Full-title: The immunoglobulin G antibody response to malaria merozoite antigens in asymptomatic children co-infected with malaria and intestinal parasites

--Manuscript Draft--

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| Article Type:      | Research Article   |
| Full Title:        | Full-title: The immunoglobulin G antibody response to malaria merozoite antigens in asymptomatic children co-infected with malaria and intestinal parasites |
| Short Title:       | Malaria antibodies in children with intestinal parasites |
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| Keywords:          | malaria, intestinal-parasites, antibody, Giardia lamblia, Entamoeba histolytica |
| Abstract:          | BackgroundCo-infection with malaria and intestinal parasites is common in children in Africa and may affect their immune response to a malaria parasite infection. Prior studies suggest that co-infections may lead to increased susceptibility to malaria infection and disease severity; however, other studies have shown the reverse. Knowledge on how co-morbidities specifically affect the immune response to malaria antigens is limited. Therefore, this study sought to determine the prevalence of co-infection of malaria and intestinal parasites and its association with antibody levels to malaria merozoite antigens. Methods A cross sectional study was carried out in two villages with high transmission of malaria in Cameroon (Ngali II and Mfou). Children aged 1-15 years were enrolled after obtaining parental consent. A malaria rapid diagnostic test was used on site. Four (4) ml of peripheral blood was collected from each participant to determine Plasmodium falciparum infections by microscopy, haemoglobin levels and serology. Fresh stool samples were collected and examined by wet mount, Kato-Katz method and modified Ritchie concentration techniques. A Multiplex Analyte Platform assay was used to measure antibody levels. ResultsA total of 320 children were enrolled. The prevalence of malaria by blood smear was 76.3% (244/320) and prevalence of malaria and intestinal parasites was 16.9% (54/320). Malaria prevalence was highest in young children; whereas, intestinal parasites (IP+) were not present until after 3 years of age. All children positive for malaria had antibodies to MSP142, MSP2, MSP3 and EBA175. No difference in antibody levels in children with malaria-co infections compared to malaria alone were found, except for antibody levels to EBA-175 were higher in children co-infected with intestinal protozoa (p = 0.018), especially those with Entamoeba histolytica infections (p=0.0026). Conclusion Antibody levels to EBA175 were significantly higher in children co-infected with malaria and E. histolytica compared to children infected with malaria alone. It is important to further investigate why and how the presence of these protozoans can modulate the immune response to malaria antigens. |
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Rose G.F. Leke |
| Response to Reviewers: | Reviewer #1: Dr. Mbe-cho and colleagues sought to determine the prevalence of co-infection of malaria and intestinal parasites and its association with antibody levels to malaria merozoite antigens. The authors report that there was no difference in antibody prevalence or levels in malaria-infected and co-infected children, except antibody |
levels to EBA-175 were significantly higher in children co-infected with malaria and E. histolytica. Overall, the study is well-designed but these results do not significantly alter or impact our understanding of the association of malaria and helminths on antibody to malaria merozoite antigens.

1. The limitation of the study is that the parasite testing in children was not followed by sensitive diagnostic techniques like PCR, and light infections may have been missed which may have resulted in misclassification of the groups. Light infections may boost the antibody responses while children remain asymptomatic.

Reply: We understand the concern. When the study was conducted (2017) in the rural villages, the prevalence of slide-positive malaria was 75.6%. In a prior study conducted in the village (Leke et al 2010), an equivalent prevalence was found of P. falciparum (50-85%) in children aged 5-15 years over a 5-year period. The estimated entomological inoculation rate (EIR) was 0.7 infectious bites/person/nightly thought out the year (~257 IB/P/Y). Based on the more recent malaria prevalence, it appears that the current EIR is similar. Thus, children were most likely being bitten approximately every-other night by an infectious mosquito, since bednets were not routinely used. With this high level of transmission, most of the slide-negative children would be PCR-positive for malaria, i.e., have enough immunity to reduce malaria to submicroscopic levels. Unfortunately, in very high transmission areas like the one reported herein, everyone will have some circulating P. falciparum parasites. So, classifying subjects as slide-positive vs slide-negative may not reflect presence/absence of parasites, but provide information on the immune status of the person. In revising the MS, information from the study by Leke et al. was included as well as a discussion of submicroscopic infections in the revised Discussion.

2. In this study, only 3.4% children were infected with helminths alone to get any meaningful data for antibody response to malaria in this group.

Reply: We agree, the sample size of children with helminth infections is too small to provide meaningful information. Accordingly, Ab levels in children with helminth infections were not analyzed. To explain the low prevalence of helminths, information on the Ministry of Health’s policy for biannual treatment of children for worms was provided.

3. Very few children are positive for E. histolytica.

Reply: True, the prevalence of Entamoeba in our study was only 5.9%, which is lower than that reported in studies in these areas of ~23% (T. E. Kwenti et al., 2016). In our study, the prevalence was lower, probably due to rigorous mass drug administration (MDA) programs implemented by the Ministry of Health and other regular or seasonal health campaigns.

4. The data on the children's anthropomorphic measurements are not mentioned. Thus, there is not much point describing how they were collected.

Reply: This section was removed from the Methods section.

5. There is no data on hookworm infection in the results.

Reply: The prevalence of hookworm infections was considered in this study during stool exams and, surprisingly, we did not find hookworms in the samples collected, most likely due to regular deworming and improved hygiene in the area. No invasive methods were used for diagnosis of adult worms. From a paper published by E. Kwenti et al. (2016) the prevalence of hookworm was 7% in south west region Cameroon.

6. The number of eggs per gram of stool were estimated for the parasites listed. Did the authors look at the responses in children with high or low intensity of the parasites?

Reply: In this study, after obtaining the prevalence of parasites and comparing with antibody response, no significant difference was observed between the malaria antibodies levels and parasites eggs counts.

7. Table 2 is not necessary, it can be written as text.

Reply: Thanks for the comment, but we think Table 2 summarizes the data more clearly and allows readers to easily compare results from different groups than presenting them in the text. Table 2 has been revised.

8. Page 21, reference # 54, year of publication is missing.
Reply: Year of publication has been included.

Please check spelling and typographical errors scattered through the manuscript (page and lines are given from word document):
1. Page 2, line 3, change led to lead in the sentence.
Reply: The word “led” has been changed to “lead”.

2. Page 2, line 14, correct the spelling of Rietchi concentration method
Reply: Spelling has been corrected to “Ritchie”

3. Page 6, line 21: The bracket has to be closed here: (AB Leo Diagnostics, Helsingborg, Sweden.
Reply: The bracket has been closed.

4. Page 7, line 17 and 18: Correct 50ul to 50µl
Reply: The change has been made.

5. Page 9 and 10: In the text, the p value for anemia (MAL+,IP-) is p=0.034; p value for the same in Table 1 is p=0.032; it needs to be corrected.
Reply: P value has been corrected to P=0.032 (correct value) in the text.

6. Page 10: In Table 1, % sign is missing in column 5 for children with Hb.
Reply: The % symbol has been included in table 1, column 5.

7. Page 10, line 3: In the sentence, change major to majority.
Reply: The word “major” has been changed to “majority”.

8. Page 14, line 27: In the sentence, MSL- should be MAL-
Reply: In Line 27 of page 14, MSL- has been changed to MAL-

9. Page 17, line 15: change beats to beads
Reply: The spelling of beads has been corrected.

10. Re-write the following sentences, they are not very clear:

Page 4, line 8: However, with most children getting infected with several episodes of infections in a short period, this renders them more prone to having clinical symptoms since the immune systems doesn’t fully recover.
Reply: The sentence has been deleted because the information is not directly relevant to the study.

Page 4, line 20: Concomitant infections in humans have suggested that Ascaris lumbricoides infection may protect against cerebral malaria (11,12), while other studies, children infected by S. mansoni were more susceptible to P. falciparum infection and develop acute malaria episodes.
Reply: The sentence has been revised to read: “Studies on concomitant infections in humans suggest that A. lumbricoides infection may protect against cerebral malaria (11,12), while other studies suggest that children infected by S. mansoni may be more susceptible to P. falciparum infections and develop acute malaria episodes (13,14).”

Page 15, line 3: In essence, the immune response in individuals who are repeatedly infection would be similar to that produce during chronic infections.
Reply: To clarify the statement, the text has been revised to read: “Because of high transmission, the children are becoming infected almost daily and are either in the process of eliminating the new infection or reducing it to a submicroscopic level. Because of constant re-exposure, the resulting immune response will be similar to that produced by a chronic infection.

Reviewer #2: The answer to the questions is divided into Major comments, Minor
comments. Additionally, I wrote minor observations that, I hope, will help this manuscript to improve readability and consistency.

1. Is the manuscript technically sound, and do the data support the conclusions?
2. Has the statistical analysis been performed appropriately and rigorously?
3. Have the authors made all data underlying the findings in their manuscript fully available?
4. Is the manuscript presented in an intelligible fashion and written in standard English?

Major comments:

• Given that there were no differences in the IgG response between age groups, it would be interesting to join these data, evaluate all the coinfected individuals, and then split the data into Giardia, E. hystolitica.
Reply: We are confused by this comment, because Fig 1 shows an increase in both Ab prevalence (Fig. 1A) and Ab levels (Fig 1 B-E) with age in Ab-positive children (Kruskal-Wallis test p values were p<0.001 MSP2 and p=0.05-0.086 (borderline) for the other antigens).

We believe combining all MAL+,IP+ children into single a group is unwise, since they were infected with a conglomerate of intestinal helminths, cestodes and protozoa (see Table 2). Combining children with such heterogenous infections is unlikely to provide meaningful information.

• I strongly suggest dividing the age of individuals in 0-5, 5-10, 10-15 years-old to partially solve the "N" problem of the groups.
Reply: Thanks for the comment. Initially, children were groups into 5-year categories as suggested by the Reviewer, i.e., 0-5, 5-10, 10-15 years old. However, when the data set showed that children aged 1 to 2 did not have intestinal parasites, the results were grouped into 2-year intervals, that allowed us to more closely define the increase in Ab prevalence (Fig. 1A) and Ab levels (Fig 1 -B,C,D,E) with age. The purpose of Fig 1 was to determine if age was a variable that needed to be taken into consideration during data analysis.

• Because of the absence of molecular Diagnosis and considering that the authors mention the possibility of oh having low parasitemia infections in the MAL- group. It is important to include MAL- individuals in Figure 1.
Reply: We are sorry if we didn’t make the point clear. ALL children who were Ab-positive are included in Fig 1, including those who are MAL+ and MAL-. Because malaria transmission is high in the area, all children in the study had been exposed to P. falciparum and many of the MAL- children were Ab-positive.

• It is necessary to compare parasite data with similar regions in Cameroon. Please compare and cite:
  • (Malaria and Helminth Co-Infection in Children Living in a Malaria Endemic Setting of Mount Cameroon and Predictors of Anemia from Theresa K Nkuo-Akenji et al. 2006)
  • Malaria, Helminths, Coinfection and Anaemia in a Cohort of Children From Mutengene, South Western Cameroon from Clarisse Njua-Yafi et al. 2016.
Reply: We thank the Reviewer for pointing out the omission of key references. Information from these studies have been included in the revised Discussion. The text now reads, “…..to those found in other highly [malaria] endemic regions of the country (32), and the prevalence of co-infections was 19.1%, which is similar to the prevalence of co-infections of 18 – 27% reported in other regions of Cameroon (9,44). The references have been added to the reference section.

• Do the authors have information about malaria and intestinal parasites last treatments? On page 17, it was commented that Albendazole treatment was frequent in these children. Deworming information will help the readers to understand why the prevalence of intestinal parasites was low compared with other studies in Cameroon. Additionally, reinforce in the discussion section that collecting/reporting that information is valuable for coinfection studies.
Reply: In response to the Reviewer’s suggestion, the following information has been added to the Methods section. “Currently, mass drug administration with albendazole is being performed twice a year by the Ministry of Health, that is usually conducted in
schools and symptomatic cases are sent to the local clinic or hospital for follow up treatment.”

- (Figure 1 B, C, D, E) use the same scale limits for all plots. This is also useful to understand differences in levels of antigenicity between proteins.
  
  Reply: We understand the comment, but we do not wish to change the Y-axis on Fig 1, since it is risky to make a direct comparison of Ab levels between antigens in serological assays. A number of variables, including parasite strain, the system to produce recombinant proteins, protein purity, the amount of antigen used, number of exposed epitopes, dilution of plasma, etc., influence the overall results. Even when Luminex beads are covalently-coupled with saturating amounts of antigen, it is questionable if direct comparison of MFI can be made between antigens. Although our assays have been optimized and equivalence amounts of antigen used during bead-coupling, comparisons among the antigens may not provide accurate information about immunogenicity. In Figs1 B, C, D, E, the Y-Axis was selected to show the best distribution of the MFI results.

- (table 3) How could the authors explain increased eosinophilia with low levels of helminth infection? This mainly applies to the age group > 9 years-old.
  
  Reply: After age 2, children start becoming infected with helminths, resulting in an increase in eosinophil counts. During the biannual drug treatment campaign, helminthic infections are eliminated, but eosinophilia persists for a period of time. With increasing age, more children in the area become i) infected and ii) re-infected, resulting in an increase in prevalence of eosinophilia.

- (Page 17) The authors argue, “First, children living in moist or wet environments where mosquitoes breed and E. histolytica are more abundant would have a high risk of acquiring both infections, that would result in frequent boosting of the Ab response.” This explanation for intestinal parasite influence on antibody production alteration is not viable since Giardia’s frequency is higher than E. histolytica in the studied population.
  
  Reply: The sentence has been deleted from the Discussion.

- (Page 17) The affirmation “Secondly, since malaria and E. histolytica are both amoebae, they might share common antigens, for example, EBA-175 could share homology with an E. histolytica antigen.” is false. Plasmodium falciparum is not an is a protozoan. This group belongs to Apicomplexa organisms. For that reason, the hypothesis about correlating Plasmodium and E. histolytic is wrong.
  
  Sorry, “amoebae” was a typo. Both Plasmodium falciparum and E. histolytica are protozoans. The Discussion has been revised to read “parasitic protozoa.”

