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Population level changes in schistosome-specific antibody levels following chemotherapy

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Summary
Aims: Previous studies have reported that chemotherapy of schistosomiasis by praziquantel in humans boosts protective antibody responses against S mansoni and S haematobium. A number of studies have reported schistosome-specific antibody levels before and after chemotherapy. Using these reports, a meta-analysis was conducted to identify predictors of population level change in schistosome-specific antibody levels after chemotherapy.

Methods and results: Following a systematic review, 92 observations from 26 articles published between 1988 and 2013 were included in this study. Observations were grouped by antigen type and antibody isotypes for the classification and regression tree (CART) analysis. The study showed that the change in antibody levels was variable: (a) between different human populations and (b) according to the parasite antigen and antibody isotypes. Thus, while anti-worm responses predominantly increased after chemotherapy, anti-egg responses decreased or did not show a significant trend. The change in antibody levels depended on a combination of age and infection intensity for anti-egg IgA, IgM, IgG1, IgG2 and anti-worm IgM and IgG.

Conclusion: The study results are consistent with praziquantel treatment boosting anti-worm antibody responses. However, there is considerable heterogeneity in post-treatment changes in specific antibody levels that is related to host age and pre-treatment infection intensity.

Keywords
antibody levels, human, meta-analysis, population, praziquantel, schistosomiasis

1 INTRODUCTION

Schistosomiasis is a water-borne parasitic disease of great public health importance mainly in sub-Saharan African countries.¹ Currently, there is a major effort to have schistosome control programmes in all schistosome endemic countries in Africa.²-⁴ There is a need to understand all of the potential effects and implications of schistosome antihelminthic treatment in affected populations. One such potential impact of treatment is on the host immune response against schistosome parasites.⁵,⁶

It is well documented that naturally acquired immunity against schistosome infections reduces both prevalence and infection intensity in the older age groups in endemic areas.⁷,⁸ However, this protective immunity takes years of chronic infection to develop naturally. There are two reasons why anti-schistosome immunity takes long to develop: first; the parasite is capable of evading host immunity and second; the host requires a threshold of antigens to mount
Non-English articles were included in this study, and several library and the Inter Library Loan of the University of Edinburgh.

Native Chinese speaker for the systematic literature review. Chinese articles were identified and translated into English by a

Praziquantel (PZQ) treatment has been reported to enhance host protective immunity by exposing the parasite's hidden antigens in large amounts and by removing the immunomodulatory effects of adult worms. PZQ treatment allows hosts' immune systems to recognize schistosome parasite adult worm antigens that is required to develop protective immunity. In 2001, a review of seven field studies reported high levels of heterogeneity in the type and magnitude of change in antibody levels after chemotherapy between different human populations. To date, many potential factors have been suggested to explain this variation, such as pre-treatment infection intensity, level of schistosome endemicity, age, sex and co-infection with human immunodeficiency virus (HIV). This is important for understanding the consequences of the epidemiological transition occurring in populations subject to national schistosome control programs through mass drug administration both for the host and the parasite. However, there has not been a systematic review of published studies investigating the potential determinants of the heterogeneity in post-treatment immune responses. Therefore, the objectives of this study are (a) to investigate the pattern of antibody levels change across different human populations and (b) to identify host and parasite factors that affect the antibody levels change after PZQ treatment in the population levels.

2 | METHODS

2.1 | Systematic review

An electronic literature search was conducted using six databases. Web of Science Core Collection, BIOSIS Citation Index and MEDLINE all of which were searched through Web of Science (www.webofknowledge.com). In addition, EMBASE, Global Health and Ovid Medicine were searched through Ovid (ovidsp.tx.ovid.com). The search terms were as follows: "schistosom" AND ("antibod" OR "IgA" OR "IgE" OR "IgM" OR "IgG") AND ("albendazole" OR "mefloquine" OR "artesunate" OR "antihelmint" OR "chemotherap" OR "praziquantel" OR "oxamniquine" OR ("drug" AND "treatment")). This electronic literature search was completed in January 2014. After removing duplicates, a total of 1366 unique articles were identified for consideration in the present study. Titles and abstracts of articles were screened to exclude those that were clearly not relevant. Full texts of potentially relevant articles were then reviewed for further selection. Full texts of the relevant articles were sourced through the Web of Science, Ovid, Google Scholar (scholar.google.com), the University of Edinburgh library and the Inter Library Loan of the University of Edinburgh. Non-English articles were included in this study, and several Chinese articles were identified and translated into English by a native Chinese speaker for the systematic literature review.

