Trypanosoma cruzi Immune Response Modulation Decreases Microbiota in Rhodnius prolixus Gut and Is Crucial for Parasite Survival and Development

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Abstract

Trypanosoma cruzi in order to complete its development in the digestive tract of Rhodnius prolixus needs to overcome the immune activity reactions and microbiota trypanolytic activity of the gut. We demonstrate that in R. prolixus following infection with epimastigotes of Trypanosoma cruzi clone Dm28c and, in comparison with uninfected control insects, the midgut contained (i) fewer bacteria, (ii) higher parasite numbers, and (iii) reduced nitrite and nitrate production and increased phenoloxidase and antibacterial activities. In addition, in insects pre-treated with antibiotic and then infected with Dm28c, there were also reduced bacteria numbers in a midgut, and a higher parasitic load compared with insects solely infected with parasites. Furthermore, and in contrast to insects infected with Dm28c, infection with T. cruzi Y strain resulted in a slight decreased numbers of gut bacteria but not sufficient to mediate a successful parasite infection. We conclude that infection of R. prolixus with the T. cruzi Dm28c clone modifies the host gut immune responses to decrease the microbiota population and these changes are crucial for the parasite development in the insect gut.

Introduction

Chagas disease is an endemic parasitic disease important in large areas of Latin America [1]. The majority of triatomine species found in this region are potential vectors for Trypanosoma cruzi, the causative agent of the disease [1,2]. Rhodnius prolixus is the main vector of Chagas disease in the northern part of South America and in some areas of Central America [3].

The T. cruzi – triatomine vector interactions are complex and mediated by numerous immune and insect factors [4–8]. The parasite circulates between humans, domestic and sylvatic mammalian reservoirs, and insect vectors [6–8]. T. cruzi infection of the insect vector starts with the ingestion of an infective blood meal. A successful transmission depends upon the parasite multiplication and differentiation in the triatomine gut [7–9].

Many factors have been implicated in the establishment of T. cruzi infection in the insects vector gut including a crop lytic factor, lectins, gut enzymes and microbiota-derived factors, antimicrobial compounds and nitric oxide [4,6–12]. In addition, the gut structural organization is important for parasite attachment to the perimicrovillar membranes and their subsequent development [5,9,13]. The success of parasite infection also depends upon the T. cruzi strain used and the insects species infected. For example, in R. prolixus, the T. cruzi Dm28c clone and CL strains are capable of completing their developmental cycles in contrast to the Y strain which rapidly disappears from the insect’s gut [14,15].

Regarding the role of the diverse gut microbiota in modulating T. cruzi development in the gut of R. prolixus, we have previously identified a Serratia sp. isolated from this insect vector which interferes with some strains of parasite development and lyces parasites in in vitro experiments [7,14,16–18]. The microbiota has also previously been shown to modulate the vectorial competence of the insect host by means of direct interaction with parasites or by competition for resources in the gut [7–11]. In addition, microbiota can constrain pathogen development indirectly by inducing vector antiparasitic activity and humoral immune defense factors in particular [9,12,16,19–21].

Humoral defenses in insects are characterized by a battery of potent antimicrobial peptides (AMPs), reactive intermediates of nitrogen or oxygen, and complex enzymatic cascades, such as the phenoxdiazide system, that contribute to clotting or hemolymph melanization [22]. These humoral factors may be secreted directly into the hemocoele or into the gut lumen as components of the immune response to eliminate potential pathogens acquired during feeding [7,8,23].

In this paper, we have examined the relationship in the gut of R. prolixus between the microbiota and the insect’s humoral antibacterial compounds with the level of infection following
feeding with *T. cruzi*. In particular, we investigated the role of *T. cruzi* infection in the modulation of the microbiota population in the *R. prolixus* gut. We demonstrated, for the first time, that infection with *T. cruzi* Dm28c clone, but not with the Y strain, changes the microbiota population in the digestive tract by modulating the host immune responses and that this contributes to parasite development in the gut of *R. prolixus*.