- How different are the two Villages Ngali II and Mfou in the central region of Cameroon? Does it exist a difference in humidity and soil moist, once the authors claimed that this variable could explain differences of Entamoeba histolytica?
  
  Reply: The two villages are very similar with no major differences in humidity or soil moisture. The estimated annual average rainfall measures 1600 mm3 with an annual average temperature of 23°C for Ngali II and for Mfou. According to the National Meteorology agency, the average humidity for the center regions is 83%. Ngali and Mfou are both in the center region of Cameroon about 60km apart. Note: as mentioned above, the words “humidity and soil moisture” have been deleted from the MS.

Minor comments:
- What criteria were used to divide the population into seven groups according to age?
  
  Reply: The fact that Intestinal parasite (IP) infections was only observed in children >2 years, helped guide separation of the children into seven groups.

- Please specify how anthropometric parameters were used in the study, once they were described but not used in the study. If this information was not used, please remove these sentences.
  
  Reply: The sentence has been removed.

- Has the studied region presence of Schistosoma haematobium? If the authors have register if this parasite in the area, Did they examined urine samples to discard infections with this parasite?
  
  Reply: Detection of S. haematobium was not included in the study design because of
low prevalence in the study area. A study conducted in this area (and other regions of Cameroon) by Louis-Albert Tchuem Tchuenté et al., (2012) reported a prevalence of S. haematobium of only 1.72%. Since a large sample size would be required to assess the impact of this pathogen on the Ab response to malaria, S. haematobium was not included in the study.

• Were the individuals asymptomatic to intestinal parasites infection too? No diarrhea, abdominal pain, etc.? Please clarify.

Reply: Yes. To make the point clear, the Methods section has been revised and states that all children with clinical cases of malaria or intestinal parasites were not included in the study and referred to the local clinic/hospital by the attending physician for treatment. Thank you for the comment.

• (Page 6) It was mentioned that Plasmodium parasitemia was quantified. Did the authors observe any correlation between the Plasmodium parasite burden and the levels of IgG responses to the antigens?

Reply: As expected, there was no correlation between parasitemia and malaria antibody levels.

• (End of Page 7) Please specify: If the cut-off is MFI+3*SD, how the standard deviation was calculated if the negative controls were pooled? Was this experiment repeated or used replicates? Traditionally, the negative controls are tested simultaneously in different wells of the plate, and the cut-off is calculated from those values.

Reply: Pooled negative control plasma sample were run in triplicates on the same plates as the test samples in all experiments, as well as the positive controls. The cut-off was obtained by calculating MFI+3 SD of the triplicates on all plates in the experiment.

• Did the authors analyze the effect of helminth parasite burden (number of eggs/gram of stool) in those individuals with helminths? This valuable information was commented on but never included in the analysis. If not used, I do not see the necessity of describing in the methods section

Reply: The information has been deleted from the Methods section.

• For data analysis:
  • Before using ANOVA, did the authors checked for the normality of the variables? If yes, please specify, if not, calculate the normality of the variables and the other ANOVA assumptions.

Reply: Yes, ANOVA was used to compare difference in age across the 4 groups (Table 2). However, comparisons of Ab MFI, which are not normally distributed, with age (Fig. 1) were performed using the Kruskal-Wallis test. The Methods section (Data analysis) has been revised. Information in Fig. 1 legend was correct.

• If the authors have not-normal variables, they should use the Kruskal-Wallis non-parametric, and Dunn posthoc tests to verify differences between groups.

Reply: Sorry for the mistake in the Methods section. The Kruskal-Wallis nonparametric test was performed in Fig 1 and 2. A posthoc test was not performed, as the goal was not to determine when peak Ab levels were obtained, but to determine if age had an influence on Ab levels. Since age was a variable, data for all age groups could not be combined, but rather age was taken into consideration during data analysis.

• Please check frequencies described in table 1 (MAL+IP- 58.8%) vs. the values reported in the second line page 9. (59.4%).

Reply: 59.4% is the correct value. The text has been revised.

• Sum of 58.8%+16.9% = 75.7% not 75.6%.

Reply: Thank you for catching the error. The values in Table 1 and text have been revised and are now consistent.

• In table 1, please add a column with P-values to facilitate the interpretation of the differences between groups. Please report statistics of multiple comparisons between groups too.

Reply: The comparisons requested by the reviewer were originally provided in the
Table legend. To comply with the request, the p values have been moved to a column labeled “p values” and the method of analysis was retained in the Table legend.

- What is the potential hypothesis to explain the increased values of parasitemia in the coinfected group?
  Reply: There is no significant difference in parasitemia between the two groups (p=0.1599). In fact, the higher parasitemia was found in young children who were intestinal parasite-negative (probably because very young children were in this group).

- Please comment in the text the presence of multi-parasitism in the studied individuals.
  Reply: We thank the reviewer for the comment. The following sentence has been added to the Results section. “Interestingly, all of the children had single parasite infections, and polyparasitism was not found.”

- (Page 11 table 3). Please include values of anemia and eosinophilia in individuals coinfected. In the current configuration is constructed is hard to determine the coinfected impact in anemia and eosinophilia values.
  Reply: Table 3 was designed to evaluate the influence of age on malaria, IP, anemia and eosinophilia. The number of co-infections are too small to be divided by age. In an attempt to address the Reviewer’s comment, a separate Table was designed that compares the influence of no infections, malaria-positive only, and co-infections on percent with anemia and eosinophilia. The Table will be uploaded as supplemental Table 1. It essentially showed that same results as expected, anemia was associated with malaria and eosinophils were associated with co-infections.

- (Page 11). In the sentence, “Thus, as children living in these villages increased with age, they developed partial immunity to malaria and anemia declined; whereas, the prevalence of IP and eosinophilia increased.” In this sentence, it is necessary to specify that “protection” is protection against malaria symptoms. The table clearly shows that the frequency of malaria does not decrease with age, only the anemia.
  Reply: The sentence has been revised to read: “Thus, as children living in these villages increased with age, they began developing partial immunity to malaria symptoms and anemia declined; whereas, the prevalence of IP and eosinophilia increased.”

- Please plot Age vs. Antibody levels for each protein to verify the correlation for each protein studied.
  Reply: The figure on the right confirms that Ab levels increase with age. The figure shows a linear regression analysis of Ab levels for MSP1, MSP2, MSP3 and EBA-175 using data from all 320 children, and includes the equation for the regression line, the R2 value (all positive), and p value (all significant). Thus, the figure confirms that Ab levels increase with age. We do NOT wish to include this figure in the MS since it is essentially identical to the one shown in Fig 1 B, C, D and E. In fact, we feel that the information in Fig 1B-E is easier for the reader to understand.
  Note: If the figure is not shown, it is provided in a separate document.

- As an exploratory analysis, I suggest joining all data and make a boxplot comparing MFI between MAL-PI-, MAL-PI+, MAL+PI-, and MAL+PI+. Mainly for MSP1, MPS2, and MSP3 group age 3-10 and 11-15 to check.
  Reply: We thank the Reviewer Thanks for the suggestion concerning exploratory analysis. A comparison of Ab levels in two of the above groups (MAL-,IP-, and MAL+,IP-) is shown in Fig 2. Unfortunately, the number of children in the MAL-,PI+ group is too small to provide valuable information. As stated above, children in the MAL-,PI+ group (n=54) are infected with a variety of intestinal helminths, cestodes and protozoa (see Table 2). With such a diverse range of pathogens, plotting the data as a boxplot will not provide useful information. In Fig. 2, the distribution of Ab levels in children co-infected with malaria and single intestinal pathogens is provided. We feel this approach is more informative than “dumping all pathogens together.”

- The sentence “E. histolytica is a gut amoeba that causes both intestinal and extraintestinal infections such as amebic colitis (dysentery) and liver or brain abscess. The protozoa cause a marked down-regulation of macrophage functions rendering the cells incapable of antigen presentation and unresponsive to cytokine stimulation (57)”
does not explain the increase of antibody production in E. histolytica infected group. Why could a diminishing antigen presentation generate higher levels of anti-Plasmodium antigens?  
Reply: Very true! Not sure why that statement wasn’t caught. The Discussion has been changed significantly. It now reads, “The decrease in macrophage function does not explain the increase in Ab to EBA-175. One possible explanation is that since malaria and E. histolytica…”

Other observations/questions:

• In the title, add "IgG" to Antibody response. Reply: IgG has been added to title (although not all of the co-authors agree this is necessary).

• Check all scientific names of parasite species for correct formatting in italics. (Example Entamoeba histolytica in the Results section in the abstract)  
Reply: The scientific name has been checked and are now in italics.

• Please, mention in the background the region where the study was performed. Reply: This information was included in the background section of the Abstract. It is also included in the Materials section.

• It is necessary to describe and discuss the role of MSP1, MPS2, MSP3, and EBA-175 as markers in serological studies.  
Reply: This information has been added to the Discussion.

• Considering that coinfection prevalence is relatively low, I consider that it is important to discriminate with colors or point shapes the individuals MAL-IP-, MAL+IP-, MAL-IP+, MAL+IP+ in Figure 1 B-C-D-E  
Reply: We thank the Reviewer for the suggestion. However, information in Fig 1B-E is designed to address the question, are Ab prevalence and levels influence by age? Whereas, Fig 2 provides comparisons between individuals infected with malaria alone or co-infected with specific intestinal parasites. Thus, colored dots or symbols are not needed in Fig 1 (and could be confusing to the reader).

• In page 6 subtitle "Laboratory detection, quantification and speciation of malaria parasites.", I will not use speciation here. I suggest “Diagnosis and quantification of Plasmodium sp. parasites.”  
Reply: The header has been changed to read: “Laboratory detection of malaria parasites.”

• (Page 14-15) What type of parasite is "Amoeba"? What is the difference between "Amoeba" and E. histolytica? Traditionally, E. histolytica is considered an amoeba too.  
Reply: The figure has been revised to read Intestinal Protozoa. Thanks for pointing out the mis-classification.

• In table 1, to facilitate reading, please remove symbols % and /ul located in cells with data and add to the columns describing the variables.  
Reply: The symbols in the data cells have been removed.

• For consistency, unify parasitemia vs. parasitemia, anemia vs. anaemia in the text and plots.  
Reply: The British spelling of parasitaemia, anaemia, and haemoglobin have been used through out the MS.

• (Page 10) change "The major of helminth parasites" to "The most frequent helminth species detected."  
Reply: The change was made as suggested.

• (Table 2) Check all the total numbers for the "Total IP+" column. For example, for protozoans, the sum is 29+19+4 = 48, and it was reported 47  
Reply: This has been verified and corrected to 48 in Table 2

• (Page 13) In plot titles Change Ab (Antibody) to IgG  
Reply: We thank the Reviewer for the comment, but decide not to make the change. Our rationale is that by definition, IgG is a class of immunoglobulin found in the blood;
whereas, Ab are plasma proteins that bind specifically with an antigen. What was measured was IgG Ab. Since the serological assay measured IgG Ab that were recorded as MFI (median fluorescence intensity), we think the labels on the Y-Axis (Ab levels -MFI) reflect what was done. The Methods section makes it clear that the Ab were of the IgG class. [Note: Serum IgG levels (which implies mg/ml) were not measured.]

• (Figure 1E) Add, Change from EBA to EBA-175.
Reply: Change has been made.

• Please verify all references formatting (For example, reference 42 is all in capital letters)
Reply: References have been edited as requested by the reviewer.

Review #3: Comments were in the attachment.
Reply: In revising the MS, all requested changes were made and additional information provided in the text, including information on the BLAST search. The only request we would not fully address is the prevalence of bednet use in the villages. The only information available is that very few children use bednets. Since the slide-positivity rate of 75.6% for P. falciparum, it is unlikely the bednets are having a major influence on the current study. The following information has been added to the MS in the Results section. “To determine if higher Ab levels in children co-infected with P. falciparum and E. histolytica might be due to cross-reactive epitopes, a BLAST search for sequence homology between EBA-175 and E. histolytica proteins was made. No similarities were found using Metablast, and only one hit was found using discontinuous metablast which had a span of only 38 nucleotides (~12 amino acids). Thus, there does not appear to be shared epitopes between these two pathogens that would explain the increase in Ab to EBA-175 in children with co-infections.”

Figure for Reviewer #2 confirming an increase in antibody levels with age.

Additional Information:

| Question                | Response                                                                 |
|-------------------------|--------------------------------------------------------------------------|
| Financial Disclosure    | The author(s) received no specific funding for this work. Funding used in for this research was mentors (Prof Leke Rose) and a Gift of the magnetic beats from Dr Anna Babakhanyan. No other specific funding were received. |
Unfunded studies
Enter: The author(s) received no specific funding for this work.

Funded studies
Enter a statement with the following details:
• Initials of the authors who received each award
• Grant numbers awarded to each author
• The full name of each funder
• URL of each funder website
• Did the sponsors or funders play any role in the study design, data collection and analysis, decision to publish, or preparation of the manuscript?
• NO - Include this sentence at the end of your statement: The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.
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* typeset

Competing Interests
Use the instructions below to enter a competing interest statement for this submission. On behalf of all authors, disclose any competing interests that could be perceived to bias this work—acknowledging all financial support and any other relevant financial or non-financial competing interests.

This statement will appear in the published article if the submission is accepted. Please make sure it is accurate. View published research articles from PLOS ONE for specific examples.

The authors have declared that no competing interests exist.
NO authors have competing interests

Enter: The authors have declared that no competing interests exist.

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I have read the journal's policy and the authors of this manuscript have the following competing interests: [insert competing interests here]

Ethics Statement

Enter an ethics statement for this submission. This statement is required if the study involved:

- Human participants
- Human specimens or tissue
- Vertebrate animals or cephalopods
- Vertebrate embryos or tissues
- Field research

Write "N/A" if the submission does not require an ethics statement.

General guidance is provided below. Consult the submission guidelines for detailed instructions. Make sure that all information entered here is included in the Methods section of the manuscript.

