed, and the number of SpI present in each snail (readily observable as translucent white bodies within an opaque gray tissue background) was determined.

Dose-response curves were obtained by challenging individual snails (30-40 snails per treatment) with doses of 1, 10, 20, 30 and 50 miracidia. Compatibility was measured for the two sympatric combinations (SmBRE versus BgBRE and SmGH2 versus BgGUA) and the two allopatric combinations (SmBRE versus BgGUA and SmGH2 versus BgBRE).

2.4. Cytotoxicity of \( \text{H}_2\text{O}_2 \) on \( S. \text{mansoni} \) sporocysts

\( \text{H}_2\text{O}_2 \) cytotoxicity was measured using the Roche Cytotoxicity Detection Kit (Roche Diagnostics, Mannheim, Germany), which is based on the measurement of lactate dehydrogenase (LDH) activity released from dead and lysed cells into the supernatant. Four hundred miracidia each of \( \text{SmBRE} \) and \( \text{SmGH2} \) were submitted to in vitro transformation to obtain primary sporocysts (Sp1). Briefly, the miracidia were cultured for 24 h in sterile Chernin’s balanced salt solution (CBSS) (Chernin, 1963), containing an antibiotic/antimycotic mixture (penicillin 100 units/ml, streptomycin 0.1 mg/ml, amphotericin B 0.25 \( \mu \text{g/ml} \); Sigma). The sporocysts were then exposed to four different concentrations of \( \text{H}_2\text{O}_2 \) (0, 75, 150 and 200 \( \mu \text{M} \)) for 2 h and cytotoxicity was examined according to the manufacturer’s instructions. As a positive control, we measured LDH release from Sp1 that had been lysed with the provided lysis solution (high control, HC); this was taken as 100% LDH release. To correct for the background, we measured LDH levels in Sp1-free \( \text{H}_2\text{O}_2 \)-treated culture medium (substance control, SC). All measured values were assayed in
triplicate. The percentage of specific H$_2$O$_2$-induced LDH release was determined as: % cytotoxicity

\[ \text{% cytotoxicity} = \frac{\text{experimental result} - \text{SC}}{\text{HC} - \text{SC}} \times 100. \]

2.5. Effect of H$_2$O$_2$ on S. mansoni sporocyst mortality

Two independent experiments were conducted in triplicate on 24-well plates containing 20 Sp1 (representing SmBRE or SmGH2) per well. The Sp1 were in-vitro transformed as described above (see Materials and methods section 2.2), and exposed to 0, 200, 400, 800 or 1,600 µM of H$_2$O$_2$ (Hydrogen peroxide 35%, FLUKA, Germany) for 4 h. Mortality was assessed under a light microscope, with the Sp1 considered “dead” when we failed to observe motility and/or the beating of the flame-cell flagella.

2.6. The total antioxidant capacity of S. mansoni sporocysts

The cumulative (total) antioxidant capacity of the sporocysts was quantified for the two parasite strains, SmBRE and SmGH2, using an Antioxidant Assay Kit (Sigma). For each test, 2,000 sporocysts were in-vitro transformed as described above. After 24 h, fully transformed sporocysts were recovered by gentle centrifugation (800 g, 5 min, 4°C). The samples were then disrupted by sonication (three pulses of 20 s each) and pelleted by centrifugation (12,000 g, 15 min, 4°C), and the antioxidant capacity of each supernatant was determined following the manufacturer’s recommendations. The amount of protein in each supernatant was determined using a Bradford protein assay kit and used as a correcting factor. The experiment was performed six times per strain.

2.7. Reverse Transcription-quantitative PCR (RT-qPCR)

RT-qPCR analyses were conducted to compare the expression of parasite antioxidant enzymes suspected to play key roles in the detoxification of host-induced oxidative stress. Real-time PCR analyses were performed using a LightCycler 2.0 system (Roche Applied Science) and a LightCycler Faststart DNA Master SYBR Green I kit (Roche Applied Science). Total RNA extractions from miracidia were performed using the Trizol Reagent (Life Technologies, USA) and
the manufacturer’s protocol. Reverse transcription was performed according to previously described procedures (Guillou et al., 2004). qPCR amplification was performed using 2.5 μl of cDNA in a final volume of 10 μl containing 3 mM MgCl₂, 0.5 μM of each primer and 1 μl of master mix. The primers were designed using either the LightCycler probe design software or the web-based Primer3 plus interface (http://www.bioinformatics.nl/CGIBIN/primer3plus/primer3plus.cgi) and are given in Table 1. The following Light-Cycler run protocol was used: denaturation at 95°C for 10 min, followed by 40 cycles of amplification and quantification at 95°C for 10 s, 60°C for 5 s and 72°C for 16 s, a melting curve of 60–95°C with a heating rate of 0.1°C/s and continuous fluorescence measurement, and then a cooling step to 40°C. For each reaction, the cycle threshold (Ct) was determined using the “Fit Point Method” of the LightCycler Software, version 3.3. The PCR reactions were performed in duplicate and the mean Ct value was calculated. For each sample, the expression level of the target gene was normalized with regard to the expression of two constitutively expressed genes (28S rRNA and α tubulin). The expression ratio (R) was calculated according to the formula: $R = 2^\Delta C_t$, where $\Delta C_t$ represents Ct (target gene) – Ct (constitutively expressed gene).