**Results**

**Trypanosoma cruzi** infection and microbiota population in the digestive tract

Parasite infection and microbiota population in *R. prolixus* 5th-instar nymphs whole digestive tract were analyzed from 5 to 29 days after treatment. In this period, the median number of *T. cruzi* Dm28c clone in the whole digestive tract (CC) ranged from 1 to 3 × 10^1 parasites/ml (Figure 1A). Regarding the microbiota, the median number of gut bacteria in the control insects fed blood alone (C) showed a rapid increase and by 8 days after feeding reached a peak of 1.4 × 10^11 before reducing to 8 × 10^10 bacterial/digestive tract by day 29 after feeding (Figure 1B) and then remained stable in numbers until the next feeding. Significant differences in bacterial numbers between parasite-infected (CC) and control (C) groups were observed at 8 days after initiating the experiments. At this time in control insects, the microbiota reached 1.4 × 10^14 bacteria (CFU) whereas parasite-infected insects presented only 1.3 × 10^9 CFU/digestive tract (Figures 1B, 1C; p < 0.001).

To investigate the role of the reduction of the microbiota upon *T. cruzi* Dm28c infection in *R. prolixus*, one group of 5th instar nymphs was treated with antibiotic solution alone added to the blood meal (A), and another group of insects was treated with antibiotic and then infected with parasites (AC). With this non-toxic dose of antibiotic in the blood meal, we observed an increase in *T. cruzi* Dm28c infection (AC), in comparison with insects fed with parasites alone (CC), ranging from 3.2 × 10^5 parasites/ml to 1.5 × 10^7 from 8–13 days to 25–29 days, respectively, after infective feeding (Figure 1A). Antibiotic treated and infected insects (AC) presented 10 times more parasites than in control insects (C) after 11 h, a significant difference in antibacterial activity was observed in the anterior midgut of control insects (C) only after 4, 5 and 6 h of incubation (Figure 3B; p < 0.05). For example, parasite-infected insects (CC), after 4 h incubation, recorded an antibacterial activity of 10.92 (±0.71), significantly higher than control groups (C) with activity of 7.45 (±0.47) (Figure 3C; p = 0.001). These results were confirmed by following bacterial growth on BHI-agar plates where control (C) and infected insects (CC) presented low numbers of bacteria (not shown).

To test the hypothesis that the higher parasitemia and lower microbiota recorded in antibiotic-treated insects resulted from modulation of the antibacterial immune responses was also investigated further. ZI assays with insects treated just with antibiotics (A) or with antibiotics and then infected with parasites (AC) showed inhibition zones much bigger than with the non-antibiotic C and CC groups with diameters larger than 2.0 cm after 18 h but these could not be measured accurately as the zones overlapped with adjacent samples (not shown). In TB assays, the antibiotic-treated and infected insects (AC) had the highest activity (Figure 1C; p > 0.1).

The results of the ZI assays showed significantly smaller zones of inhibition (ZI) in control insects (C) (0.78 cm ± 0.03) when compared with control infected insects (CC) (1.0 cm ± 0.02) (Figure 3A; p < 0.0001). The effect of *T. cruzi* Dm28c infection on antibacterial activities in the anterior midgut was also tested using a turbidimetric assay (TB). Experiments incubating the extracts with bacteria for 11 h at 37°C showed significantly higher antibacterial activity in infected insects (CC) compared with control insects (C) only after 4, 5 and 6 h of incubation (Figure 3B; p < 0.05). As expected, upon antibiotic treatment alone (A), the bacterium population was significantly reduced (p < 0.001) compared with the controls (C) (Figure 1C). In the group of insects treated with antibiotic and infected with parasites (AC), we noted few bacteria in the digestive tract as in the group treated with antibiotic alone (A) (Figure 1C; p > 0.1).