Ethical clearance used for the study was obtained from the Cameroon National Ethics Committee (IRB approval: No2016/12/845/CE/CNERSH/SP). Administrative authorizations were obtained from authorities of the Ngali II and Mfou health districts. Written Informed consents were obtained from parents of all participants.
Format for specific study types

**Human Subject Research (involving human participants and/or tissue)**
- Give the name of the institutional review board or ethics committee that approved the study
- Include the approval number and/or a statement indicating approval of this research
- Indicate the form of consent obtained (written/oral) or the reason that consent was not obtained (e.g. the data were analyzed anonymously)

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Full-title: The immunoglobulin G antibody response to malaria merozoite antigens in asymptomatic children co-infected with malaria and intestinal parasites

Running title: Malaria antibodies in children with intestinal parasites

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ABSTRACT

Background

Co-infection with malaria and intestinal parasites is common in children in Africa and may affect their immune response to a malaria parasite infection. Prior studies suggest that co-infections may lead to increased susceptibility to malaria infection and disease severity; however, other studies have shown the reverse. Knowledge on how co-morbidities specifically affect the immune response to malaria antigens is limited. Therefore, this study sought to determine the prevalence of co-infection of malaria and intestinal parasites and its association with antibody levels to malaria merozoite antigens.

Methods

A cross sectional study was carried out in two villages with high transmission of malaria in Cameroon (Ngali II and Mfou). Children aged 1-15 years were enrolled after obtaining parental consent. A malaria rapid diagnostic test was used on site. Four (4) ml of peripheral blood was collected from each participant to determine Plasmodium falciparum infections by microscopy, haemoglobin levels and serology. Fresh stool samples were collected and examined by wet mount, Kato-Katz method and modified Ritchie concentration techniques. A Multiplex Analyte Platform assay was used to measure antibody levels.

Results

A total of 320 children were enrolled. The prevalence of malaria by blood smear was 76.3% (244/320) and prevalence of malaria and intestinal parasites was 16.9% (54/320). Malaria prevalence was highest in young children; whereas, intestinal parasites (IP+) were not present until after 3 years of age. All children positive for malaria had antibodies to MSP1, MSP2, MSP3 and EBA175. No difference in antibody levels in children with malaria-co infections compared to malaria alone were found, except for antibody levels to EBA-175.
were higher in children co-infected with intestinal protozoa \((p = 0.018)\), especially those with *Entamoeba histolytica* infections \((p=0.0026)\).

**Conclusion**

Antibody levels to EBA175 were significantly higher in children co-infected with malaria and *E. histolytica* compared to children infected with malaria alone. It is important to further investigate why and how the presence of these protozoans can modulate the immune response to malaria antigens.

**Key words:** malaria, intestinal-parasites, antibody, *Giardia lamblia, Entamoeba histolytica*

**Introduction:**

In sub-Saharan Africa, malaria caused by *Plasmodium falciparum* (Pf) remains an important public health threat, killing over 271,000 children under the age of five each year \((1)\). In malaria endemic areas, individuals exposed to malaria infections gradually develop clinical immunity \((2)\) and commonly experience asymptomatic infections without fever or symptoms and do not require antimalarial treatment. Asymptomatic infection results from partial immunity that controls, but does not completely eliminate, malaria parasites, thus allowing for constant presence of circulating parasites \((2)\).

The prevalence of intestinal parasitic infections in children is fairly constant across sub-Saharan Africa with an average prevalence of 26\% \((3,4)\). In Cameroon, the prevalence in children less than 18 years is 26.8\% \((5)\), while that for the general population is more than 28\% The major intestinal parasites are *Ascaris lumbricoides, Trichuris trichuria and Entamoeba histolytica* \((6-8)\), but many cases of intestinal parasites go undetected.
Co-infections with malaria and intestinal parasites (IP) are common in malaria endemic areas in sub-Saharan Africa (7,8) and infections with IP and Pf are both ranked among the major cause of mortality and morbidity in sub-Saharan Africa. Several studies conducted on IP (not including amoebas) and Pf have shown conflicting results. Some helminths suppress different T-helper types and favor an increase in regulatory T (Treg) cell (9). Studies on concomitant infections in humans suggest that *A. lumbricoides* infection may protect against cerebral malaria (10,11), while other studies suggest that children infected by *Schistosoma mansoni* may be more susceptible to *P. falciparum* infections and develop acute malaria episodes (12,13). Also, it has been shown that the levels of TNF-α, IL-2, IL-10, IL-6 in *Plasmodium*-helminth co-infected individuals were significantly higher than the malaria-positive (MP) group (14) dampening the immune response to malaria. However, little known regarding host immune responses to malaria in children co-infected with protozoan pathogens. Studies suggest that children co-infected with malaria and intestinal helminths had significantly decreased antibody levels to the malarial antigen apical merozoite antigen 1 (AMA-1) compared to those with *P. falciparum* or IP alone(15). Hence, infections with intestinal helminths can stifle protective anti-plasmodial antibody responses (15). However, increase in MSP3 IgG1–4 levels were significantly associated with children infected with malaria alone compared to children co-infected with both parasites(15).

Malaria and other intestinal parasites overlap extensively in their epidemiological distributions causing polyparasitism. Polyparasitism with intestinal parasites has been reported as one of the contributing factors to hypo-responsiveness (16), dampening of the immune response by inducing a strong Treg response, which could in turn, blunt a strong response to vaccines (17). Equally, some studies have suggested an effect of IP on antibody responses to *P. falciparum* gametocyte antigens that may have consequences on transmission-blocking immunity (18).
Effective elimination and future eradication of malaria will require not only vector control, but also managing asymptomatic malaria patients and developing an effective vaccine. Given the high burden and concomitant nature of both malaria and intestinal parasites in the same geographical setting, conflicting data shows polyparasitism could interfere with the efficacy of malaria vaccines (19). To our knowledge, since limited information is available on whether and how co-infections of intestinal parasites and malaria affect the specific immune response to malaria antigens (20), the goal of this study was to investigate the prevalence and relationship between co-infections of malaria (MAL+) and intestinal parasites (IP+) (nematodes, trematodes, and protozoans) on naturally acquired antibodies to malaria.

**Methods**

**Study area description**

The study was conducted in Ngali II and Mfou, two villages in the central region of Cameroon (located at 4°27′N and 11°38′E) with a total population of about 1,000 children per squared Km (about 4000 in Ngali II and 6000 in Mfou) under the age of 15 years. The climate is typically equatorial with two discontinuous dry and rainy seasons. The annual average rainfall measures about 1600 mm with an annual average temperature of 23°C (21).

Most children in Ngali II and Mfou over 3 years of age accompany their parents to the farm and return home late at night. The use of mosquito bed nets is rare in the two villages and residents have minimal access to portable water with approximately one well per 500 inhabitants. Currently, mass drug administration with albendazole is being performed twice a year by the Ministry of Health, that is usually conducted in schools and symptomatic cases are sent to the local clinic or hospital for follow up treatment.
Study population

A cross sectional study was carried out in Ngali II and Mfou from January to May 2017, a transitional period from the dry to wet season. Children who had lived in either of the villages for at least six months and whose parents gave informed consent were included in the study. All participants were systematically examined by a physician for clinical systems of malaria and IP. Children who presented with symptoms of malaria, e.g., fever, headaches or intestinal illnesses, e.g., diarrhea, vomiting were not enrolled. A total of 320 participants (140 from Ngali II and 180 from Mfou) aged 1-15 years participated in the study. Since both villages have the same demographic features, data for the two villages were combined.

Blood collection and on-site testing for malaria

Venous peripheral blood (about 4mL) was collected by venipuncture using a butterfly needle (G22) and a 5mL labeled EDTA tube from all 320 participants. Haemoglobin (Hb) was measured using the HemoCue (AB Leo Diagnostics, Helsingborg, Sweden). On site, after collecting the venous blood from the participants, a drop from the same collected blood was placed on a CareStart™ Malaria pLDH/HRP-2 Combo Test (Access Bio Inc. USA) to detect histidine-rich protein-2 (HRP-2) specific to *Plasmodium falciparum* and *Plasmodium lactate dehydrogenase* (pLDH) pan-specific to *Plasmodium spp* (*falciparum, P. vivax, P. malariae, P. ovale*). Results were read according to manufacturer instructions and recorded after 5 minutes.

Laboratory detection of malaria parasites

Ten microliters of whole blood were used to prepare thick and thin smears for malaria parasite identification, speciation and quantification. The slides were air-dried overnight, and the thin blood smears were fixed in absolute (100%) methanol. Both thick and thin smears were stained using 10% Giemsa solution, washed with water and air-dried. Slides were then microscopically examined (thin and thick smear) for the presence of malaria parasites by two
experienced microscopists. The parasite density was determined by counting the number of parasites against 200 leucocytes. The counts were expressed as the number of P. falciparum-infected erythrocytes (IE) per microliter of blood (Pf IE/µl), assuming an average leukocyte count of 8,000 cells/µl of blood (22). When the difference in parasitaemia between the two readers was greater than 5 Pf IE/µl of blood, a third reader re-examined the slide and the mean of the two closest values were considered. Also, a differential count for eosinophil, lymphocytes, monocytes, neutrophils was obtained alongside parasitaemia and different malaria species (by microscopy)

**Antibody Analysis**

Plasma samples were tested for antibodies against the merozoite antigens MSP-142, MSP-2, MSP-3 and EBA-175 using a multi-analyte platform assay with antigen-coupled magnetic beads with different spectral addresses. Details of this assay used has been described previously (23) (24). In brief, plasma samples were diluted 1:100 with PBS, 50µl of plasma was incubated with 50µl antigen-coupled microspheres (2000 microspheres/test) for 60 minutes in the dark, washed with PBS, and then incubated at 500rpm for 60minutes at 25 °C on a rotating shaker and using a magnet plate separator. Then, 100 µl of secondary Ab (R-phycoerythrin-conjugated, Affini Pure F(ab')2 fragment, Goat anti-human IgG Fc fragment specific, Jackson Immuno-research, West Grove, PA, USA, Cat no. 109-116-170) diluted to 2 µg/ml in PBS-1 % BSA was added to each well and incubated as above in the dark for 1 h. The mixture is then washed and a minimum of 100 beads were read in a MAGPIX® reader. A minimum bead count of 100 per spectral address recorded as Median Fluorescence Intensity (MFI).

Controls included on each plate were: PBS to determine background fluorescence, the negative control (NC) consisted of pooled plasma from four malaria-naïve US individuals, and the positive control (PC) was pooled plasma from Cameroonian with high antibody levels to *Plasmodium falciparum*. Results were exported to Excel for analysis. The cut-off for
positivity was calculated as mean of MFI +3 standard deviation of the negative control as shown in the results sections.

Stool sample collection and analysis

Sterile labelled stool collection vials were given to the parents along with instructions for proper stool collection. All samples were analyzed within 7 hours of collection to avoid missing hookworm eggs and minimize chances of under reporting. Approximately, 4 mg of feces was suspended in 5ml PBS and a drop examine by wet mount. The Kato Katz technique was used for morphological identification of helminths eggs, e.g., A. lumbricoides, T. trichiura, or larval stage of Strongyloides stercoralis (25) while the modified Ritchie’s concentration stool technique was used to identify all protozoans and cestodes (26). The smears were read at objective 10X for eggs and larvae and objective 40X for cysts and vegetative forms of protozoan. All stool slides were read by 2 technicians and in 2 different laboratories under supervision of a microbiologist and parasitologists.

Data analysis

Data were analyzed using Microsoft Excel 2013, and GraphPad® prism 8. Standard summary statistics were used to describe the study population and results are presented as proportions. Fischer’s exact test was used to compare antibody levels between the malaria-negative, IP-positive (MAL-,IP+) and malaria-positive, IP-negative (MAL+,IP-) groups, because of the small sample sizes of the groups. The one-way-ANOVA test was used to compare all 4 groups after checking for normality (e.g., age). An unpaired t test was used to compare the means of the MAL-,IP- vs. MAL+,IP- groups. Kruskal-Wallis test was used to compare antibody levels, which are not normally distributed, among the groups or within the MAL+IP+ groups. An individual was considered to have a co-infection if at least one IP species and P. falciparum were present. Anaemia was considered when Hb values were < 11.5 g/dL and classified according to WHO (27,28). To search DNA sequences of P.
falciparum EBA-175 and those of *E. histolytica* for possible cross-reactive epitopes, PIEBA175 ([ncbi.nlm.nih.gov/gene/2654998](https://ncbi.nlm.nih.gov/gene/2654998)) was compared with *E. histolytica* ([ncbi.nlm.nih.gov/assembly/GCF_000208925.1](https://ncbi.nlm.nih.gov/assembly/GCF_000208925.1)) using Megablast for highly similar sequences and discontinuous megablast for more dissimilar sequences.

**Results**

**The study population**

A total of 320 children were enrolled (Table 1). Among the children, 76.3% were slide-positive for malaria (MAL+), with 59.4% having malaria without intestinal parasites (MAL+,IP⁻) and 16.9% being coinfected with malaria and intestinal parasites (MAL+, IP+). All subjects who tested positive for malaria using the rapid diagnostic field test were confirmed positive by microscopy. Among children who were infected with malaria, 71.3% were infected with only *P. falciparum* and 5% had *P. falciparum* and *P. malariae*. Interestingly, only 2.2% of the children had IP without malaria and 21.6% were negative for both malaria and IP.