2.8. ROS detection in single cells

The cell-permeable fluorescent oxygen probe, 1-pyrenebutyric acid (PBA), can be used for the measurement of free radicals in solution (Oter and Ribou, 2009) and in living cells (Ribou et al., 2004; Rharass et al., 2006), with the fluorescence intensity and lifetime of PBA decreasing proportionately to the free-radical concentration. Measurement of the fluorescent lifetime offers many advantages over intensity based measurements when working in vivo, not the least that the measurements are independent of the absolute intensity of emitted light and the fluorophore concentration, thereby avoiding artifacts arising from optical losses. Moreover, these probes do not require a reaction with ROS, are usually stable and the fluorescent lifetime is not modified by probe degradation or variations in its intracellular accumulation.
2.8.1. Staining and fixation

Hemolymph samples were recovered from BgBRE and BgGUA snails, and aliquots (150 µl) were put in a Sykes-Moore chamber. After 4 h, the adhered hemocytes were rinsed with Hank's buffered salt solution (HBSS) and stained for 20 min with PBA (Acros Organics, Belgium; 0.10 µM in 1% ethanol). The hemocytes were then rinsed three times and placed in HBSS for measurements. For fixation experiments, hemocytes were treated as described above except that after the final rinsing step, the cells were killed by incubation for 10 min in Baker solution (10% paraformaldehyde in 1% aqueous calcium chloride). In the latter case, the experiments were performed within 1 h after cell killing, in order to avoid probe reorganization (Ribou et al., 2004).

2.8.2. ROS quantification by fluorescent-lifetime measurement of single cells

The fluorescent decay of single living cells loaded with PBA was recorded using time-resolved microfluorimetry, as previously described (Ribou et al., 2003). Briefly, a laser (nitrogen laser NL100; Stanford Research Systems, USA) delivered monochromatic 337-nm pulses, each with a half-amplitude pulse-width of 3 ns, and an objective (40×; Unitron) was used to concentrate the excitation beam on the microscopic sample. Emitted photons were collected and focused on a photomultiplier 1P28 (Hamamatsu Corporation, Japan). A diaphragm placed on the emission pathway allowed the selection of signals from single cells, while a 404-nm bandpass filter (half bandwidth; 40 nm) also located along the emission pathway was used to select the pyrene emission. Each signal was digitalized by a digital oscilloscope (TDS 3032C; Tektronix, USA). The fluorescent decay of single PBA-loaded cells selected by the 404-nm filter could be resolved into three exponential curves. The time constants (i.e. lifetimes) and amplitude values of each exponential curve in the decay were obtained using the downhill simplex method (Nelder and Mead, 1965). The first two decays corresponded to the intrinsic fluorescence of the cell attributed to the reduced form of NAD(P)H. The third long-time constant (> 100 ns), which was characteristic of
pyrene derivatives, was related to the ROS concentration through the Stern-Volmer equation (Stern and Volmer, 1919) that describes collisional fluorescent quenching of a probe (i.e. PBA) by a quencher (i.e., free radicals). The method has been described in several papers (Ribou et al., 2003; Ribou et al., 2004; Rharass et al., 2006). We calculated the variation of intracellular ROS concentrations as follows:

\[
\frac{[\text{ROS}]}{[\text{ROS}]_m} = \frac{[t_m (t_0 - t)]}{[t(t_0 - t_m)]} \quad \text{Equation 1}
\]

where \( t \) is the fluorescent lifetime measured for 108 single hemocytes originating from eight \( Bg\text{BRE} \) snails and 111 cells from eight \( Bg\text{GUA} \) snails; \( t_m \) is the mean of all lifetimes; and \( t_0 \) is the fluorescent lifetime in the absence of ROS (measured from dead hemocytes fixed with Baker solution). In this equation, \([\text{ROS}]_m\) is the mean of the concentrations from all tested cells (219 cells). We assumed that fixation ended all cellular activity and ROS production. In the presented data, the mean ROS concentration has been assigned an arbitrary value of 1.

2.9. \( \text{H}_2\text{O}_2 \) production and release by \( B. \text{glabrata} \) hemocytes

The \( \text{H}_2\text{O}_2 \) production by hemocytes was measured using Amplex® Red (Invitrogen). Hemolymph was collected from the head-foot regions of \( Bg\text{BRE} \) and \( Bg\text{GUA} \) snails (7-11 mm in diameter) as previously described (Bouchut et al., 2006), and the number of hemocytes per \( \mu l \) of hemolymph was quantified using a cell counter (Z Series Coulter Counter; Beckman Coulter); 226.2 ± 50.6 cells /\( \mu l \) and 241.2 ± 102.1 cells /\( \mu l \) were obtained for \( Bg\text{BRE} \) and \( Bg\text{GUA} \), respectively. The Hemolymph of four snails was pooled and 45,000 hemocytes per well were dispensed to a 96-well plate for each strain. The hemocytes were allowed to adhere and spread for 1 h at 26°C. The plate was then centrifuged (600g for 10 min), the plasma was removed, the adhered hemocytes were washed three times with HBSS, and the wells were treated with Amplex® Red reaction mixture (100 \( \mu L \) per well, prepared according to the manufacturer’s instructions). Optical density was measured with a microplate reader at 570 nm during the following 1 h (at 5, 10, 15, 20, 30, 40, 50, 60 min). The results are expressed as Amplex Red O.D. at 570 nm / 45,000 cells.
2.10. Biomphalaria glabrata superoxide anion plasma content