**Bacterial growth inhibition**

The anterior midgut antimicrobial activities at 9 days after feeding in the ZI assay (0.84 ± 0.04 cm diameter) were significantly higher than posterior midgut (0.09 ± 0.09 cm diameter) of control infected insects (CC) (Figure 2A; p < 0.001). Therefore, the bacterial growth inhibition and immune reactions experiments were standardized using samples of anterior midgut. We also compared the ZI assay, the antibacterial activities of the anterior midgut at days 5, 9 and 16 after infective feeding (CC), and observed that day 9 activity (0.84 ± 0.04 cm diameter) was significantly higher compared with day 5 (0.67 ± 0.02 cm diameter) and day 16 (0.64 ± 0.02 cm diameter) (Figure 2B; p < 0.001).

**Gut antimicrobial activities**

In order to obtain a better understanding of the regulation of microbiota in the vector, we investigated the antibacterial activities in the anterior midgut of control *R. prolixus* (C) and insects infected with *T. cruzi* Dm28c clone (CC) were investigated.

The results of the ZI assays showed significantly smaller zones of inhibition (ZI) in control insects (C) (0.78 cm ± 0.03) when compared with control infected insects (CC) (1.0 cm ± 0.02) (Figure 3A; p < 0.0001). The effect of *T. cruzi* Dm28c infection on antibacterial activities in the anterior midgut was also tested using a turbidimetric assay (TB). Experiments incubating the extracts with bacteria for 11 h at 37°C showed significantly higher antibacterial activity in infected insects (CC) compared with control insects (C) only after 4, 5 and 6 h of incubation (Figure 3B; p < 0.05).

**Phenoloxidase (PO) activities**

PO activities were also measured in the gut of *R. prolixus* 5th instar nymphs 9 days after feeding. PO specific activities were the highest in the insects infected with *T. cruzi* Dm28c clone (CC) (0.024 abs/mg protein) compared with the control group (C) (0.012 abs/mg protein) (Figure 4A; p < 0.001). Antibiotic-treated and infected insects (AC) recorded results (0.016 abs/mg protein) higher than insects treated with antibiotic alone (A) (0.006 abs/mg protein)
Figure 1. Parasite infection and microbiota population in *Rhodnius prolixus* 5th-instar nymphs digestive tract challenged by *Trypanosoma cruzi* Dm28c clone. (A) Parasite infection at different days after infection. (B) Microbiota population at different days after feeding. (C) Microbiota population 8 days after feeding. Treatments: C – control insects fed on blood alone; CC – insects infected with *T. cruzi* Dm28c clone; A – insects treated with antibiotic alone; AC – insects treated with antibiotic and infected with the Dm28c parasites; Y – insects infected with *T. cruzi* Y strain; AY – insects treated with antibiotic and infected with *T. cruzi* Y strain. In figures A and C each point represents the number of parasites or
bacteria in an individual digestive tract, and horizontal lines indicate the median. In figure B each point represents the median. In figure C the median for antibiotic treated and infected insects (AC) is zero and therefore overlaps the x-axes. Treatments were repeated 3–5 times with 6–10 insects in each experiment reaching a total of 25 to 35 insects for each group. Medians were analyzed with 1 way ANOVA and Mann Whitney test.

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Figure 2. Antibacterial activity using inhibition zone (ZI) assay in Trypanosoma cruzi Dm28c clone. (A) Antibacterial activity with anterior and posterior midgut regions at 9 days after feeding; (B) Antibacterial activity of anterior midgut at 5, 9 and 16 days after feeding. Bars represent mean ± SEM. Means were analyzed by using t Test and 1 way ANOVA. Each bar represents the mean (+/− SEM) of four separate experiment, n = 6–10 insects for each determination.