The mean age of the children changed with infection status among the 4 groups (p = 0.0001) with the lowest age found in uninfected children (6.4 years) and highest in children with co-infections (9.3 years) (Table 1). Malaria infections were found in all age groups; whereas, none of the children under age 4 years had intestinal parasites. Mean haemoglobin levels were lower in children infected with malaria, but the difference was of marginal significance (p = 0.08; MAL⁻,IP⁻ vs MAL+,IP+) the prevalence of anaemia was higher in children who were infected with malaria (MAL+,IP⁻)(p=0.032), but not those with co-infections (p >0.999) compared to children who were parasite-negative (MAL⁻,IP⁻).

| Table 1: Description of 320 children infected with malaria and intestinal parasites (IP) |
|-----------------------------------------------|---------------------------------|---------------------------------|---------------------------------|---------------------------------|
| MAL⁻, IP⁻ | MAL⁺, IP⁻ | MAL⁻,IP⁺ | Co-infections (Mal⁻,IP⁺) | P values |
| Malaria and intestinal parasites | Malaria only | Malaria and *P. malariae* | Malaria and *P. falciparum* | Malaria and *E. histolytica* |
|-----------------------------------------------|---------------------------------|---------------------------------|---------------------------------|---------------------------------|
| Mean age (years) | 6.4 | 7.1 | 8.2 | 9.3 |
| Prevalence of anaemia | 45% | 50% | 55% | 58% |
| Mean haemoglobin levels (g/dL) | 11.5 | 10.8 | 10.3 | 9.8 |


Prevalence of intestinal parasites

Overall, 19.1% (61/320) of the children were positive for intestinal parasites, 16.9% of whom were also infected with malaria and 2.2% were IP+ but MAL- (Table 2). The most frequent helminthic parasites detected were A. lumbricoides (2.8%) and single cases of Trichura sp. and Strongyloides sp. Among the 320 children, 14.7% had detectable protozoan infections, including 7.8% infected with Giardia lamblia, 5.9% with E. histolytica, and 0.9% with Isospora sp. Very few children had intestinal cestodes (Table 2). Interestingly, all of the children had single parasite infections, and polyparasitism was not found.
Table 2: Prevalence of Intestinal Parasites (IP+) in the 320 Children, Ages 1 to 15 years

| Intestinal Parasites | Number of Children | Total IP+ (% positive) |
|----------------------|--------------------|------------------------|
| **Mal**              | MAL-, IP+ | MAL+, IP+ | Total IP+ |
| **Helminths**        | 11 (3.4%)    |            |           |
| *Ascaris lumbricoides*| 2             | 7          | 9 (2.8%)  |
| Others*              | 0             | 2          | 2 (0.87%) |
| **Protozoans**       | 48 (14.7%)    |            |           |
| *Giradia lamblia*    | 3             | 22         | 25 (7.8%) |
| *Entamoeba histolytica complex* | 1 | 18 | 19 (5.9%) |
| Other**              | 1             | 3          | 4 (0.9%)  |
| **Cestodes**         | 2 (0.63%)     |            |           |
| *Hymenolepis nana*   | 0             | 2          | 2 (0.63%) |
| **Total IP**         | 7 (2.2%)      | 54 (16.9%) | 61 (19.1%) |

Others*: 1 Trichura sp. and 1 Strongyloides sp.
Others**: 3 Isospora sp.

Influence of age on malaria, intestinal parasites, anaemia and moderate eosinophilia

As expected, children aged 1 through 2 years did not have soil-transmitted IP and had normal eosinophil levels; whereas, 63% of 1-2-year old children were infected with malaria and had the highest prevalence of anaemia (Table 3). In contrast, in children 9-15 years of
age ~80% were slide-positive for malarial parasites, 24%-29% had intestinal parasites, and 10-38% had moderate eosinophilia. Thus, as children living in these villages increased with age, they began developing partial immunity to malaria symptoms and anaemia declined; whereas, the prevalence of IP and eosinophilia increased.

Table 3: Influence of Age on Malaria, Intestinal Parasites, Anaemia and Percentage of Peripheral Eosinophils

| Age (years) | N  | % Mal+ | % IP+ | % with anaemia | % with eosinophilia |
|------------|----|--------|-------|----------------|------------------|
| 1 - 2      | 27 | 63.0   | 0     | 55.6           | 0                |
| 3 - 4      | 47 | 61.7   | 6.4   | 48.9           | 4.3              |
| 5-6        | 40 | 62.5   | 15.0  | 40.0           | 7.5              |
| 7-8        | 63 | 88.9   | 28.6  | 36.5           | 9.5              |
| 9-10       | 55 | 83.6   | 21.8  | 38.2           | 20.0             |
| 11-12      | 54 | 79.6   | 25.9  | 38.9           | 22.2             |
| 13-15      | 34 | 82.4   | 23.5  | 26.5           | 38.3             |

*Anemia: Children with haemoglobin less than 11.5 g/dL. **Moderate eosinophilia: ≥1,500 eosinophils/mm³ or ≥18.7% peripheral eosinophils

A comparison of anaemia and eosinophilia among the 4 groups of children shown in Table 1 was made (S1 Table). Results showed that anaemia was associated with malaria infections and eosinophilia was associated with IP.

Antibody Levels to Malaria Merozoite Antigens

With repeated exposure to malaria, Ab prevalence and levels increased with age to the four merozoite antigens (Fig 1). Among 1- to 2-year-olds, only 25% of the infants had Ab to EBA-175 and MSP3, 30% had Ab to MSP2, but 80% had Ab to MSP1 (Fig 1). However, by age 13-15 years, 60% had acquired Ab to MSP3 and >80% had Ab EBA-175, MSP2 and MSP3 (Fig 1A). Among Ab-positive children, Ab levels also increased with age (Fig 1B-E).
Although different amounts of Ab were ultimately obtained to the different antigens, the overall trend was for an increase in median Ab with age. Thus, it was important to take age into consideration when making comparison between children infected with malaria (MAL+,IP-), co-infected with malaria and IP (MAL+,IP+) and those who were not infected (MAL-,IP-) at the time the study was conducted.

[Insert Figure 1]

**Fig1: Prevalence and amount of Ab in different age groups.** (A) Prevalence of Ab to the 4 merozoite antigens. The number of participants in each age group is provided in Table 3. Fig1 B – E show Ab levels (MFI) for children who were Ab-positive for each age group. Horizontal bars represent median Ab levels. Kruskal-Wallis test (nonparametric comparison among groups) values were for MSP1 (p=0.067); MSP2 (p<0.001); MSP3 (p=0.086) and EBA (p=0.056). MFI = Median fluorescence intensity; MSP = merozoite surface proteins; EBA= erythrocytes binding antigen

**Comparison of Ab levels in participants with and without malaria and IP**

Since children below 3 years of age were not infected with IP, they were not included in the comparative studies described below. Given that Ab prevalence and levels increased with age, the study population was divided into 2 groups: children aged 3-10 years, a time period when children were becoming infected with IP (Table 3) and those 11-15 years, mainly children who had been infected repeatedly with malaria and may had lived with IP for a period of time. As predicted, Ab levels were slightly higher in MAL+ children due to current boosting compared to MAL-, but the differences were not significant (all p values >0.05) (Fig 2).

A comparison between Ab levels in children infected with malaria and co-infected with IP was conducted. Children with helminths and cestodes were not included in the
analysis because the sample sizes were too small. Ab levels were compared between
children aged 3-10 years infected with malaria (n=112) and co-infected with flagellate and
intestinal amoeba (n= 25 children), including G. lamblia (n= 15) and E. histolytica (n = 10
children) (Fig 2). Antibody levels did not differ between malaria-infected children with or
without intestinal amoeba for MSP1, MSP2 and MSP3; however, there were significantly
higher Ab levels to EBA-175 in children co-infected with malaria and intestinal amoeba (p =
0.018) (Fig 2D). The higher Ab levels were due to E. histolytica infections (p=0.0026), and
not G. lamblia (p=0.3844). No differences were found between children aged 11 to 15 years
for any of the antigens between children with malaria (single infection) and co-infected with
any of the IP.

To determine if higher Ab levels in children co-infected with P. falciparum and E.
histolytica might be due to cross-reactive epitopes, a BLAST search for sequence homology
between EBA-175 and E. histolytica proteins. No similarities were found using Metablast,
and only one hit was found using discontinuous metablast which had a span of only 38
nucleotides (~13 amino acids) that had 82% similarity. Thus, there does not appear to be
shared epitopes between these two pathogens that would explain the increase in Ab to EBA-
175 in children with co-infections.

[Insert Figure 2]

Fig 2. Antibody levels in children ages 3 to 10 for all antibody-positive individuals
Distribution of Ab levels in MFI among malaria negative (MAL-) and malaria-positive (MAL+)
and those co-infected with malaria plus Intestinal (Int.) protozoa (n=25); malaria plus G.
lamblia (n=15); and malaria plus E. histolytica (n=10). The number of datapoints varied
because not all participants had Ab to all antigens. Horizontal lines represent medians for
the group. MFI = median fluorescence intensity; MSP = merozoite surface proteins, EBA =
erythrocytes binding antigen (EBA)
Discussion

Malaria and polyparasitism (cestodes, protozoans, trematodes) are still common conditions throughout Africa (29,30). In the 1-15-year-old children living in the rural Cameroonian villages surveyed, the prevalence of slide-positive malaria was 76.3% and 19.1% had intestinal parasites, with 16.9% co-infections (Table 1-2). This prevalence of malaria is similar to those found in other highly endemic regions of the country (31), and the prevalence of co-infections was 19.1%, which is similar to a prevalence of 18 – 27% reported in other regions of Cameroon (32,33). This high transmission is related to geo-ecological and climatic conditions at the time of the study which was the transition from the dry to wet season, a period that favors vector breeding and distribution (34).

From Table 3, the prevalence of slide-positive malaria ranged from 61% to 90% in different age groups implying that children in these villages were repeatedly exposed to malaria throughout their lives. The current prevalence of malaria in 2017 is similar to that recorded previously for Ngali II between 1998-2004, that ranged from 50% to 85% in 5-15 year old children with an estimated entomological inoculation rate of 0.7 infectious bites/per/night (~257 infectious bites annually)(35). Prior studies have established that repeated exposure induces immunity to malaria, with development of anti-disease immunity followed by anti-parasite immunity (36–39). As a result, the highest prevalence of 56% anaemia was found in young children (2,40,41) with a decline to 27% in 13 to 15-year-olds (Table 3). On the other hand, Infections with IP only occurred later in life from 3 years onward with a mean infection age of 8.1 years. Increase in intestinal parasites was associated with an age-related increase in eosinophil counts (42,43), a known innate immune response to helminthic and other soil-transmitted organisms (Table 3). In this study, only 11/320 (3.4%) children were infected with helminths. Although some epidemiological studies have demonstrated an increased risk of infection by P. falciparum in individuals co-infected with helminths, other results are conflicting (44,45). The low prevalence of helminths is explained, in part by, the
fact that mass community de-worming is done biannually following the national infectious
disease guide-line for IP control program. The most prevalent intestinal parasites were the
protozoans, *G. lamblia* and *E. histolytica* (48). These protozoa are commonly found in damp
soil and contaminated water with a prevalence of 2-20% in Cameroon (50–53). These
results suggest children acquire their intestinal infections after learning to walk and interact
with the environment. Thus, children in the study population were exposed to malaria early
in life and began developing anti-malaria immunity prior to exposure to intestinal parasites.

Generally, both Ab prevalence and Ab levels increased with age in 1 to 15-year-olds living in
this high transmission area (Fig 1). Often, the presence of Ab is used as markers of
infection, including the merozoite antigens used in this study. This study compared antibody
levels with age in four main groups of children, MAL+, IP+, MAL-, IP-, MAL-, IP+ and MAL+, IP-
children to four (MSP1, MSP2 MSP3, EBA17) malaria antigens (54–56). Over 80% of
1-2-year-olds had Ab to MSP1, humoral immunity began to develop early in life and
continued to mature as children developed into adolescents (Fig 1). Often individuals who
are MAL+ have higher Ab levels than MAL- individuals due to boosting of the Ab response
(36,38,39). In the current study, Ab levels did not differ significantly between MAL+ and
MAL- individuals, neither those who were 3-10 years nor 11-15 years-old. This result was
not surprising, since 75% of the children were slide-positive for malaria (Table 1). Because
of high transmission, children are becoming infected almost on a daily basis and either are in
the process of eliminating the new infection or reducing it to submicroscopic levels. Thus,
most children living in areas with high perennial transmission will test positive for malaria by
PCR. *Because of constant re-exposure, the resulting immune response will be similar to that
produced by a chronic infection.*

Prior studies have demonstrated that malaria-helminths co-infections can down regulate
malaria and orient the immune response via the Th2 response hence, making patients less
sick (20,36,57,58) whereas, others have demonstrated on the contrary that IP and malaria
co-infections increase malaria disease (13,56). Unfortunately, the current study could not
resolve the controversy because very few children had helminthic infections, due to frequent
treatment with albendazole via the mass drug administration program conducted by the
Ministry of Health and other random health campaigns. However, co-infections with malaria
and amoeba were relatively common. Ab levels to MSP1, MSP2 and MSP3 were similar in
children infected with *P. falciparum* alone and those with amoeba (Fig 2); however,
significantly higher Ab levels to EBA-175 were found in children co-infected with malaria and
intestinal amoeba (p = 0.018). The higher Ab levels were due to *E. histolytica* infections (p =
0.0026), and not *G. lamblia* (p = 0.384). This result was unexpected. *E. histolytica* is a gut
amoeba that cause both intestinal and extraintestinal infections such as amebic colitis
(dysentery) and liver or brain abscess. This protozoa can cause a marked down‐regulation
of macrophage functions rendering the cells incapable of antigen presentation and
unresponsive to cytokine stimulation (59). This decrease in macrophage function does not
explain the increase in Ab to EBA-175. One possible explanation is that since malaria and
*E. histolytica* are both protozoan pathogens, they might share common antigens, for
example, EBA-175 could share homology with an *E. histolytica* antigen. To investigate this
possibility, a blast search of the NCIB gene bank was conducted for EBA-175 and the *E.
histolytica* genome. However, this search revealed only a ~13 amino acid sequence with
82% similarity, which is clear too small to explain the increase in Ab levels of co-infected
children. Finally, an alternative explanation could be that this result is a spurious
observation by chance. Clearly the association between malaria and *E. histolytica* merits
further study.

 Altogether, a keen observation needs to be repeated with a larger sample size as *E.
histolytica* boosting of Ab to EBA-175 – co-infection might not only be limited to EBA-175,
but other antigens as well. Children in these villages began to acquire an Ab response to the
4 merozoite antigens early in life, prior to infection with IP. There was no evidence that
infection with IP influenced Ab levels or negatively-altered the already established Ab
response to the 4 merozoite antigens.
CONCLUSION

The prevalence of malaria was high in children 1-2 years old; whereas, intestinal parasite infections occurred in children over 3 years old. Thus, immunity to *P. falciparum* began prior to infection with soil-transmitted parasites. No differences were found in antibody prevalence or levels in malaria-infected and co-infected children, except antibody levels to EBA175 were significantly higher in children co-infected with malaria and *E. histolytica*. This is the first report of an interaction between malaria and *E. histolytica* and antibodies to EBA-175 and merits further evaluation.