The plasma content of superoxide anion was monitored via the superoxide-mediated reduction of nitroblue tetrazolium (NBT), which results in the precipitation of an insoluble blue formazan that can be quantified spectrophotometrically. Briefly, hemolymph was collected from BgBRE and BgGUA snails as described above. Hemocytes were removed by centrifugation (1,500 g for 15 min), and then 50 µL of plasma from each snail was mixed with 50 µl of 0.1% NBT (Sigma) dissolved in PBS; (Na₂HPO₄ 8.41 mM, NaH₂PO₄ 1.65 mM, NaCl 45.34 mM, pH 7.45).

Formazan blue formation was measured with a microplate reader at 620 nm over the course of 3 h (at 5, 10, 15, 30, 60, 90, 180 min). NBT-free plasma was used as a control, and triplicate experiments were conducted for 10 individuals per strain.

2.11. Statistical analyses

The normality of our experimental data was assessed using the Shapiro-Wilk normality test (Shapiro and Wilk, 1965). Our data on the effect of H₂O₂ cytotoxicity on S. mansoni sporocysts (LDH test), B. glabrata hemocyte H₂O₂ production, and superoxide anion plasma content were all found to be normally distributed ($P > 0.05$), and were subsequently analyzed using the student's $t$-test. Our data on the effect of H₂O₂ on S. mansoni sporocyst mortality and the total antioxidant capacity of sporocysts were not normally distributed ($P < 0.05$), and were subsequently analyzed using the Mann-Whitney test. The results of the ROS concentration assays in each mollusk strain were analyzed using the Mann-Whitney test. The Kolmogorov-Smirnov two-samples test was utilized to determine whether the ROS concentrations were similarly distributed in hemocytes from BgBRE and BgGUA.

3. Results

3.1. Effect of H₂O₂ on S. mansoni sporocysts
Two different assays were conducted to test the effect of H$_2$O$_2$ on the two strains of *S. mansoni* sporocysts (SmBRE and SmGH2) (Fig. 1). First, an LDH test was used to examine the cytotoxicity of H$_2$O$_2$ on sporocysts of each strain. Our results revealed that the susceptibility to H$_2$O$_2$ was significantly higher for SmGH2 than SmBRE (Fig. 1A). When exposed to 200 µM H$_2$O$_2$, SmGH2 sporocysts showed 25.8% cytotoxicity (i.e., 25.8% of the cells had lysed and released their LDH content), whereas no changes were observed for SmBRE at the same H$_2$O$_2$ concentration (Fig. 1A). However, although cell lysis occurred in SmGH2, the sporocysts were still alive at this concentration. To investigate possible between-strain differences in mortality, we next exposed sporocysts to increasing concentrations of H$_2$O$_2$ and examined motility and the beating of the flame-cell flagella, which were taken as distinguishing between living and dead larvae. No difference between the two strains was observed until the concentration of H$_2$O$_2$ reached 1,600 µM (Fig. 1B). At this concentration, 31.2% and 2.6% of the SmGH2 and SmBRE sporocysts were dead, respectively; this difference is statistically significant (student’s *t*-test; *P* = 0.017). These results suggest that SmGH2 sporocysts are more susceptible to H$_2$O$_2$ than SmBRE sporocysts.

3.2. The total antioxidant capacity of *S. mansoni* sporocysts

To investigate potential differences in the constitutive antioxidant abilities of sporocysts from SmBRE and SmGH2, we measured the cumulative antioxidant activities of these two strains (Fig. 2). Our results revealed that the antioxidant ability of SmGH2 was significantly lower than that of SmBRE (approximately 13% less; Mann-Whitney test, *P* = 0.0001).

3.3. Reverse Transcription-quantitative PCR of ROS-scavenger expression among *S. mansoni* strains

RT-qPCR was used to compare the expression of parasite antioxidant enzymes suspected to play key roles in the detoxification of host-induced oxidative stress (Table 1) (Guillou et al., 2007). Notably, Cu-Zn SOD (Smp_176200.2) was found to be expressed at a significantly higher level in
Sm BRE than in Sm GH2 (2.7-fold; \( P = 0.017 \)) (Fig. 3). In contrast, no difference was observed in the expression levels of glyceraldehyde-3-phosphate dehydrogenase (GAPDH, Smp_056970.1), GST omega (Smp_152710.1), GST 28 kD (Smp_054160), GST 26 kD (Smp_163610), glyoxalase II (Smp_091010) or thioredoxin peroxidase (TPX, Smp_158110) (data not shown).