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Discussion

In triatomeine insects, there is a close association of gut microorganisms with parasites. Different parasite species or strains infecting Rhodnius prolixus can increase or decrease the midgut bacterial flora. Thus, Eichler and Schaub [19] showed that R. prolixus infected with T. rangeli strain Choachi, but not with T. cruzi strain ‘Chile 5’, reduced the symbiont Rhodococcus rhodnii concentration by 50%. Furthermore, we demonstrate herein that T. cruzi Dm28c clone establishes a parasite infection and reduces R. prolixus microbiota while strain Y fails to infect R. prolixus and the microbiota remain at significantly higher levels than with the Dm28c clone. These observations on parasite and insect microbiota interactions are probably related to the success of parasite development in insects. The difference of our results from Eichler and Schaub may be related to different strains/clones of R. prolixus [19]. In addition, hemolytic bacteria isolated from the stomach of R. prolixus, and identified as Serratia marcescens, killed epimastigotes of T. cruzi Y but not the Dm28c strain [14,16]. The S. marcescens lytic activity against T. cruzi Y strain is inhibited by D-mannose since the bacteria possess mannose-sensitive fimbriae that mediate adherence and lysis of trypanosomes [17,18]. This intimate interaction of gut microorganisms with parasites emphasizes both the complexity of the triatomeine gut environment and the importance of the maintenance of gut homeostasis since many bacteria, fungi and parasites, such as T. cruzi, survive and develop in the digestive tract without killing the vector host [7,8,15,19].

One remarkable result of infecting R. prolixus with T. cruzi Dm28c was the significant decrease in the vector gut microbiota. This was probably related to increases in the antibacterial and PO activities in the gut of infected insects. It is likely that T. cruzi Dm28c succeeds in completing development and successfully establishes infection in the vector as a consequence of these changes. In mosquitoes too there is a correlation between microbiota and parasite development [24,25]. Challenges Anopheles stephensi, An. gambiae, and An. albimanus with bacteria significantly reduces oocyst infection rates and densities in Plasmodium falciparum-infected mosquito cohorts [24]. Subsequently, studies demonstrated that An. gambiae microbial flora up-regulates immune genes related to anti-Plasmodium factors and in microbe-free aseptic mosquitoes there is an increase in susceptibility to Plasmodium infection [25]. Also, in Tsetse flies, bacterial infection enhanced immunity in insects and blocked the trypanosome’s ability to establish infections, indicating that bacteria can act to block parasite transmission by up-regulation of some immune responsive genes [26].

Pre-treating the insects with antibiotics before infecting with T. cruzi clone Dm28c demonstrates the influence of the insect’s microbiota on parasite infection, since a reduced microbiota population caused an increase in parasite numbers. Similarly, in An. gambiae, antibiotic-treated insects became more susceptible to P. falciparum infection compared with the control septic mosquitoes [25]. In addition, the immune reactions of R. prolixus treated with antibiotics were altered in consequence of the higher parasitemia or the reduced microbiota. Thus, antibacterial activity was higher in insects treated with antibiotics and infected with parasites in comparison to both control and infected control insects. However, both nitrite/nitrate production and phenoloxidase were decreased in insects receiving antibiotic alone and in those given antibiotic and then infected with T. cruzi Dm28c.

Regarding nitric oxide (NO) production, a small but significant decrease was observed in the levels of nitrite/nitrate in insects infected with T. cruzi Dm28c. It is generally recognized that NO

Nitrite/nitrate productions

The last immune reaction investigated for modulation by the T. cruzi Dm28c infection was the production of NO (nitrite/nitrate) in the gut. This activity showed a significant decrease in the infected group (CC) compared with the controls (C) and the levels varied from 10.3 μMols/mg protein to 11.5 μMols/mg protein, respectively (Figure 4B; p<0.001). In antibiotic-treated insects, NO (nitrite/nitrate) levels were also lower for infected insects so that in antibiotic-treated and infected insects (AC) 7.2 μMols/mg protein were recorded compared with 8.6 μMols/mg protein for the insects treated with antibiotic alone (A) (Figure 4B; p<0.001).
reactions together with oxygen intermediates and derivatives have antiparasitic effects [27–29]. Inducible NO has been shown to modulate R. prolixus responses to infection by T. rangeli in the hemolymph [30] and also demonstrate that NO regulates too the infection by T. cruzi in Rhodnius [31]. Whitten et al. [31] hypothesized that NO synthase gene expression and NO production may be modulated differently by T. rangeli and T. cruzi infections in various tissues of R. prolixus. Their findings showed that in R. prolixus infected with T. cruzi presented high nitrite concentration in the crop after 1-2 days of blood feeding but nitrite levels decreased after 2 weeks of parasite infection [31]. Our results of low nitrite and nitrate production in the crop of R. prolixus infected by T. cruzi after nine days of infection are in agreement with these data and demonstrate the complexity of the modulation of NO in this insect. Alterations in NO-derived substances or NOS expression in response to infection have also been described in mosquitoes infected with Plasmodium sp [32,33]. Particularly interesting too was the significant enhanced antibacterial activity and high PO levels in the anterior midgut at day nine after T. cruzi Dm28c infection as these paralleled...
decreases in the microbiota population at this time. The presence of PO in the gut epithelial membrane of Rhodnius prolixus with higher levels in the anterior midgut was also identified by Genta et al. [22]. A previous report has also shown that two strains of T. rangeli parasites can differentially activate the prophenoloxidase system in hemolymph in vivo and that this activation may be involved in modulation of the infections [31]. In the present study, a non-invasive parasite, T. cruzi Dm28c, stimulates the PO activity in the digestive tract but the parasite population is refractory to killing and completes development in the insect vector. Therefore, the PO, at the concentration recorded, may not significantly inhibit parasite growth, but may differentially kill bacteria that may potentially lyse and compete with the parasites.