Declarations

Ethical consideration

Ethical clearance used for the study was obtained from the Cameroon National Ethics Committee (IRB approval: N°2016/12/845/CE/CNERSH/SP). Administrative authorizations were obtained from authorities of the Ngali II and Mfou health districts.

Informed consents were obtained from parents of all participants. A clinical examination was performed for all eligible participants by a medical doctor.

All participants positive for any *Plasmodium* spp by RDT at the time of blood collection and those who were found to have PI by stool analysis were treated for free following the protocol recommended by the Cameroonian Ministry of Health. All children with mild anaemia were given an iron supplement free of charge.

Authors’ contributions

GFLR supervised the study. JDB and WM co-supervised the study. CMN and ELG conceived and designed the work. CMN, EFL, DJC, AEW, MBN carried out experiments.
Data was collected and analyzed by CMN. The first draft of this manuscript was written by CMN, critically read and edited by DWT, YML. DWT and GFLR reviewed the final draft.

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Competing interest

Authors declare no competing interests.

Data Availability:

The database for the study can be found in the “Supporting Material File.”

Consent for publication

All authors give their consent for publication of this manuscript.

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Running title: Malaria antibodies in children with intestinal parasites

7/20/20 – this is the original MS with Reviewer #3 tracks included.

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ABSTRACT

Background

Co-infection with malaria and intestinal parasites is common in children in Africa and may affect their immune response to a malaria parasite infection. Prior studies suggest that co-infections may lead to increased susceptibility to malaria infection and disease severity; however, other studies have shown the reverse. Knowledge on how co-morbidities specifically affect the immune response to malaria antigens is limited. Therefore, this study sought to determine the prevalence of co-infection of malaria and intestinal parasites and its association with antibody levels to malaria merozoite antigens.

Methods

A cross sectional study was carried out in two villages with high transmission of malaria in Cameroon (Ngali II and Mfou). Children aged 1-15 years were enrolled after obtaining parental consent. A malaria rapid diagnostic test was used on site. Four (4) ml of peripheral blood was collected from each participant for microscopy to determine Plasmodium falciparum infections by microscopy and species, haemoglobin levels and serology. Fresh stool samples were collected and examined by wet mount, Kato-Katz method and modified Ritchie-Ritchi concentration techniques. A Multiplex Analyte Platform (MAP) assay was used to measure antibody levels.

Results

A total of 320 children were enrolled. The prevalence of malaria by blood smear was 76.3% (244/320 (75.6%)) and prevalence of malaria and intestinal parasites was co-infections 16.9% (54/320) (16.9%). Malaria prevalence was highest in young children; whereas, intestinal parasites (IP+) were not present until after 3 years of age. All children positive for malaria had antibodies to MSP1α, MSP2, MSP3 and EBA175. No difference in antibody levels in children with malaria-co infections compared to malaria alone were found, except for antibody levels to EBA175 were higher in children co-infected with intestinal
protozoa (amoeba) (p = 0.018), especially those with *Entamoeba histolytica* infections
(p=0.0026).

**Conclusion:**

Antibody levels to EBA175 were significantly higher in children co-infected with malaria and
*E. histolytica* compared to children infected with malaria alone. It is important to further
investigate why and how the presence of these protozoans can modulate the immune
response (Th1/Th2)-to malaria antigens.

**Key words:** malaria, intestinal-parasites, antibody, *Giardia lamblia, Entamoeba histolytica*
**Background/Introduction:**

In sub-Saharan Africa, malaria caused by *Plasmodium falciparum* (Pf) remains an important public health threat, killing over 271,000-292,000 children under the age of five each year (1). In malaria endemic areas, individuals exposed to malaria infections gradually develop clinical immunity (2)(3) and commonly experience asymptomatic infections without fever or symptoms and do not require antimalarial treatment. A symptomatic infection results from partial immunity that controls, but does not completely eliminate, malaria parasites, thus allowing for constant presence of circulating parasites (2)(3). However, with most children getting infected with several episodes of infections in a short period, this renders them more prone to having clinical symptoms since the immune systems doesn’t fully recover.

The prevalence of intestinal parasitic infections in children is fairly constant across sub-Saharan Africa with an average prevalence of 26% (3,4)(4,5). In Cameroon, the prevalence in children less than 18 years is 26.8% (5)(6), while that for the general population is more than 28%. The major soil-transmitted intestinal parasites are *Ascaris lumbricoides*, *Trichuris trichuria* and *Entamoeba histolytica* (6–8)(7–9), but many cases of intestinal parasites go undetected.

Co-infections with malaria and intestinal parasites (IP) are common in malaria endemic areas in sub-Saharan Africa (7,8)(8,9) and infections with IP and Pf are both ranked among the major cause of mortality and morbidity in sub-Saharan Africa. Several studies conducted on IP (not including amoebas) and Pf have shown conflicting results. Some helminths suppress different T-helper types and favor an increase in regulatory T (Treg) cell (9)(10). Studies on concomitant infections in humans have suggested suggest that *A. scaris lumbricoides* infection may protect against cerebral malaria (10,11)(11,12), while other studies suggest that children infected by *Schistosoma mansoni* may be more susceptible to *P. falciparum* infections and develop acute malaria episodes (12,13)(13,14). Also, it has been shown that the levels of TNF-α, IL-2, IL-10, IL-6 in *Plasmodium-helminth* co-infected individuals were significantly higher than the malaria-positive (MP) group (14)(15).
Studies suggest that children co-infected with malaria and intestinal helminths had significantly decreased antibody levels to the malarial antigen apical merozoite antigen 1 (AMA-1) compared to those with *P. falciparum* or IP alone [15][46]. Hence, infections with intestinal helminths can stifle protective anti-plasmodial antibody responses [15][46]. However, increase in MSP3 IgG1–4 levels were significantly associated with children infected with malaria alone compared to children co-infected with both parasites [15][46].

Malaria and other intestinal parasites overlap extensively in their epidemiological distributions causing polyparasitism. Polyparasitism with intestinal parasites has been reported as one of the contributing factors to hypo-responsiveness [16][47], dampening of the immune response by inducing a strong Treg response, which could in turn, blunt a strong response to vaccines [17][48]. Equally, some studies have suggested an effect of IP on antibody responses to *P. falciparum* gametocyte antigens that may have consequences on transmission-blocking immunity [18][49].

Effective elimination and future eradication of malaria will require not only vector control, but also managing asymptomatic malaria patients and developing an effective vaccine. Given the high burden and concomitant nature of both malaria and intestinal parasites in the same geographical setting, conflicting data shows polyparasitism could interfere with the efficacy of malaria vaccines [19][20]. To our knowledge, since limited information is available on whether and how co-infections of intestinal parasites and malaria affect the specific immune response to malaria antigens [20][44], the goal of this study was to investigate the prevalence and relationship between co-infections of malaria (MAL+) and intestinal parasites (IP+) (nematodes, trematodes, and protozoans) on naturally acquired antibodies to malaria merozoite.

**Methods.**

**Study Area description:**

The study was conducted in Ngali II and Mfou, two villages, in the central region of Cameroon (located at 4°27′N and 11°38′E) with a total population of about 1,000 children per squared Km (about 4000 in Ngali II and 6000 in Mfou) under the age of 15 years. The climate
is typically equatorial with two discontinuous dry and rainy seasons. The annual average rainfall measures about 1600 mm with an annual average temperature of 23°C. Most children in Ngali II and Mfou over 3 years of age accompany their parents to the farm and return home late at night. They seldom sleep under The use of mosquito bed nets is rare in the two villages and are geographically similar, residents are relatively poor and have minimal access to portable water; with approximately one well per 500 inhabitants. Currently, mass drug administration with albendazole is being performed twice a year by the Ministry of Health, that is usually conducted in schools and symptomatic cases are sent to the local clinic or hospital for follow up treatment.

**Study Population**

A cross sectional study was carried out in Ngali II and Mfou from January to May 2017, a transitional period from the dry to wet season. Children who had lived in either of these two villages for at least six months and whose parents gave informed consent were included in the study. Since both villages (Ngali II & Mfou) were very similar in all features, data for both village were combined. Vital parameters (temperature, pulse) and anthropometric parameters (weight, height) were measured by assisting attendant nurses. These parameters were used to calculate body mass index (BMI) and advice was given to the parents of the participating children, as part of a service for participation. All participants were systematically examined by a physician for clinical systems of malaria and IP. Only asymptomatic participants were included in the study. Children who presented with symptoms of malaria, e.g., fever, headaches or intestinal illnesses, e.g., diarrhea, vomiting were not enrolled. A total of 320 participants (140 from Ngali II and 180 from Mfou) aged 1-15 years participated in the study. Since both villages have the same demographic features, data for the two villages were combined.

**Blood Collection and On-site Testing for Malaria**

Venous peripheral blood (about 4mL) was collected by venipuncture using a butterfly needle (G22) and a 5mL labeled EDTA tube from all 320 participants. Hæmoglobin (Hb) was measured using the HemoCue (AB Leo Diagnostics, Helsingborg, Sweden). On site, after
collecting the venous blood from the participants, a drop from the same collected blood was placed on a CareStart™ Malaria pLDH/HRP-2 Combo Test (Access Bio Inc. USA) to detect histidine-rich protein-2 (HRP-2) specific to *Plasmodium falciparum* and *Plasmodium lactate dehydrogenase* (pLDH) pan-specific to *Plasmodium spp* (*falciparum, P. vivax, P. malariae, P. ovale*). Results were read according to manufacturer instructions and recorded after 5 minutes.

**Laboratory detection, quantification and speciation of malaria parasites.**

Ten microliters of whole blood were used to prepare thick and thin smears for malaria parasite identification, speciation and quantification. The slides were air-dried overnight, and the thin blood smears were fixed in absolute (100%) methanol. Both thick and thin smears were stained using 10% Giemsa solution, washed with water and air-dried. Slides were then microscopically examined (thick and thick smear) for the presence of malaria parasites by two experienced microscopists. The parasite density was determined by counting the number of parasites against 200 leucocytes. The counts were expressed as the number of *P. falciparum*-infected erythrocytes (IE) parasites per micro-liter of blood (*Pf IE*/µl), assuming an average leukocyte count of 8,000 cells/µl of blood (22)(23). When the difference in parasitaemia between the two readers was greater than 5 *Pf IE*/µl of blood, a third reader re-examined the slide and the mean of the two closest values were considered. Also, a differential count for eosinophil, lymphocytes, monocytes, neutrophils was obtained alongside parasitaemia and different malaria species (by microscopy)

**Antibody Analysis**

Plasma samples were tested for antibodies against the merozoite antigens MSP-1₄₅, MSP-2, MSP-3 and EBA-175 using a multi-analyte platform (MAP) assay with antigen-coupled magnetic beads with different spectral addresses. Details of this assay used has been described previously (23)(24)(24)(25). In brief, plasma samples were diluted 1:100 with PBS, 50µl of plasma was incubated with 50µl antigen-coupled microspheres (2000 microspheres/test) for 60 minutes in the dark, washed with PBS, and then incubated at
500rpm for 60 minutes at 25 °C on a rotating shaker and using a magnet plate separator.

Then, 100 µl of secondary Ab (R-phycoerythrin-conjugated, Affini Pure F(ab’): fragment, Goat anti-human IgG Fc fragment specific, Jackson Immuno-research, West Grove, PA, USA, Cat no. 109-116-170) diluted to 2 µg/ml in PBS-1 % BSA was added to each well and incubated as above in the dark for 1 h. The mixture is then washed and a minimum of 100 beads were read in a MAGPIX® reader. A minimum bead count of 100 per spectral address recorded as Median Fluorescence Intensity (MFI).

Controls included on each plate were: PBS to determine background fluorescence, the negative control (NC) consisted of pooled plasma from four malaria-naïve US individuals, and the positive control (PC) was pooled plasma from Cameroonian with high antibody levels to *Plasmodium* falciparum. Results were exported to Excel for analysis. The cut-off for positivity was calculated as mean of MFI +3 standard deviation of the negative control as shown in the results sections.

**Stool sample collection and analysis**

Sterile labelled stool collection vials were given to the parents along with instructions for proper stool collection. All samples were analyzed within 7 hours of collection to avoid missing hookworm eggs and minimize chances of under reporting. Approximately, 4 mg of feces was suspended in 5ml PBS and a drop examine by wet mount. The Kato Katz technique was used for morphological identification of helminths eggs, e.g., *A. scaris*, *lumbricoides, T. trichura trichiura*, or larval stage of *Strongyloides stercoralis* [25][26] while the modified Ritchie’s concentration stool technique was used to identify all protozoans and cestodes [26][27]. The smears were read at objective 10X for eggs and larvae and objective 40X for cysts and vegetative forms of protozoan. The number of eggs per gram of stool were estimated for the parasites listed. Helminth eggs and protozoans were counted in about 4mg of stool and counts were extrapolated as the number of eggs per gram of stool. All stool slides were read by 2 technicians and in 2 different laboratories under supervision of a microbiologist and parasitologists.

**Data analysis**


Data were analyzed using Microsoft Excel 2013, and GraphPad Prism 8. Standard summary statistics were used to describe the study population and results are presented as proportions. Fischer’s exact test was used to compare antibody levels between the malaria-negative, IP-positive (MAL-, IP+) and malaria-positive, IP-negative (MAL+, IP-) groups, because of the small sample sizes of the groups. The one-way ANOVA test was used to compare all 4 groups after checking for normality (e.g., age). An unpaired t test was used to compare the means of the MAL-, IP- vs. MAL+, IP- groups. Kruskal-Wallis test was used to compare antibody levels, which are not normally distributed, among the groups or within the MAL+IP+ groups. An individual was considered to have a co-infection if at least one IP species and P. falciparum were present. Anemia was considered when Hb values were < 11.5 g/dL and classified according to WHO [27, 28]. To search DNA sequences of P. falciparum EBA-175 and those of E. histolytica for possible cross-reactive epitopes, PFEBA175 (ncbi.nlm.nih.gov/gene/2654998) was compared with E. histolytica (ncbi.nlm.nih.gov/assembly/GCF_000208925.1) using Megablast for highly similar sequences and discontinuous megablast for more dissimilar sequences.