3.4. Intracellular ROS measurements in single B. glabrata hemocytes

We monitored intracellular ROS levels in single hemocytes, using PBA. This method allows global ROS to be measured without interference from the reactive hydroxyl radical or \( \text{H}_2\text{O}_2 \). PBA fluorescent lifetimes were measured for 108 and 111 individual hemocytes originating from eight Bg BRE and eight Bg GUA snails, respectively. Bg BRE hemocytes produced significantly (11.2%) more ROS than Bg GUA hemocytes (Mann-Whitney test; \( P = 0.009 \)) (Fig. 4A). Fig. 4B shows the distribution of hemocytes from both strains according to their ROS concentrations, which were calculated from the ratio given in equation 1 (see Materials and methods section 2.8.2.). Although the cells from both Bg BRE and Bg GUA samples were distributed around the mean ROS concentration, their distributions were significantly different (Kolmogorov-Smirnov test, \( P = 0.012 \)). Among the hemocytes producing more than 1.5-fold of the mean ROS concentration, 68.8% were from Bg BRE strain, while only 31.2% were from Bg GUA. Conversely, among the hemocytes that showed the lowest ROS concentrations (< 0.7-fold of the mean ROS concentration) 38.1% were from Bg BRE and 62% were from Bg GUA (Fig. 4B). By recording the fluorescent lifetimes of single cells loaded with PBA, we also obtained the relative concentrations of free and bound NAD(P)H (an indicator of metabolic change) in each cell. However there was no significant difference in the quantity of bound and free NAD(P)H (mean ratio = 0.60 for both strains; data not shown), suggesting that the strains had similar levels of metabolism. Thus, the only molecular difference observed between the two strains was the level of ROS production.

3.5. \( \text{H}_2\text{O}_2 \) production and release by B. glabrata hemocytes
We assessed H$_2$O$_2$ production and secretion by hemocytes of both strains using Amplex Red. Fig. 5 shows the cumulative amount of H$_2$O$_2$ constitutively released by hemocytes of each strain over 1 h. The maximum level of H$_2$O$_2$ production was reached at 20 min for BgGUA and at 40 min for BgBRE after addition of Amplex Red substrate. In total, BgBRE hemocytes produced significantly more (1.44-fold; mean value) H$_2$O$_2$ than BgGUA hemocytes (student’s t-test; P < 0.05) (Fig. 5).

3.6. Biomphalaria glabrata superoxide anion plasma content

To our knowledge, all spectrophotometric methods currently available for the determination of H$_2$O$_2$ are based on the measurement of red or orange pigments, making these methods unsuitable for use on B. glabrata plasma samples, which are already tinted red by hemoglobin. Consequently, we used NBT to measure the amount of superoxide anion (O$_2^-$; a precursor of H$_2$O$_2$) produced in both strains. As shown in Fig. 6, at 3 h after addition of NBT BgBRE plasma contained significantly more (44% more) superoxide anion than BgGUA plasma (student’s t-test; P = 0.0007).

3.7. Compatibility of sympatric and allopatric S. mansoni/B. glabrata combinations

Sympatric pairings of S. mansoni and B. glabrata originating from Brazil and Guadeloupe were previously shown to display different levels of compatibility that remained remarkably stable across laboratory generations (Théron et al., 2008). We first used dose-response curves obtained by challenging snails with increasing doses of miracidia to confirm that similar differences could be observed between our strains. At doses of 10 or more miracidia/snail, SmBRE/BgBRE showed an infection rate of 100%, while SmGH2/BgGUA had an infection rate of approximately half that, at around 55%. Interestingly, differences were also observed for the number of parasites (SpI) that develop within the snails. The infection intensity rose gradually as the challenge doses increased for SmBRE/BgBRE, reaching 16.18 ± 0.86 parasites/snail at the 50-miracidia dose. In contrast, the
infection intensity for *Sm*GH2/*Bg*GUA remained low regardless of the challenge dose, varying between 1.6 ± 0.20 and 3.2 ± 0.64 parasites/snail (Fig. 7).

When we tested the heterologous combinations, we found that the *Sm*BRE/*Bg*GUA pairing showed a substantial level of compatibility, with infection rates of 80-90% (not significantly different from the 100% achieved by the *Sm*BRE/*Bg*BRE pairing), but with lower parasite intensities (9.8 ± 0.89 for the 50-miracidia dose) compared with the sympatric combination (16.18 ± 0.86 parasites/snail at the 50-miracidia dose). In contrast, the *Sm*GH2/*Bg*BRE combination showed very little infectivity, with infection rates < 6% and ~1 parasite/snail regardless of the challenge dose (Fig. 7).

3.8. ROS, ROS scavengers and compatibility in sympatric and allopatric *S. mansoni/B. glabrata* combinations

The above-described results indicated that levels of ROS and ROS scavengers were correlated in both sympatric combinations, with high-level ROS/ROS scavenger production in the Brazilian combination, but lower-level ROS/ROS scavenger production in the Guadeloupean combination. If high levels of *S. mansoni* ROS scavenger are correlated with better resistance of the intramolluskan stage of the parasite (as we hypothesized), we would expect *Sm*GH2 to have a relatively low ability to infect the allopatric *Bg*BRE snails, while *Sm*BRE would have a high ability to infect the allopatric *Bg*GUA snails. This hypothesis was verified in our model, as shown in Fig. 8.