Regarding the role of the enhanced antimicrobial activity recorded for the T. cruzi Dm28c-infected R. prolixus, triatomine vectors have been shown to produce endogenous antimicrobial peptides (defensin and lysozyme) in the gut/fat body following infection with T. cruzi or bacteria [34,35]. Waniek et al. [36] demonstrated a marked increase of a defensin transcript in Triatoma brasiliensis in response to T. cruzi infection and suggested a potential function of gut defensin in parasite population control. Insect antibacterial peptides are often described as molecules that kill bacteria and fungi infections and, as such, they potentially could protect T. cruzi from these microorganisms. Direct killing activities have been recorded for some antibacterial compounds against parasites [12,35]. In these studies, however, high concentrations of heterologous antibacterial peptides have often been used in in vitro experiments and T. cruzi in the insect vector might never be exposed to such high levels of these compounds or these activities may be affected by gut digestive protease.

Even more significant is the recent description of a novel antimicrobial peptide, prolixin, from R. prolixus with activity against both Gram-negative and Gram-positive bacteria but with no toxicity for T. cruzi [37]. This factor may well explain, in the present work, the elevated levels of antibacterial activity recorded in T. cruzi infected R. prolixus with the differential capability of killing the midgut microbiota but not the T. cruzi parasites.

In experimental infections, not all T. cruzi strains succeed to grow and complete development in all species of triatomines [15]. Therefore, to test the hypothesis that the parasites infection success is closely related to the microbiota population, we infected R. prolixus with T. cruzi Y strain which is incapable of completing development in Rhodnius digestive tract [14,16]. The results showed, in contrast to T. cruzi Dm28c that T. cruzi Y strain could not infect R. prolixus and also failed to reduce the microbiota population of the gut. Even when the microbiota population was reduced by treating the insects with antibiotics, T. cruzi Y strain did not successfully infect the insects gut. It therefore seems that the parasite infection depends on both the parasite’s ability to diminish the bacteria population, which may have trypanolytic activity, and also to modulate the insect’s immune reactions.

The present study demonstrates that the activation of PO and antibacterial activities, and the reduction of NO, are related to decreases in the microbiota and increases in T. cruzi Dm28c density in the gut. These data, together with previous studies showing other factors that can influence the establishment of parasite infection, demonstrate the molecular complexity involving the interactions of T. cruzi with R. prolixus.

Materials and Methods

Parasite infection, microbiota population and immune responses in R. prolixus 5th-instar nymphs digestive tract were analyzed from 5 to 29 days after treatment.

Ethics Statement

Defibrinated rabbit blood was provided by the Laboratory Animals Creation Center (Cecal) which breed and maintain animals for the laboratory. All research programs in Fiocruz respect the guidelines of the Ethics Committee on Animal Use (Cecal) composed by Fiocruz researchers and external consultants.