**Results**

**The Study Population**

A total of 320 children were enrolled (Table 1). Among the children, 76.3% were slide-positive for malaria (MAL+), with 59.4% having malaria without intestinal parasites (MAL-, IP-) and 16.9% being coinfected with malaria and intestinal parasites (MAL+, IP+). All subjects who tested positive for malaria using the rapid diagnostic field test were confirmed positive by microscopy. Among children who were infected with malaria, 71.3% were infected with only P. falciparum and 5% had P. falciparum and P. malariae. Interestingly, only 2.2% of the children had IP without malaria and 21.6% were negative for both malaria and IP.

The mean age of the children changed with infection status among the 4 groups (p = 0.0001) with the lowest age found in uninfected children (6.4 years) and highest in children with co-infections (9.3 years) (Table 1). Malaria infections were found in all age groups;
whereas, none of the children under age 4 years had intestinal parasites. Mean haemoglobin levels were lower in children infected with malaria, but the difference was of marginal significance ($p = 0.08; \text{MAL}-,\text{IP}- \text{ vs } \text{MAL+},\text{IP}-$). The prevalence of anaemia was higher in children who were infected with malaria (MAL+,IP-) ($p=0.03$), but not those with co-infections ($p >0.999$) compared to children who were parasite-negative (MAL-,IP-).

Table 1: Description of 320 children infected with malaria and intestinal parasites (IP)

|                  | MAL- IP- | MAL+ IP- | MAL+ IP+ Co-infections (Mal+,IP+) | Total P values |
|------------------|----------|----------|---------------------------------|----------------|
| Number (%) of children | 69 (21.6%) | 190 (59.4%) | 7 (2.2%) | 54 (16.9%) | 320 |
| Mean years of age (range) | 6.4 (1-14) | 7.9 (1-15) | 8.6 (4-12) | 9.3 (4-15) | 0.0001* |
| Parasitaemia: (median # infected erythrocytes/µl (range) | 0 (40-96,000) | 420/µl (40-96,000) | 0 | 900/µl (40-30,970) | 0.1599** |
| Measures of anaemia | | | | | |
| Hb (g/dL) (mean ±SD) | 12.1 ±1.6 | 11.6 ± 2.2** | 12.2 ± 1.4 | 12.4±1.8 | 0.0658* |
| Prevalence of anaemia | | | | | |
| # (%) of children with Hb <11.5 g/dL | 21 (30.4%) | 87 (45.6%)** | 2 (28.6%) | 17 (31.5) | 131 |

*p = 0.0001*, comparison among the 4 groups (ordinary one-way ANOVA)

** comparison among the 4 groups (Mann-Whitney test)

***p = 0.087, comparison between MAL- IP - vs. MAL+ IP- (unpaired t test)

***p = 0.032, comparison between MAL-,IP - vs. MAL+,IP- (Fisher’s exact test)
Overall, 19.1% (61/320) of the children were positive for intestinal parasites, 16.9% of whom were also infected with malaria and 2.2% were IP+ but MAL- (Table 2). The most frequent major of helminthic parasites detected were *Ascaris lumbricoides* (2.8%) and single cases of *Trichura sp.* and *Strongyloides sp.* Among the 320 children, 14.7% had detectable protozoan infections, including 7.8% infected with *Giardia lamblia*, 5.9% with *Entamoeba histolytica*, and 0.9% with *Isospora sp.* Very few children had intestinal cestodes (Table 2).

Interestingly, all of the children had single parasite infections, and polyparasitism was not found.

**Table 2: Prevalence of Intestinal Parasites (IP+) in the 320 Children, Ages 1 to 15 years**

| Intestinal Parasites | Number of Children |
|----------------------|--------------------|
| **Helminths**        |                    |
| *Ascaris lumbricoides* | 2 (7) 9 (2.8%)     |
| Others*              | 0 (2) 2 (0.87%)    |
| **Protozoans**       | 48 (14.7%)         |
| *Giardia lamblia*    | 3 (22) 25 (7.8%)   |
| *Entamoeba histolytica complex* | 1 (18) 19 (5.9%) |
| Other**              | 1 (3) 4 (0.9%)     |
| **Cestodes**         | 2 (0.63%)          |
| *Hymenolepis nana*   | 0 (2) 2 (0.63%)    |
| **Total IP**         | 7 (2.2%) 54 (16.9%) 61 (19.1%) |

Others*: 1 *Trichura sp.* and 1 *Strongyloides sp.*
Others**: 3 *Isospora sp.*
Influence of Age on Malaria, Intestinal Parasites, Anaemia and Moderate Eosinophilia

As expected, children aged 1 through 2 years did not have soil-transmitted IP and had normal eosinophil levels; whereas, 63% of 1-2-year olds children were infected with malaria and had the highest prevalence of anaemia (Table 3). In contrast, in children 9-15 years of age ~80% were slide-positive for malarial parasites, 24%-29% had intestinal parasites, and 10-38% had moderate eosinophilia. Thus, as children living in these villages increased with age, they began developing partial immunity to malaria symptoms and anaemia declined; whereas, the prevalence of IP and eosinophilia increased.

Table 3: Influence of Age on Malaria, Intestinal Parasites, Anaemia and Percentage of Peripheral Eosinophils

| Age (years) | N = | % Mal+ | % IP+ | *% with anaemia | **% with eosinophilia |
|------------|-----|--------|-------|-----------------|----------------------|
| 1-2        | 27  | 63.0   | 0     | 55.6            | 0                    |
| 3-4        | 47  | 61.7   | 6.4   | 48.9            | 4.3                  |
| 5-6        | 40  | 62.5   | 15.0  | 40.0            | 7.5                  |
| 7-8        | 63  | 88.9   | 28.6  | 36.5            | 9.5                  |
| 9-10       | 55  | 83.6   | 21.8  | 38.2            | 20.0                 |
| 11-12      | 54  | 79.6   | 25.9  | 38.9            | 22.2                 |
| 13-15      | 34  | 82.4   | 23.5  | 26.5            | 38.3                 |

*Anaemia: Children with haemoglobin less than 11.5 g/dL. **Moderate eosinophilia: ≥1,500 eosinophils/mm³ or ≥18.7% peripheral eosinophils

A comparison of anaemia and eosinophilia among the 4 groups of children shown in Table 1 was made (S1 Table). Results showed that anaemia was associated with malaria infections and eosinophilia was associated with IP.

Antibody Levels to Malaria Merozoite Antigens
With repeated exposure to malaria, Ab prevalence and levels increased with age to the four merozoite antigens (Fig. 1). Among 1- to 2-year-olds, only 25% of the infants had Ab to EBA-175 and MSP3, 30% had Ab to MSP2, but 80% had Ab to MSP1 (Figure 1). However, by age 13-15 years, 60% had acquired Ab to MSP3 and >80% had Ab EBA-175, MSP2 and MSP3 (Fig. 1A). Among Ab-positive children, Ab levels also increased with age (Fig. 1B-E). Although different amounts of Ab were ultimately obtained to the different antigens, the overall trend was for an increase in median Ab with age. Thus, it was important to take age into consideration when making comparison between children infected with malaria (MAL+,IP-), co-infected with malaria and IP (MAL+,IP+) and those who were not infected (MAL-,IP-) at the time the study was conducted. 

[Figure 1 – revised]
Figure 1: Prevalence and amount of Ab in different age groups. (A) Prevalence of Ab to the 4 merozoite antigens. The number of participants in each age group is provided in Table 4. Figure 1B–E shows Ab levels (MFI) for children who were Ab-positive for each age group. Horizontal bars represent median Ab levels. Kruskal-Wallis test (nonparametric comparison among groups) values were for MSP1 (p=0.067); MSP2 (p<0.001); MSP3 (p=0.086) and
Comparison of Ab Levels in Participants with and without Malaria and IP

Since children below 3 years of age were not infected with IP, they were not included in the comparative studies described below. Given that Ab prevalence and levels increased with age, the study population was divided into 2 groups: children aged 3-10 years, a time period when children were becoming infected with IP (Table 3) and those 11-15 years, mainly children who had been infected repeatedly with malaria and may have lived with IP for a period of time. As predicted, Ab levels were slightly higher in MAL+ children due to current boosting compared to MAL-, but the differences were not significant (all p values >0.05) (Figure 2).

A comparison between Ab levels in children infected with malaria and co-infected with IP was conducted. Children with helminths and cestodes were not included in the analysis because the sample sizes were too small. Ab levels were compared between children aged 3-10 years infected with malaria (n=112) and co-infected with flagellate and intestinal amoeba (n= 25 children), including *G. lamblia* (n=15) and *E. histolytica* (n=10 children) (Figure 2). Antibody levels did not differ between malaria-infected children with or without intestinal amoeba for MSP1, MSP2 and MSP3; however, there were significantly higher Ab levels to EBA-175 in children co-infected with malaria and intestinal amoeba (p = 0.018) (Figure 2D). The higher Ab levels were due to *E. histolytica* infections (p=0.0026), and not *G. lamblia* (p=0.3844). No differences were found between children aged 11 to 15 years for any of the antigens between children with malaria (single infection) and co-infected with any of the IP.

To determine if higher Ab levels in children co-infected with *P. falciparum* and *E. histolytica* might be due to cross-reactive epitopes, a BLAST search for sequence homology between EBA-175 and *E. histolytica* proteins. No similarities were found using Metablast, and only one hit was found using discontinuous metablast which had a span of only 38 nucleotides (~13 amino acids) that had 82% similarity. Thus, there does not appear to be
shared epitopes between these two pathogens that would explain the increase in Ab to EBA-175 in children with co-infections.

[FIGURE 2 - revised]

**Fig 2.** Antibody levels in children ages 3 to 10 for all antibody-positive individuals

Distribution of Ab levels in MFI among malaria negative (MAL-) and malaria-positive (MAL+) and those co-infected with malaria plus *Intestinal (Int.) amoebaprototiza* (n=25); malaria plus *G. lamblia* (n=15); and malaria plus *E. histolytica* (n=10). The number of datapoints varied because not all participants had Ab to all antigens. Horizontal lines represent medians for the group. MFI = median florescence intensity; MSP = merozoite surface proteins, EBA = erythrocytes binding antigen (EBA)

**Discussion**
Malaria and polyparasitism (cestodes, protozoans, trematodes) are still common conditions throughout Africa (29,30)(36,34). In the 1-15-year-old children living in the two rural Cameroonian villages surveyed, the prevalence of slide-positive malaria was 76.35.6% and 19.1% had intestinal parasites, with 16.9% co-infections (Table 1-2). This prevalence of malaria is similar to those found in other highly endemic regions of the country (31)(32), and reported in other regions of Cameroon (32,33)(3,4). This high transmission is related to the geo-ecological and climatic conditions at the time of the study which was the transition from the dry to wet season, a period that favors vector breeding and distribution (34)(33).

From Table 3 above, the prevalence of slide-positive malaria ranged from 61% to 90% in different age groups implying that children in these villages were repeatedly exposed to malaria throughout their lives. The current prevalence of malaria in 2017 is similar to that recorded previously for Ngali II between 1998-2004, that ranged from 50% to 85% in 5-15 year olds, with an estimated entomological inoculation rate of 0.7 infectious bites/per/night (~257 infectious bites annually) (35).

Prior studies have established that repeated exposure induces immunity to malaria, with development of anti-disease immunity followed by anti-parasite immunity (36-39)(44-37). As a result, the highest prevalence of 56% anaemia was found in young children (2,40,41)(3,38,39) with a decline to 27% in 13 to 15-year-olds (Table 3). On the other hand, Infections with IP only occurred later in life from 3 years onward with a mean infection age of 8.1 years. Increase in intestinal parasites was associated with an age-related increase in eosinophil counts (42,43)(40,44), a known innate immune response to helminthic and other soil-transmitted organisms (Table 3). In this study, only 11/320 (3.4%) children were infected with helminths. Although some epidemiological studies have demonstrated an increased risk of infection by P. falciparum in individuals co-infected with helminths, other results are conflicting (44,45)(42,43). The low prevalence of helminths This could be is explained, in part by, the fact that mass community de-worming is done biannually following the national infectious disease guide-line for IP control program.

The most prevalent intestinal parasites were the protozoans, G.tardia lamblia intestinalis and E. histolytica (48). These protozoa are commonly found in damp soil and contaminated water with a prevalence of 2-20% in Cameroon (50–53)(48–51). These results
suggest children acquire their intestinal infections after learning to walk and interact with the
environment. Thus, children in the study population were exposed to malaria early in life and
began developing anti-malaria immunity prior to exposure to intestinal parasites."

Generally, both Ab prevalence and Ab levels increased with age in 1 to 15-year-olds living in
this high transmission area (Fig. 1). Often, the presence of Ab is used as markers of
infection, including the merozoite antigens used in this study. This study compared antibody
levels with age in four main groups of children, MAL+,IP+, MAL-,IP-, MAL-,IP- and
MAL+,IP- children to four (MSP1, MSP2 MSP3, EBA17) malaria antigens.(54–56),(52–54).
Since over 80% of 1-2-year-olds had Ab to MSP1, humoral immunity began to develop early
in life and continued to mature as children developed into adolescents (Fig. 1). Often
individuals who are MAL+ have higher Ab levels than MAL- individuals due to boosting of
the Ab response (36,38,39)(34,36,37). In the current study, Ab levels did not differ
significantly between MAL+ and MAL- individuals, neither those who were 3-10 years
nor 11-15 years-old. This result, however, was not surprising, since 75% of the children
were slide-positive for malaria (Table 1). Because of high transmission. Therefore, it is likely
that children are becoming infected almost on a daily basis and either are in the process of
eliminating the new infection or reducing it to either slide-negative or had either been
recently infected or had submicroscopic infection. Thus, most children living in
areas with high perennial transmission will test positive for malaria by PCR. In essence, the
immune response in individuals who are repeatedly infected would be similar to that produced
during chronic infections. Because of constant re-exposure, the resulting immune response
will be similar to that produced by a chronic infection.