4. Discussion

Snail-schistosome compatibility and infection rates result from a complex interplay between the host’s defense mechanisms and the parasite’s infectivity strategies. Due to selective pressures exerted by the parasite on the host and vice versa, co-evolutionary dynamics may be observed (Janzen, 1980; Howard, 1991). Between-population or between-strain differences in the outcomes
of such evolutionary processes may be expected due to differences in the epidemiological and environmental conditions, and/or genetic architectures. Such differential selection patterns could explain, at least in part, the geographic and/or strain-specific compatibility variations seen in snail-schistosome interactions (Théron et al., 2008). At present, however, there is relatively little empirical evidence demonstrating reciprocal molecular adaptations in both host and parasite.

Here, we investigated the interaction between *S. mansoni* and the snail, *B. glabrata*, as this interaction is a popular model for the study of co-evolutionary dynamics (Beltran and Boissier, 2008; Beltran et al., 2008; Bouchut et al., 2008; Roger et al., 2008a; Roger et al., 2008b; Roger et al., 2008c; Steinauer, 2009). We confirmed that there are different levels of compatibility between two geographic strains of *S. mansoni* and their sympatric snail hosts, *B. glabrata* (Fig. 7) that both have co-evolved independently. We compared the host oxidant and parasite antioxidant abilities that appear to form the core of the attack/defense interactions of these two pairings.

ROS are the main effectors of the snail immune system; they are highly reactive and can trigger irreversible cell damage. Indeed, ROS produced by the hemocytes of *B. glabrata* are known to play a crucial role in the killing of *S. mansoni* (Hahn et al., 2000; Hahn et al., 2001a, b; Bender et al., 2005; Bayne, 2009). Conversely, *S. mansoni* possess antioxidant systems capable of counteracting the ROS produced by their host’s immune system. *Schistosoma mansoni* is exposed to ROS in both their intermediate (snail) and definitive (human or mammalian) hosts, and produce oxidative-stress scavengers in their excretory-secretory products (ESP) during all stages of their life cycle (Mei and LoVerde, 1997; Curwen et al., 2004; Zelek and Von Janowsky, 2004; Knudsen et al., 2005; van Balkom et al., 2005; Bernal et al., 2006; Dzik, 2006; Perez-Sanchez et al., 2006; Cass et al., 2007; Guillou et al., 2007; Mourao et al., 2009a; Wu et al., 2009). Therefore, the success or failure of host invasion by *S. mansoni* depends at least in part on its ability to defend itself against oxidative damage (Mourao et al., 2009a). In this system, therefore, ROS and ROS scavengers should be involved in a co-evolutionary arms race, and we would expect their respective production levels in sympatric host/parasite combinations to be closely related.
As previous studies have established that H$_2$O$_2$ is the main ROS involved in killing *S. mansoni* sporocysts, probably due to its stability and capacity to cross cell membranes (Hahn et al., 2001b; Bienert et al., 2006), we studied the susceptibility of two strains of *S. mansoni* to H$_2$O$_2$. Our results showed a clear intrinsic difference between parasites isolated from two different geographic regions: the intramolluskan stages of Guadeloupean *S. mansoni* (SmGH2) were more sensitive to H$_2$O$_2$ than those of the Brazilian strain (SmBRE) (Fig. 1). Moreover, we observed a difference in antioxidant potential between strains, with SmGH2 displaying a lower level of antioxidant activity than SmBRE (Fig. 2). Thus, SmBRE has a more efficient antioxidant system, which would seem to explain its higher level of resistance to H$_2$O$_2$-mediated oxidative damage.

In order to identify the molecular pathways involved in these differential antioxidant properties, we investigated the strain-specific transcription levels of genes encoding various antioxidant enzymes, including GAPDH (Smp_056970.1), GST omega (Smp_152710.1), GST28 (Smp_054160), GST26 (Smp_163610), glyoxalase II (Smp_091010), thioredoxin peroxidase, and Cu-Zn SOD (Smp_176200.2) (Guillou et al., 2007; Vermeire and Yoshino, 2007; Roger et al., 2008c; Mourao et al., 2009a; Wu et al., 2009). Among these candidates, only the Cu-Zn SOD mRNA displayed differential expression, with expression levels that were 2.7-fold higher in SmBRE than in SmGH2 (Fig. 3). This finding is consistent with our protein-level results from a previous proteomic study (Roger et al., 2008c), and these observations collectively suggest that Cu-Zn SOD plays a key role in the antioxidant strategy of *S. mansoni*. The involvement of Cu-Zn SOD in ROS detoxification is a recurring and intriguing question, because it is capable of dismutating the superoxide anion (O$_2^-$) to produce H$_2$O$_2$ (Zelck and Von Janowsky, 2004; Guillou et al., 2007; Mourao et al., 2009a). The hypothesis currently used to explain the role of *S. mansoni* Cu-Zn SOD in ROS detoxification is based on a suspected peroxidative function (Yim et al., 1993; Yim et al., 1996; Kim and Kang, 1997; Bayne et al., 2001). In short, it has been proposed that *S. mansoni* Cu-Zn SOD could use its own dismutation product (H$_2$O$_2$) to produce hydroxyl radicals (HO') that are less toxic for sporocysts (Bayne et al., 2001).
In a co-evolutionary context, the between-strain differences in ROS susceptibility and antioxidant activity of these *S. mansoni* strains suggest that there could be comparable differences in the ROS production capabilities of the host snail strains. To test this hypothesis, we investigated ROS production by the two snail strains. First, we used a fluorescence-based method (Rharass et al., 2006) to investigate the hemocyte production of free-radicals such as nitric oxide and superoxide anion. This approach revealed that *BgBRE* snails produced more free radicals than *BgGUA* snails (Fig. 4A). Moreover, a distribution analysis of free-radical concentrations in single hemocytes showed that the cells producing higher concentrations of ROS came from *BgBRE* individuals, while those producing lower levels of free radicals were from *BgGUA* snails (Fig. 4B).