2.2 | Inclusion criteria

An article was included in this study if it met all of the following criteria: (a) human study (either sex), (b) infection with *S mansoni* and/or *S haematobium* diagnosed by parasitological egg count, (c) participants treated with PZQ, (d) number of participants reported, (e) schistosome-specific antibody levels reported before and after PZQ treatment, (f) follow-up study conducted within 14-180 days after PZQ treatment, (g) schistosome worm antigen (WWA: whole worm homogenate and/or soluble worm antigen), and/or soluble egg antigen (SEA) used to measure antibody isotype levels, (h) participants potentially exposed to schistosome infection for their life-time and/or longer than 1 year before the study, (i) participants’ ages could be categorized as child (0-10 years old), adolescent (11-21 years old) or adult (≥21 years old).

2.3 | Exclusion criteria

Articles were excluded based on the following exclusion criteria: (a) participants had a previous history of antihelminthic treatment prior to the study participation, (b) participants were treated with any antihelminthic drug other than PZQ (eg, oxamniquine), (c) participants were specifically selected because of co-infection with HIV and/or soil transmitted helminths and/or plasmodium, (d) purified and/or recombinant schistosome antigens were used to measure antibody isotype levels, (e) participants were temporary visitors to endemic areas (ie, travellers), (f) participants were originally from endemic areas but had moved to non-endemic areas prior to the study (eg, immigrants), (g) participants were diagnosed with acute schistosomiasis, (h) clinical case reports from a single patient.

In schistosomiasis endemic areas, co-infection with soil transmitted helminths is frequently reported. These studies were kept in the analysis only if they did not specifically select co-infected participants. Schistosome-specific antibody isotype levels before (baseline) and after (follow-up) treatment with PZQ were extracted from the selected articles. For those articles that reported results only in graphical format, the software DataThief III (2006) was used to extract the raw data, whenever the graph format allowed it.

In addition to antibody levels, the following information was also extracted from each article: the year of publication, article title, names of authors, study area (country, region and village), schistosome parasite species, co-infection status, co-infecting pathogen species, number of participants, age or age range, sex, height, weight, days between the treatment and follow-up, pre- and post- treatment infection intensity and prevalence, PZQ dose and cure rate. Several articles reported results from multiple different groups of participants in the same study area, such as from different age groups. In such cases, the result from each group was recorded as a separate observation. For the purpose of classification and regression tree (CART) analysis, they were treated as independent observations. For articles that reported results from multiple follow-up time points, the first follow-up after 14 days was selected and included in this study. A total of 92 observations...
from 26 articles (published 1988-2013) met all the inclusion criteria and were considered for the final meta-analysis (Figure 1, Tables 1, Appendix S1 and Table S1).

Potential predictors were selected according to their biological importance, as suggested by earlier studies 19,21 and if they were reported by the majority of articles included in this study. The following predictors were considered: age groups (0-10 years old, 11-21 years old, ≥21 years old), pre-treatment infection intensity [low (S mansoni: 1-99 eggs/1 g faeces, S haematobium: 1-49 eggs/10 ml urine) or high (S mansoni: >100 eggs/1 g faeces, S haematobium: >50 eggs/10 mL urine)], schistosome species (S mansoni, S haematobium, or co-infection of S mansoni and S haematobium), disease prevalence (low/moderate: <50% or high: ≥50%), and days between chemotherapy and follow-up. The “low” infection intensity in the current study is equivalent to “light” in the WHO's infection intensity categories both for S mansoni and S haematobium. However, “high” infection intensity is WHO's “heavy” for S haematobium but “moderate” or “heavy” for S mansoni. This was due to the small sample size of S mansoni studies: there were only eight observations from four articles that reported >400 eggs/1 g faeces that is “heavy” infection by WHO guideline.

2.4 | Software

Articles identified by the systematic review were recorded using Thomson Reuters EndNote, and the extracted data were entered in a spreadsheet using Microsoft Excel 2010. B. Tummers, DataThief III. 2006 (http://datathief.org/) was used to extract data from published graphs. IBM SPSS Statistics Version 21.0 was used for statistical analysis. GraphPad Software GraphPad Prism version 6.03 was used to create graphs.