Prior studies have demonstrated that malaria-helminths co-infections can down regulate
malaria and orient the immune response via the Th2 response hence, making patients less
sick (20,36,57,58)(21,34,55,56) whereas, others have demonstrated on the contrary that IP
and malaria co-infections increase malaria disease (13,56)(14,54). Unfortunately, the current
study could not resolve the controversy because very few children had helminthic infections,
due to frequent treatment with albendazole via the mass drug administration program
conducted by the Ministry of Health and other random health campaigns. However, co-infections with malaria and amoeba were relatively common. Ab levels to MSP1, MSP2 and MSP3 were similar in children infected with *P. falciparum* alone and those with amoeba (Fig. 2); however, significantly higher Ab levels to EBA-175 were found in children co-infected with malaria and intestinal amoeba (*p* = 0.018). The higher Ab levels were due to *E. histolytica* infections (*p* = 0.0026), and not *G. lamblia* (*p* = 0.384). This result was unexpected. *E. histolytica* is a gut amoeba that cause both intestinal and extraintestinal infections such as amebic colitis (dysentery) and liver or brain abscess. These protozoa can cause a marked down-regulation of macrophage functions rendering the cells incapable of antigen presentation and unresponsive to cytokine stimulation (59,57). This decrease in macrophage function does not explain the increase in Ab to EBA-175. One possible explanation is that in addition to a possible immunological interaction, there are at least 2 other explanations as to why *E. histolytica* infections might be associated with higher Ab levels to EBA-175. First, children living in moist or wet environments where mosquitoes breed and *E. histolytica* are more abundant would have a high risk of acquiring both infections, that would result in frequent boosting of the Ab response. Secondly, since malaria and *E. histolytica* are both protozoan pathogens amoebae, they might share common antigens, for example, EBA-175 could share homology with an *E. histolytica* antigen. To investigate this possibility, a blast search of the NCIB gene bank was conducted for EBA-175 and the *E. histolytica* genome. However, this search revealed only a ~13 amino acid sequence with 82% similarity, which is clear too small to explain the increase in Ab levels of co-infected children. showed no significant similarity between both gene sequences. Finally, an alternative explanation could be that this result is a spurious observation by chance. Clearly the association between malaria and *E. histolytica* merits further study. Altogether, a keen observation needs to be repeated with a larger sample size as *E. histolytica* boosting of Ab to EBA-175 – co-infection might not only be limited to EBA-175, but other antigens as well. Children in these villages began to acquire an Ab response to the 4 merozoite antigens early in life, prior to infection with IP. -There was no evidence that
infection with IP influenced Ab levels or negatively-altered the already established Ab response to the 4 merozoite antigens.

CONCLUSION

The prevalence of malaria was high in children 1-2 years old; whereas, intestinal parasite infections occurred in children over 3 years old. Thus, immunity to *P. falciparum* began prior to infection with soil-transmitted parasites. No differences were found in antibody prevalence or levels in malaria-infected and co-infected children, except antibody levels to EBA175 were significantly higher in children co-infected with malaria and *E. histolytica*. This is the first report of an interaction between malaria and *E. histolytica* and antibodies to EBA-175 and merits further evaluation.

**Declarations**

**Ethical consideration**

Ethical clearance used for the study was obtained from the Cameroon National Ethics Committee (IRB approval: No 2016/12/845/CE/CNERSH/SP). Administrative authorizations were obtained from authorities of the Ngali II and Mfou health districts.

Informed consents were obtained from parents of all participants. A clinical examination was performed for all eligible participants by a medical doctor.

All participants positive for any *Plasmodium* spp by RDT at the time of blood collection and those who were found to have PI by stool analysis were treated for free following the protocol recommended by the Cameroonian Ministry of Health. All children with mild anemia were given an iron supplement free of charge.

**Authors’ contributions**
GFLR supervised the study. JDB and WM co-supervised the study. NCMN and ELG conceived and designed the work. NCMN, EFL, DJC, AEW, MBN carried out experiments. Data was collected and analyzed by NCMN. The first draft of this manuscript was written by NCMN, critically read and edited by DWT, YML. DWT and GFLR reviewed the final draft.

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Competing interests:
Authors declare no competing interests.

Data Availability:
The database for the study can be found in the “Supporting Material File.” The authors approve of the availability of all data underlying the findings and without restriction upon reasonable request from the corresponding authors. All-important data are within the paper.
All authors give their consent for publication of this manuscript.

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**Reviewer #1:** Dr. Mbe-cho and colleagues sought to determine the prevalence of co-infection of malaria and intestinal parasites and its association with antibody levels to malaria merozoite antigens. The authors report that there was no difference in antibody prevalence or levels in malaria-infected and co-infected children, except antibody levels to EBA-175 were significantly higher in children co-infected with malaria and E. histolytica. Overall, the study is well-designed but these results do not significantly alter or impact our understanding of the association of malaria and helminths on antibody to malaria merozoite antigens.

1. The limitation of the study is that the parasite testing in children was not followed by sensitive diagnostic techniques like PCR, and light infections may have been missed which may have resulted in misclassification of the groups. Light infections may boost the antibody responses while children remain asymptomatic.

   **Reply:** We understand the concern. When the study was conducted (2017) in the rural villages, the prevalence of slide-positive malaria was 75.6%. In prior study conducted in the village (Leke et al 2010), an equivalent prevalence was found of P. falciparum (50-85%) in children aged 5-15 years over a 5-year period. The estimated entomological inoculation rate (EIR) was 0.7 infectious bites/person/nightly thought out the year (~257 IB/P/Y). Based on the more recent malaria prevalence, it appears that the current EIR is similar. Thus, children were most likely being bitten approximately every-other night by an infectious mosquito, since bednets were not routinely used. With this high level of transmission, most of the slide-negative children would be PCR-positive for malaria, i.e., have enough immunity to reduce malaria to submicroscopic levels. Unfortunately, in very high transmission areas like the one reported herein, everyone will have some circulating P. falciparum parasites. So, classifying subjects as slide-positive vs slide-negative may not reflect presence/absence of parasites, but provide information on the immune status of the person. In revising the MS, information from the study by Leke et al. was included as well as a discussion of submicroscopic infections in the revised Discussion.

2. In this study, only 3.4% children were infected with helminths alone to get any meaningful data for antibody response to malaria in this group.

   **Reply:** We agree, the sample size of children with helminth infections is too small to provide meaningful information. Accordingly, Ab levels in children with helminth infections were not analyzed. To explain the low prevalence of helminths, information on the Ministry of Health’s policy for biannual treatment of children for worms was provided.

3. Very few children are positive for E. histolytica.

   **Reply:** True, the prevalence of Entamoeba in our study was only 5.9%, which is lower than that reported in studies in these areas of ~23% (T. E. Kwenti et al., 2016). In our study, the prevalence was lower, probably due to rigorous mass drug administration (MDA) programs implemented by the Ministry of Health and other regular or seasonal health campaigns.

4. The data on the children's anthropomorphic measurements are not mentioned. Thus, there is not much point describing how they were collected.

   **Reply:** This section was removed from the Methods section.

5. There is no data on hookworm infection in the results.

   **Reply:** The prevalence of hookworm infections was considered in this study during stool exams and, surprisingly, we did not find hookworms in the samples collected, most likely due to regular deworming and improved hygiene in the area. No invasive methods were used for diagnosis of adult worms. From a paper published by E. Kwenti et al. (2016) the prevalence of hookworm was 7% in south west region Cameroon.
6. The number of eggs per gram of stool were estimated for the parasites listed. Did the authors look at the responses in children with high or low intensity of the parasites?
Reply: In this study, after obtaining the prevalence of parasites and comparing with antibody response, no significant difference was observed between the malaria antibodies levels and parasites eggs counts.

7. Table 2 is not necessary, it can be written as text.
Reply: Thanks for the comment, but we think Table 2 summarizes the data more clearly and allows readers to easily compare results from different groups than presenting them in the text. Table 2 has been revised.

8. Page 21, reference # 54, year of publication is missing.
Reply: Year of publication has been included.

Please check spelling and typographical errors scattered through the manuscript (page and lines are given from word document):
1. Page 2, line 3, change led to lead in the sentence.
Reply: The word “led” has been changed to “lead”.

2. Page 2, line 14, correct the spelling of Rieties concentration method
Reply: Spelling has been corrected to “Ritchie”

3. Page 6, line 21: The bracket has to be closed here: (AB Leo Diagnostics, Helsingborg, Sweden.
Reply: The bracket has been closed.

4. Page 7, line 17 and 18: Correct 50ul to 50µl
Reply: The change has been made.

5. Page 9 and 10: In the text, the p value for anemia (MAL+,IP-) is p=0.034; p value for the same in Table 1 is p=0.032; it needs to be corrected.
Reply: P value has been corrected to P=0.032 (correct value) in the text.

6. Page 10: In Table 1, % sign is missing in column 5 for children with Hb.
Reply: The % symbol has been included in table 1, column 5.

7. Page 10, line 3: In the sentence, change major to majority.
Reply: The word “major” has been changed to “majority”.

8. Page 14, line 27: In the sentence, MSL- should be MAL-
Reply: In Line 27 of page 14, MSL- has been changed to MAL-

9. Page 17, line 15: change beats to beads
Reply: The spelling of beads has been corrected.

10. Re-write the following sentences, they are not very clear:

Page 4, line 8:
However, with most children getting infected with several episodes of infections in a short period, this renders them more prone to having clinical symptoms since the immune systems doesn’t fully recover.
Concomitant infections in humans have suggested that Ascaris lumbricoides infection may protect against cerebral malaria (11,12), while other studies, children infected by S. mansoni were more susceptible to P. falciparum infection and develop acute malaria episodes.

Reply: The sentence has been revised to read: “Studies on concomitant infections in humans suggest that A. lumbricoides infection may protect against cerebral malaria (11,12), while other studies suggest that children infected by S. mansoni may be more susceptible to P. falciparum infections and develop acute malaria episodes (13,14).”

In essence, the immune response in individuals who are repeatedly infection would be similar to that produce during chronic infections.

Reply: To clarify the statement, the text has been revised to read: “Because of high transmission, the children are becoming infected almost daily and are either in the process of eliminating the new infection or reducing it to a submicroscopic level. Because of constant re-exposure, the resulting immune response will be similar to that produced by a chronic infection.

Reviewer #2: The answer to the questions is divided into Major comments, Minor comments. Additionally, I wrote minor observations that, I hope, will help this manuscript to improve readability and consistency.

1. Is the manuscript technically sound, and do the data support the conclusions?
2. Has the statistical analysis been performed appropriately and rigorously?
3. Have the authors made all data underlying the findings in their manuscript fully available?
4. Is the manuscript presented in an intelligible fashion and written in standard English?

Major comments:

• Given that there were no differences in the IgG response between age groups, it would be interesting to join these data, evaluate all the coinfected individuals, and then split the data into Giardia, E. hystolitica. Reply: We are confused by this comment, because Fig 1 shows an increase in both Ab prevalence (Fig. 1A) and Ab levels (Fig 1 B-E) with age in Ab-positive children (Kruskal-Wallis test p values were p<0.001 MSP2 and p=0.05-0.086 (borderline) for the other antigens).

  We believe combining all MAL+,IP+ children into single a group is unwise, since they were infected with a conglomerate of intestinal helminths, cestodes and protozoa (see Table 2). Combining children with such heterogenous infections is unlikely to provide meaningful information.

• I strongly suggest dividing the age of individuals in 0-5, 5-10, 10-15 years-old to partially solve the "N" problem of the groups. Reply: Thanks for the comment. Initially, children were groups into 5-year categories as suggested by the Reviewer, i.e., 0-5, 5-10, 10-15 years old. However, when the data set showed that children aged 1 to 2 did not have intestinal parasites, the results were grouped into 2-year intervals, that allowed us to more closely define the increase in Ab prevalence (Fig. 1A) and Ab levels (Fig 1 B,C,D,E) with age. The purpose of Fig 1 was to determine if age was a variable that needed to be taken into consideration during data analysis.
Because of the absence of molecular Diagnosis and considering that the authors mention the possibility of low parasitemia infections in the MAL- group. It is important to include MAL- individuals in Figure 1.

Reply: We are sorry if we didn’t make the point clear. ALL children who were Ab-positive are included in Fig 1, including those who are MAL+ and MAL-. Because malaria transmission is high in the area, all children in the study had been exposed to P. falciparum and many of the MAL- children were Ab-positive.

It is necessary to compare parasite data with similar regions in Cameroon. Please compare and cite:
• (Malaria and Helminth Co-Infection in Children Living in a Malaria Endemic Setting of Mount Cameroon and Predictors of Anemia from Theresa K Nkou-Akenji et al. 2006)
• Malaria, Helminths, Coinfection and Anaemia in a Cohort of Children From Mutengene, South Western Cameroon from Clarisse Njua-Yafi et al. 2016.

Reply: We thank the Reviewer for pointing out the omission of key references. Information from these studies have been included in the revised Discussion. The text now reads, “…..to those found in other highly [malaria] endemic regions of the country (32), and the prevalence of co-infections was 19.1%, which is similar to the prevalence of co-infections of 18 – 27% reported in other regions of Cameroon (9,44). The references have been added to the reference section.

Do the authors have information about malaria and intestinal parasites last treatments? On page 17, it was commented that Albendazole treatment was frequent in these children. Deworming information will help the readers to understand why the prevalence of intestinal parasites was low compared with other studies in Cameroon. Additionally, reinforce in the discussion section that collecting/reporting that information is valuable for co-infection studies.

Reply: In response to the Reviewer’s suggestion, the following information has been added to the Methods section. “Currently, mass drug administration with albendazole is being performed twice a year by the Ministry of Health, that is usually conducted in schools and symptomatic cases are sent to the local clinic or hospital for follow up treatment.”

(Figure 1 B, C, D, E) use the same scale limits for all plots. This is also useful to understand differences in levels of antigenicity between proteins.

Reply: We understand the comment, but we do not wish to change the Y-axis on Fig 1, since it is risky to make a direct comparison of Ab levels between antigens in serological assays. A number of variables, including parasite strain, the system to produce recombinant proteins, protein purity, the amount of antigen used, number of exposed epitopes, dilution of plasma, etc., influence the overall results. Even when Luminex beads are covalently-coupled with saturating amounts of antigen, it is questionable if direct comparison of MFI can be made between antigens. Although our assays have been optimized and equivalence amounts of antigen used during bead-coupling, comparisons among the antigens may not provide accurate information about immunogenicity. In Figs 1 B, C, D, E, the Y-Axis was selected to show the best distribution of the MFI results.