Insects treatment

Fifth-instar Rhodnius prolixus nymphs were obtained from a colony reared and maintained in our laboratory at a relative humidity of 50–60% and at 27±2°C. After molting, insects starved for 15–20 days and weighing 35.2±3.4 mg, were randomly chosen and then fed with defibrinated rabbit blood through a membrane feeding apparatus [38]. A control group (C) was fed on blood alone and the infected groups on blood containing 1×10⁵ Trypanosoma cruzi Dm28c clone/ml of blood (CC) or Y strain epimastigotes (Y). Insect vector infection by artificial blood feeding is successful when using the epimastigote parasite form which is naturally encountered inside the midgut and capable of adhesion to the gut epithelial cells [9,16]. In some experiments, groups of insects were fed on blood containing a mixture of three different antibiotics (A), gentamicin, penicillin and streptomycin, with final concentrations of 450, 300 and 500 μg/ml of blood, respectively. Another group of insects were fed on blood containing these antibiotics together with 1×10⁵ T. cruzi Dm28c epimastigotes/ml of blood (AC). Only fully engorged insects were used in experiments, which fed upon 160.5±22.1 mg of blood that represents approximately 2.0±10⁵ T. cruzi Dm28c epimastigotes/infected insects. All insects were raised and maintained as previously described [39].

Parasites

Trypanosoma cruzi Dm28c clone and Y strain parasites are maintained in our laboratory and grown in a brain heart infusion (BHI, DIFCO) culture medium supplemented with 10% heat-inactivated fetal calf serum at 28°C, according to Azambuja and Garcia [36]. The epimastigotes (99% purity) were obtained from the log-growth phase of the parasites [present up to day 7 of cultivation] [40]. To determine parasite infection in insects, the whole digestive tract was homogenized in 1 ml of sterile phosphate saline buffer (PBS) and parasites counted directly in a hemocytometer.

Microbiota – Colony Forming Unit (CFU) assay

The microbiota of R. prolixus digestive tract were assessed by counting colony forming units (CFU) that grew in brain heart infusion agar (BHI agar). The entire digestive tract was dissected in sterile conditions after different days of treatments and homogenized in 1 ml of sterile phosphate saline buffer (PBS). Samples were then immediately transferred to ice, diluted 10⁻⁵, 10⁻⁷ or 10⁻⁹, with PBS, 20 μl aliquots spread onto BHI agar plates and then incubated overnight at 30°C and CFUs subsequently counted. A PBS aliquot was also plated as a control to guarantee sterility. The insects, R. prolixus, maintained in our colony present a natural microbiota flora which contains different bacteria species including Serratia marcescens strain. These insects were used in our experiments without infecting with other bacteria.

Antibacterial assays

To analyze the antibacterial activity in anterior and posterior midguts, tissues were dissected and disrupted in 0.1% Triton X-100 (v/v) at 4°C. The samples were centrifuged at 8,000 g for 1 min at 4°C. Aliquots were taken from the supernatant for each assay, as described below.
Zone of inhibition (ZI) assay. Aliquots of 20 μl from the R. prolixus gut samples collected and prepared, as described above, were kept frozen at −20°C for no more than 2 weeks. The samples (5 μl) were dispensed in triplicate on to sterile filter paper discs, arranged on the surface of soft agar plates and incubated at 37°C for 18 h. Plates were prepared using 1% agarose in liquid culture medium (BHI) containing streptomycin (100 μg/ml) and ampicillin (80 μg/ml) and Escherichia coli D31 incorporated into the agar. Antibacterial activity of the gut samples was recorded as diameters (cm) of the growth inhibition zones of E. coli D31 on the agarose plates. Preliminary tests, undertaken to assess the antibacterial activities of anterior and posterior midgut samples at different days after treatment, showed the highest activity (ZI) in the anterior midgut nine days after feeding. Therefore, all experiments investigating the insects immune responses were standardized using samples of the anterior midguts (tissues and contents) taken nine days after treatments.