However, although our results revealed that global ROS production differed between *BgBRE* and *BgGUA*, oxidants can differ in their reactivity and efficient parasite killing requires that the host produce the right oxidant (Bayne et al., 2001). As previous studies have demonstrated the crucial role of hydrogen peroxide (*H*$_2$*O*$_2$) in the killing of *S. mansoni* sporocysts (Hahn et al., 2001b; Goodall et al., 2004; Bender et al., 2005; Bender et al., 2007), we investigated potential differences in hemocyte *H*$_2$*O*$_2$ production between the snail strains. Our results showed that hemocytes from *BgBRE* constitutively produced more *H*$_2$*O*$_2$ than those from *BgGUA* (Fig. 5). We then examined the *H*$_2$*O*$_2$ content of plasma from these snails. As technical restrictions make it impossible to directly measure *H*$_2$*O*$_2$ in plasma, we measured the superoxide anion, which is a precursor of *H*$_2$*O*$_2$ (Selkirk et al., 1998). Our results confirmed that *BgBRE* plasma contained significantly more superoxide anion than *BgGUA* plasma (Fig. 6). All of these data were obtained from hemocytes harvested from uninfected snails and without cell stimulation. Notably, no difference in ROS production was observed when these hemocytes were stimulated by the addition of phorbol 12-myristate 13-acetate (PMA) to culture medium (data not shown).

Taken together, our data show that: (i) the production of ROS in general and *H*$_2$*O*$_2$ (the main ROS acting against *S. mansoni* sporocysts) in particular differ between the two snail strains; and (ii) this *H*$_2$*O*$_2$ production seems to be correlated with the level of ROS scavengers produced by
sympatric parasites. BgBRE snails produce higher amounts of H$_2$O$_2$ and interact naturally with SmBRE, which have better resistance against oxidative stress, while BgGUA snails produce less H$_2$O$_2$ and are sympatric with SmGH2, which is more susceptible to ROS. If our hypothesis is accurate, therefore, we would expect our cross-infection experiments to reveal differences: (i) in the infective potential of our two S. mansoni strains; and (ii) in the resistance potential of our two B. glabrata strains.

Indeed, the results of the infection and cross-infection experiments showed significant differences in the infection rates and intensities (Fig. 7). The factors and mechanisms underlying these differences are not yet known, but may include historical epidemiological conditions, differential selective pressures in the transmission areas, genotypic diversities in the host and parasitic isolates, recognition mechanisms developed through the matching-phenotypes model, and intraspecific competition among sporocysts (for details, see (Théron et al., 1997; Théron and Coustau, 2005; Théron et al., 2008; Bech et al.). Notably, the host-parasite combination characterized by the higher infection rates and parasite intensities (SmBRE/BgBRE) was also characterized by a higher ROS-production capacity by the host and a higher ROS-scavenging ability by the parasite. In contrast, the host-parasite combination with lower infection rates and parasite intensities (SmGH2/BgGUA) showed lower ROS production by the host and lower ROS scavenging by the parasite (Fig. 8). These observations argue for the presence of reciprocal adaptation between the ROS and ROS scavenger traits. This was further supported by the results from our allopatric cross-infections. The SmGH2 strain, which had co-evolved with its sympatric snail (BgGUA) to produce lower levels of ROS, could not effectively infect high-ROS-producing BgBRE snails (Figs. 7 and 8). Conversely, the SmBRE strain, which had co-evolved with a host that produced more ROS (BgBRE), could easily infect low-ROS-producing BgGUA snails (Figs. 7 and 8). Interestingly, however, the infection success of SmBRE was lower for the allopatric combination than the sympatric pairing (Figs. 7 and 8), suggesting that the oxidative factors probably act in combination with other factors to determine the outcome of the B. glabrata/S. mansoni interaction.
Within hosts, immune effectors exert the main selective pressure on parasites (Loker and Adema, 1995; Damian, 1997). However, another factor that helps to define the interaction is the efficiency of parasite recognition by snail immune receptors, and the ability of the parasite to escape this recognition. We previously discovered a group of polymorphic antigens of *S. mansoni* (the *S. mansoni* polymorphic mucins, *Sm*PoMucs) (Roger et al., 2008a; Roger et al., 2008b; Roger et al., 2008c), and recently showed that these antigens are recognized by diversified *B. glabrata* immune receptors (the fibrinogen-related proteins, FREPs) (Moné et al., 2010a). These reports on the molecular interactions underlying snail–schistosome compatibility suggest that co-evolutionary (reciprocal adaptation) processes probably occur through a combination of changes in general resistance (ROS/ROS scavengers) and more specific interactions (FREPs/*Sm*PoMucs). In non-specific resistance/infectivity interactions involving density-dependant forces (e.g., the number of developing parasites within the host), co-evolution leads to global increases in the amount of attack/defense products, such as the interplay of ROS and ROS scavengers described herein. In highly specific genotype-by-genotype interactions, such as recognition/evasion processes, however, co-evolution leads to increases in the diversification and/or polymorphisms among specific molecules, as observed for FREPs and *Sm*PoMucs.