(table 3) How could the authors explain increased eosinophilia with low levels of helminth infection? This mainly applies to the age group > 9 years-old.

Reply: After age 2, children start becoming infected with helminths, resulting in an increase in eosinophil counts. During the biannual drug treatment campaign, helminthic infections are eliminated, but eosinophilia persists for a period of time. With increasing age, more children in the area become i) infected and ii) re-infected, resulting in an increase in prevalence of eosinophilia.
The authors argue, "First, children living in moist or wet environments where mosquitoes breed and E. histolytica are more abundant would have a high risk of acquiring both infections, that would result in frequent boosting of the Ab response." This explanation for intestinal parasite influence on antibody production alteration is not viable since Giardia's frequency is higher than E. histolytica in the studied population.

Reply: The sentence has been deleted from the Discussion.

The affirmation "Secondly, since malaria and E. histolytica are both amoebae, they might share common antigens, for example, EBA-175 could share homology with an E. histolytica antigen." is false. Plasmodium falciparum is not an is a protozoan. This group belongs to Apicomplexa organisms. For that reason, the hypothesis about correlating Plasmodium and E. histolytica is wrong. Sorry, “amoebae” was a typo. Both Plasmodium falciparum and E. histolytica are protozoans. The Discussion has been revised to read “parasitic protozoa.”

How different are the two Villages Ngali II and Mfou in the central region of Cameroon? Does it exist a difference in humidity and soil moist, once the authors claimed that this variable could explain differences of Entamoeba histolytica?

Reply: The two villages are very similar with no major differences in humidity or soil moisture. The estimated annual average rainfall measures 1600 mm with an annual average temperature of 23°C for Ngali II and for Mfou. According to the National Meteorology agency, the average humidity for the center regions is 83%. Ngali and Mfou are both in the center region of Cameroon about 60km apart. Note: as mentioned above, the words “humidity and soil moisture” have been deleted from the MS.

Minor comments:

• What criteria were used to divide the population into seven groups according to age?
  Reply: The fact that Intestinal parasite (IP) infections was only observed in children >2 years, helped guide separation of the children into seven groups.

• Please specify how anthropometric parameters were used in the study, once they were described but not used in the study. If this information was not used, please remove these sentences.
  Reply: The sentence has been removed.

• Has the studied region presence of Schistosoma haematobium? If the authors have register if this parasite in the area, Did they examined urine samples to discard infections with this parasite?
  Reply: Detection of S. haematobium was not included in the study design because of low prevalence in the study area. A study conducted in this area (and other regions of Cameroon) by Louis-Albert Tchuem Tchuenté et al., (2012) reported a prevalence of S. haematobium of only 1.72%. Since a large sample size would be required to assess the impact of this pathogen on the Ab response to malaria, S. haematobium was not included in the study.

• Were the individuals asymptomatic to intestinal parasites infection too? No diarrhea, abdominal pain, etc.? Please clarify.
  Reply: Yes. To make the point clear, the Methods section has been revised and states that all children with clinical cases of malaria or intestinal parasites were not included in the study and referred to the local clinic/hospital by the attending physician for treatment. Thank you for the comment.

• (Page 6) It was mentioned that Plasmodium parasitemia was quantified. Did the authors observe any correlation between the Plasmodium parasite burden and the levels of IgG responses to the antigens?
  Reply: As expected, there was no correlation between parasitemia and malaria antibody levels.
• (End of Page 7) Please specify: If the cut-off is MFI+3*SD, how the standard deviation was calculated if the negative controls were pooled? Was this experiment repeated or used replicates? Traditionally, the negative controls are tested simultaneously in different wells of the plate, and the cut-off is calculated from those values.

Reply: Pooled negative control plasma sample were run in triplicates on the same plates as the test samples in all experiments, as well as the positive controls. The cut-off was obtained by calculating MFI+3 SD of the triplicates on all plates in the experiment.

• Did the authors analyze the effect of helminth parasite burden (number of eggs/gram of stool) in those individuals with helminths? This valuable information was commented on but never included in the analysis. If not used, I do not see the necessity of describing in the methods section.

Reply: The information has been deleted from the Methods section.

• For data analysis:
  • Before using ANOVA, did the authors checked for the normality of the variables? If yes, please specify, if not, calculate the normality of the variables and the other ANOVA assumptions.

Reply: Yes, ANOVA was used to compare difference in age across the 4 groups (Table 2). However, comparisons of Ab MFI, which are not normally distributed, with age (Fig. 1) were performed using the Kruskal-Wallis test. The Methods section (Data analysis) has been revised. Information in Fig. 1 legend was correct.

• If the authors have not-normal variables, they should use the Kruskal-Wallis non-parametric, and Dunn posthoc tests to verify differences between groups.

Reply: Sorry for the mistake in the Methods section. The Kruskal-Wallis nonparametric test was performed in Fig 1 and 2. A posthoc test was not performed, as the goal was not to determine when peak Ab levels were obtained, but to determine if age had an influence on Ab levels. Since age was a variable, data for all age groups could not be combined, but rather age was taken into consideration during data analysis.

• Please check frequencies described in table 1 (MAL+IP- 58.8%) vs. the values reported in the second line page 9. (59.4%).

Reply: 59.4% is the correct value. The text has been revised.

• Sum of 58.8%+16.9% = 75.7% not 75.6%.

Reply: Thank you for catching the error. The values in Table 1 and text have been revised and are now consistent.

• In table 1, please add a column with P-values to facilitate the interpretation of the differences between groups. Please report statistics of multiple comparisons between groups too.

Reply: The comparisons requested by the reviewer were originally provided in the Table legend. To comply with the request, the p values have been moved to a column labeled “p values” and the method of analysis was retained in the Table legend.

• What is the potential hypothesis to explain the increased values of parasitemia in the coinfected group?

Reply: There is no significant difference in parasitemia between the two groups (p=0.1599). In fact, the higher parasitemia was found in young children who were intestinal parasite-negative (probably because very young children were in this group).
• Please comment in the text the presence of multi-parasitism in the studied individuals.  
Reply: We thank the reviewer for the comment. The following sentence has been added to the Results section. “Interestingly, all of the children had single parasite infections, and polyparasitism was not found.”

• (Page 11 table 3). Please include values of anemia and eosinophilia in individuals coinfected. In the current configuration is constructed is hard to determine the coinfection impact in anemia and eosinophilia values.  
Reply: Table 3 was designed to evaluate the influence of age on malaria, IP, anemia and eosinophilia. The number of co-infections are too small to be divided by age. In an attempt to address the Reviewer’s comment, a separate Table was designed that compares the influence of no infections, malaria-positive only, and co-infections on percent with anemia and eosinophilia. The Table will be uploaded as supplemental Table 1. It essentially showed that same results as expected, anemia was associated with malaria and eosinophils were associated with co-infections.

• (Page 11). In the sentence, "Thus, as children living in these villages increased with age, they developed partial immunity to malaria and anemia declined; whereas, the prevalence of IP and eosinophilia increased." In this sentence, it is necessary to specify that "protection" is protection against malaria symptoms. The table clearly shows that the frequency of malaria does not decrease with age, only the anemia.  
Reply: The sentence has been revised to read: “Thus, as children living in these villages increased with age, they began developing partial immunity to malaria symptoms and anemia declined; whereas, the prevalence of IP and eosinophilia increased.

• Please plot Age vs. Antibody levels for each protein to verify the correlation for each protein studied.  
Reply: The figure on the right confirms that Ab levels increase with age. The figure shows a linear regression analysis of Ab levels for MSP1, MSP2, MSP3 and EBA-175 using data from all 320 children, and includes the equation for the regression line, the R² value (all positive), and p value (all significant). Thus, the figure confirms that Ab levels increase with age. We do NOT wish to include this figure in the MS since it is essentially identical to the one shown in Fig 1 B, C, D and E. In fact, we feel that the information in Fig 1B-E is easier for the reader to understand.  
Note: If the figure is not shown, it is provided in a separate document.

• As an exploratory analysis, I suggest joining all data and make a boxplot comparing MFI between MA1-PI-, MAL-PI+, MAL+PI-, and MAL+PI+. Mainly for MSP1, MPS2, and MSP3 group age 3-10 and 11-15 to check.  
Resolution analysis of Ab levels for each protein. The Table will be uploaded as supplemental Table 1. It essentially showed that same results as expected, anemia was associated with malaria and eosinophils were associated with co-infections.
Reply: We thank the Reviewer Thanks for the suggestion concerning exploratory analysis. A comparison of Ab levels in two of the above groups (MAL-,IP-, and MAL+,IP-) is shown in Fig 2. Unfortunately, the number of children in the MAL-,PI+ group is too small to provide valuable information. As stated above, children in the MAL-,PI+ group (n=54) are infected with a variety of intestinal helminths, cestodes and protozoa (see Table 2). With such a diverse range of pathogens, plotting the data as a boxplot will not provide useful information. In Fig. 2, the distribution of Ab levels in children co-infected with malaria and single intestinal pathogens is provided. We feel this approach is more informative than “dumping all pathogens together.”

• The sentence "E. histolytica is a gut amoeba that causes both intestinal and extraintestinal infections such as amebic colitis (dysentery) and liver or brain abscess. The protozoa cause a marked down-regulation of macrophage functions rendering the cells incapable of antigen presentation and unresponsive to cytokine stimulation (57)” does not explain the increase of antibody production in E. histolytica infected group. Why could a diminishing antigen presentation generate higher levels of anti-Plasmodium antigens?
  
  Reply: Very true! Not sure why that statement wasn’t caught. The Discussion has been changed significantly. It now reads, “The decrease in macrophage function does not explain the increase in Ab to EBA-175. One possible explanation is that since malaria and E. histolytica...”

Other observations/questions:
• In the title, add "IgG" to Antibody response. Reply: IgG has been added to title (although not all of the co-authors agree this is necessary).

• Check all scientific names of parasite species for correct formatting in italics. (Example Entamoeba histolytica in the Results section in the abstract)
  
  Reply: The scientific name has been checked and are now in italics.

• Please, mention in the background the region where the study was performed.
  
  Reply: This information was included in the background section of the Abstract. It is also included in the Materials section.

• It is necessary to describe and discuss the role of MSP1, MPS2, MSP3, and EBA-175 as markers in serological studies.
  
  Reply: This information has been added to the Discussion.

• Considering that coinfection prevalence is relatively low, I consider that it is important to discriminate with colors or point shapes the individuals MAL-IP-, MAL+IP-, MAL-IP+, MAL+IP+ in Figure 1 B-C-D-E
  
  Reply: We thank the Reviewer for the suggestion. However, information in Fig 1B-E is designed to address the question, are Ab prevalence and levels influence by age? Whereas, Fig 2 provides comparisons between individuals infected with malaria alone or co-infected with specific intestinal parasites. Thus, colored dots or symbols are not needed in Fig 1 (and could be confusing to the reader).

• In page 6 subtitle "Laboratory detection, quantification and speciation of malaria parasites.”, I will not use speciation here. I suggest "Diagnosis and quantification of Plasmodium sp. parasites.
  
  Reply: The header has been changed to read: “Laboratory detection of malaria parasites.”

• (Page 14-15) What type of parasite is "Amoeba"? What is the difference between "Amoeba" and E. histolytica? Traditionally, E. histolytica is considered an amoeba too.
Reply: The figure has been revised to read Intestinal Protozoa. Thanks for pointing out the mis-classification.

• In table 1, to facilitate reading, please remove symbols % and /ul located in cells with data and add to the columns describing the variables.
  Reply: The symbols in the data cells have been removed.

• For consistency, unify parasitemia vs. parasitemia, anemia vs. anemia in the text and plots.
  Reply: The British spelling of parasitaemia, anaemia, and haemoglobin have been used throughout the MS.

• (Page 10) change "The major of helminth parasites" to "The most frequent helminth species detected."
  Reply: The change was made as suggested.

• (Table 2) Check all the total numbers for the "Total IP+" column. For example, for protozoans, the sum is 29+19+4 = 48, and it was reported 47
  Reply: This has been verified and corrected to 48 in Table 2

• (Page 13) In plot titles Change Ab (Antibody) to IgG
  Reply: We thank the Reviewer for the comment, but decide not to make the change. Our rationale is that by definition, IgG is a class of immunoglobulin found in the blood; whereas, Ab are plasma proteins that bind specifically with an antigen. What was measured was IgG Ab. Since the serological assay measured IgG Ab that were recorded as MFI (median fluorescence intensity), we think the labels on the Y-Axis (Ab levels -MFI) reflect what was done. The Methods section makes it clear that the Ab were of the IgG class. [Note: Serum IgG levels (which implies mg/ml) were not measured.]

• (Figure 1E) Add, Change from EBA to EBA-175.
  Reply: Change has been made.

• Please verify all references formatting (For example, reference 42 is all in capital letters)
  Reply: References have been edited as requested by the reviewer.

Review #3: Comments were in the attachment.
  Reply: In revising the MS, all requested changes were made and additional information provided in the text, including information on the BLAST search. The only request we would not fully address is the prevalence of bednet use in the villages. The only information available is that very few children use bednets. Since the slide-positivity rate of 75.6% for P. falciparum, it is unlikely the bednets are having a major influence on the current study. The following information has been added to the MS in the Results section. “To determine if higher Ab levels in children co-infected with P. falciparum and E. histolytica might be due to cross-reactive epitopes, a BLAST search for sequence homology between EBA-175 and E. histolytica proteins was made. No similarities were found using Metablast, and only one hit was found using discontinuous metablast which had a span of only 38 nucleotides (~12 amino acids). Thus, there does not appear to be shared epitopes between these two pathogens that would explain the increase in Ab to EBA-175 in children with co-infections.”
Figure for Reviewer #2 confirming an increase in antibody levels with age.

Distribution of MFI for all 320 children by age. Figure show the regression line +/- 95% CI. These data confirm that between the ages of 1 to 15 years, the amount of Ab increases with age, as the results of increasing Ab prevalence and Ab levels.