Turbidometric (TB) assay. The TB assay was modified from Thomas et al. [41] and Bexfield et al. [42]. Anterior midgut samples, nine days after treatment, as described above, were centrifuged at 8,000 g for 10 min at 4°C and 70 μl of supernatant transferred into tubes containing 630 μl of Milli-Q water. All samples were filter sterilized and frozen at −20°C. The assay involved incubating the midgut samples with 1% peptone plus the test bacteria, Escherichia coli k12. E. coli k12 were grown in 20 ml tryptone soy buffer (TSB) for 17 h at 30°C with oscillation and then 100 μl of the bacteria were transferred to 10 ml of TSB and incubated for a further 4 h. The bacteria were then washed in PBS and diluted in TSB to a final concentration of 1×10⁵ cells/ml. Fifty microlitres of midgut sample were then incubated with 10 μl of bacterial suspension in triplicate in the wells of a sterile flat-bottom, 96-well microtiter plate (Nunc, Fisher Scientific UK, Leicestershire, UK). Control wells without midgut extracts contained (1) bacteria, peptone and antibiotic [streptomycin (100 μg/ml) and ampicillin (80 μg/ml)]; (2) bacteria and peptone; or (3) peptone alone. The optical densities were measured at 550 nm (OD₅₅₀) during 12 h incubation at 37°C and read at hourly intervals from time zero. All data points were subsequently blanked against time zero to account for the opacity of the midgut samples and then the bacteria E. coli k12 readings were subtracted from all sample readings and multiplied by 100.

Anterior midgut samples (50 μl), from control insects (C) and infected insects (CC), were also incubated with 10 μl of E. coli k12 (1×10⁵ cells/ml) and 10 μl of peptone, in tubes at 37°C, concurrently with the TB assay. At different time during incubation, samples were plated onto BHI-agar to compare the bacterial growth with the readings in the TB assay.

Phenoloxidase (PO) activating assay
Anterior midgut samples freshly prepared were collected at nine days after insect treatments, as described above, and 10 μl of supernatant were transferred into tubes containing 90 μl of Milli-Q water. To determine the phenoloxidase (PO) activity in anterior midgut portions, all homogenates used for enzymatic assays were freshly prepared and PO activity was determined by measuring the production of dopachrome from DOPA [23,43,44]. The assay was prepared by incubating 25 μl of sample in triplicate with 10 μl of 10 mM sodium cacodylate pH 7.4 containing 10 mM CaCl₂ buffer and 25 μl of a saturated solution of DOPA (4 mg/ml). The absorbance at 490 nm was measured in a microplate reader for 120 min at 37°C with readings taken every 15 min. The values of enzymatic activity presented are expressed as abs/mg protein and refer to specific PO activity by protein concentration in anterior midgut homogenates multiplied by 100.

Nitrite and nitrate determinations
The anterior midgut samples were freshly prepared and collected at nine days after insect treatments, as described above, and 10 μl of supernatant were diluted in 90 μl of Milli-Q water. Nitrate and nitrate contents of samples, which can be indicative of reactive nitrogen intermediate (RNI) metabolism, were analyzed following the manufacturer’s instructions using the Griess Reagent System Assay Kit (Promega, WI, USA) [45], and absorbance of the product was measured at 550 nm. Nitrite and nitrate contents were quantified as µmoles using a range of sodium nitrate standards and the specific activity was calculated mg/ml of protein concentration in anterior midgut samples.

Protein determination
Protein content of samples was quantified with a protein assay kit (BCA* Protein Assay Reagent, Pierce, USA) using bovine serum albumin (BSA) standards.

Statistical analyses
The results were analyzed with GraphPad Prism 5 using 1 Way ANOVA or unpaired T test, or Mann Whitney test (non-parametric test) depending on the data distribution and number of treatments. Data are reported as mean ± standard error (SE) or as individual values with medians for parasite and microbiota populations. Differences among groups were considered not statistically significant when p>0.05. Probability levels are specified in the text and Figure legends.

Author Contributions
Conceived and designed the experiments: DPC PA NAR ESG. Performed the experiments: DPC CSM MSG. Analyzed the data: DPC ESG NAR PA. Contributed reagents/materials/analysis tools: DPC PA ESG. Wrote the paper: DPC ESG NAR.

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