Even if the success of infection is not exclusively based on the levels of ROS and ROS scavengers, our model of dynamic co-evolution predicts that a change in parasite virulence or host resistance would be associated with life history trade-offs (reallocation of resources), with increased production of a molecule under co-evolutionary pressure yielding indirect negative consequences for other functions (development, growth, fecundity, reproductive rate, etc.) (Green et al., 2000; Lohse et al., 2006; Forde et al., 2008). Indeed, this kind of trade-off has been observed in our model, as a previous study showed that cumulative cercarial production was two-fold higher for the *Sm*GH2/*Bg*GUA combination than for the *Sm*BRE/*Bg*BRE (Théron et al., 1997). This could indicate that *Sm*BRE has made a tradeoff by investing in the production of ROS scavengers at the expense of producing cercariae.
In summary, host-parasite interactions are dynamic biological systems in which the host’s
defense mechanisms face the parasite’s infectivity mechanisms, leading to a co-evolutionary arms
race (Combes, 2000; Howard and Jack, 2007).

Developing correlation approaches to studying co-evolution have some limitations. Indeed
the correlations between traits of interacting species cannot always provide unequivocal evidence
for co-evolution. Reciprocity could also occur and an absence of correlated traits is not evidence for
an absence of co-evolution (Nuismer et al., 2007; Nuismer et al., 2010; Yoder and Nuismer, 2010).
Non-co-evolutionary mechanisms could explain correlations between the traits of interacting
species. For example, the correlation could result from a colonization process in which a parasite
species with new potential arrived in a new environment and is more well-adapted to the sympatric
interacting species. In other interaction models correlated traits could evolve if the abiotic or biotic
environments favour similar traits in both of the interacting species. For example, a biotic selection
that affects only one of the interacting species can itself cause trait matching. This can occur if
interactions have potent fitness consequences for only one of the species or if the outcome of
interactions depends on the phenotype of only one of the species. These one-way interactions can
generate correlations that are indistinguishable from those that evolve due to co-evolutionary
processes (Nuismer et al., 2007; Nuismer et al., 2010; Yoder and Nuismer, 2010). In our model of
interest \textit{B. glabrata} could be infected by a lot of pathogens species (other than \textit{S. mansoni}) that
represent a selective pressure that could enhance snail ROS production. In this context,
schistosomes for which the specificity for the intermediate snail host is very high will still succeed
in infecting the snails, only if they are able to circumvent ROS by increasing their ROS scavenger
production.

However, Nuismer et al. (2007, 2010) state that correlation could occur if interactions are
mediated by a mechanism of phenotype matching such as what takes place for host-parasite
interactions. This phenotype matching process was proposed for our \textit{S. mansoni / B. glabrata} model
of interest (Théron and Coustau, 2005).
Therefore whatever are the mechanisms involved in the apparition of trait correlation between two interacting species, our present results reveal the existence of phenotypic matching between host and parasitic strains in terms of their attack (ROS production) and defense (ROS scavenging) traits. To our knowledge, this work provides the first example of a clear link between the level of oxidant and antioxidant molecules possibly resulting from sympatric co-evolution, and provides supporting evidence for a field illustration of the Red Queen Hypothesis (Van Valen, 1974) and its predictions of a functional trait in a metazoan host/parasite model. Detailed mechanistic studies will be conducted in multiple populations to fully confirm the link between correlated traits and the Red Queen context.

Acknowledgments

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**Figure legends**

**Fig. 1.** Effect of hydrogen peroxide (H$_2$O$_2$) on *Schistosoma mansoni* sporocysts. (A) Cytotoxicity among *S. mansoni* sporocysts 2 h after exposure to different H$_2$O$_2$ concentrations. The asterisk indicates a significant difference ($P < 0.05$) in the cytotoxic effect of H$_2$O$_2$ on *S. mansoni* Guadeloupean strain (*Sm*GH2) versus *S. mansoni* Brazilian strain (*Sm*BRE) sporocysts. (B) Percent sporocyst mortality after 4 h exposure to different H$_2$O$_2$ concentrations. The asterisk indicates a significant difference ($P < 0.05$) in the mortality rates of *Sm*GH2 versus *Sm*BRE sporocysts.

**Fig. 2.** Constitutive total antioxidant capacities of *Schistosoma mansoni* Guadeloupean strain (*Sm*GH2) and *S. mansoni* Brazilian strain (*Sm*BRE) sporocysts. Values are expressed as µM of antioxidant activity per 10 µg of sporocyst proteins. The asterisk indicates a significant difference ($P < 0.05$).

**Fig. 3.** Ratios of Zn-Cu superoxide dismutase (Zn-Cu SOD, Smp_176200.2) transcript levels in the two strains of *Schistosoma mansoni* miracidia (Guadeloupean strain, *Sm*GH2 and Brazilian strain, *Sm*BRE). Ratios were determined using real-time quantitative PCR and are expressed relative to the expression levels of 28s rRNA and α-tubulin. The histogram represents the average values of duplicates ± S.D. The expression ratio was calculated according to the formula: $R = 2^{(ΔCt)}$, where $ΔCt$ represents Ct (target gene) – Ct (constitutively expressed gene).
Fig. 4. Intracellular ROS measurements in single *Biomphalaria glabrata* hemocytes (A) Global reactive oxygen species (ROS) concentration in each snail strain. The histogram represents the ROS concentrations in arbitrary units (-fold mean) for the *Biomphalaria glabrata* Guadeloupean strain, BgGUA and Brazilian strain, BgBRE. The asterisk indicates a significant difference (*P* < 0.05) in ROS production by hemocytes of the two snail strains. (B) ROS concentrations in hemocyte populations from BgGUA and BgBRE snails. The histograms represent the fluorescent lifetimes of 1-pyrenebutyric acid (PBA)-loaded hemocytes from eight each of BgGUA and BgBRE; 108 single hemocytes from eight BgBRE and 111 hemocytes from eight individuals of BgGUA snails were assessed. The *x* axis represents the fluorescent lifetime in nanoseconds, while the *y* axis corresponds to the number of cells.

Fig. 5. Hydrogen peroxide (H$_2$O$_2$) production by *Biomphalaria glabrata* hemocytes. Cumulative production of H$_2$O$_2$ was measured using Amplex Red. The data are presented as the mean (± S.D.) of Amplex Red absorbance at 570 nm ($A_{570nm}$) per 45,000 cells over five replicates. The asterisk indicates a significant difference (*P* < 0.05) in H$_2$O$_2$ production from hemocytes of *B. glabrata* Guadeloupean strain, BgGUA, versus Brazilian strain, BgBRE.

Fig. 6. Constitutive superoxide anion plasma content in *Biomphalaria glabrata* Guadeloupean strain, BgGUA, versus Brazilian strain, BgBRE. The superoxide anion plasma content was assessed by spectrophotometric measurement (620 nm) of nitroblue tetrazolium (NBT) reduction. At 3 h after initiation of the reaction, the BgBRE plasma contained significantly more superoxide anion than that from BgGUA (the asterisk indicates a significant difference; *P* < 0.05).
Fig. 7. Infection rates and intensities in sympatric and allopatric *Schistosoma mansoni/Biomphalaria glabrata* combinations. The percentage of snails infected and the intensity of infection: number of mother sporocysts (SpI) developed (*n* SpI) was measured after individual snails were challenged with different miracidial doses (1, 10, 20, 30 or 50 miracidia (Mi)).

Fig. 8. Schematic representation of our reactive oxygen species (ROS)-based co-evolutionary hypothesis. The percentage of prevalence is indicated for each *Biomphalaria glabrata/Schistosoma mansoni* combination. The number of arrows represents the differential host oxidant (ROS) or parasite antioxidant (ROS scavenger) capabilities. *Biomphalaria glabrata* Brazilian strain, *Bg*BRE, and Guadeloupean strain, *Bg*GUA; *S. mansoni* Brazilian strain, *Sm*BRE, and Guadeloupean strain, *Sm*GH2.
Table 1
Primer sequences for Reverse Transcription-quantitative PCR in this study.

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<tr>
<th>Gene Name and SchistoDB ID</th>
<th>Amplicon Length</th>
<th>Smp_scaffold</th>
<th>Forward primer (5' to 3')</th>
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<tr>
<td>Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) Smp_056970</td>
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<td>000155</td>
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<tr>
<td>Glutathione S-transferase omega Smp_152710</td>
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<td>000154</td>
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<tr>
<td>Glutathione S-transferase 28 kD (GST 28) Smp_054160</td>
<td>128</td>
<td>000143</td>
<td>CGGACCGGACGTGCTGAAT</td>
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<td>Glutathione-S-transferase 26 kD (GST 26) Smp_163610</td>
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</table>

Gene names are given according to the SchistoDB accession numbers (http://schistodb.net/schistodb20/). Their respective scaffolds are included in the table. Smp_163610 primer sequences are given according to the mRNA sequence of the gene (XM_002582157.1) due to inconsistency in the genome assembly. α-tubulin and 28S primers sequences were previously published (Bahia et al., 2006; Roger et al., 2008a).
Antioxidant capacity (µM / 10µg protein)

<table>
<thead>
<tr>
<th></th>
<th>SmGH2</th>
<th>SmBRE</th>
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</thead>
<tbody>
<tr>
<td>Sporocyst</td>
<td>0.21</td>
<td>0.24</td>
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</table>

* indicates a significant difference between the two groups.