2.5 | Statistical analysis

The majority of studies included in this study used the enzyme-linked immunosorbent assay (ELISA) method to quantify antibody isotype levels and reported optical density (OD). However, OD values cannot be directly compared between studies conducted by different research groups.25 Therefore, the outcome variable was categorized according to the direction of change in antibody levels from pre-treatment baseline to levels at follow-up. That is, pre-treatment and post-treatment antibody isotype levels were compared within the same population and the outcome was categorized as “increase” if the post-treatment level was higher than the pre-treatment level, and “decrease” if it was lower. There were seven observations that reported the exact same value of antibody isotype levels both pre- and post-treatment.10,26,27 The number of those observations were too small to form their own category “no change”; therefore, they were categorized into “decrease” group in this study for analyses purposes. All observations were grouped according to the type of schistosome parasite antigens (WWA or SEA) that were used to measure antibodies and analysed separately. Prior to the data stratification, we conducted preliminary analyses. The analyses results supported the partitioning by antigen type but not parasite species; therefore, WWA and SEA were analysed separately but not parasite species.

There were 29 observations from four articles that failed to report pre-treatment infection intensity of study participants. In these cases, pre-treatment infection intensity was obtained from scientific publications describing the larger populations that contained the study populations (articles listed in Table S2). Similarly, there were three observations from two articles that did not report the schistosome infection prevalence in the study area. In these cases, prevalence was obtained from scientific publications or governmental reports from the same area or larger area that contained the study populations (Table S2).

In schistosomiasis endemic areas, infection intensity peaks in the young age group, giving a convex curve typically observed in schistosomiasis.28 Furthermore, our preliminary analyses using a mixed effects logistic regression model suggested that there is an association between age and infection intensity (data not shown). To take this nonlinear association into account, a combination predictor for age and infection intensity was generated, with format age/infection...
| References            | Parasite species | Antigen type | Antibody isotype                        | Follow-up (days) | No. of participants for each observation and antibody isotype (participants) |
|-----------------------|------------------|--------------|-----------------------------------------|------------------|--------------------------------------------------------------------------|
| Abebe et al. (2001)   | S.m              | SEA          | IgA, IgE, IgG1, IgG2, IgG3, IgG4, IgG, IgM | 35               | 66                                                                       |
| Ali et al. (1994)     | S.m, S.h         | SEA/WWA      | IgA                                      | 90               | 21, 51                                                                   |
| Feldmeier et al. (1988)* | S.m, S.h     | WWA          | IgE, IgG                                 | 150              | 19                                                                       |
| Fouda et al. (2007)*  | S.m              | WWA          | IgE                                      | 180              | 19                                                                       |
| Grogan et al. (1996)* | S.h              | SEA/WWA      | IgE, IgG4                                | 35               | 55                                                                       |
| Hamadto et al. (1990)* | S.m              | SEA/WWA      | IgA, IgE, IgG, IgM                       | 49               | 36, 40                                                                   |
| Hussein et al. (1996)* | S.m             | SEA/WWA      | IgG, IgM                                 | 60               | 26, 38                                                                   |
| Ismail et al. (1992)* | S.m, S.h         | SEA/WWA      | IgG, IgM                                 | 90               | 20, 15, 14, 11 (SEA)                                                     |
| Joseph et al. (2004)* | S.m              | SEA/WWA      | IgE, IgG1, IgG2, IgG3, IgG4              | 35               | 10                                                                       |
| Mutapi et al. (1998)  | S.m, S.h         | SEA          | IgA, IgG1                                | 84               | 132                                                                      |
| Mutapi et al. (1998b)* | S.h             | SEA          | IgA, IgE, IgG1, IgG2, IgG3, IgG4         | 126              | 37                                                                       |
| Nagaty et al. (1996)* | S.m, S.h         | SEA/WWA      | IgA, IgE, IgG, IgM                       | 180              | 21, 51                                                                   |
| Nassr et al. (2002)*  | S.m              | WWA          | IgG1, IgG4                               | 90               | 8                                                                        |
| Naus et al. (1998)*   | S.h              | SEA/WWA      | IgE, IgG1, IgG2, IgG3, IgG4, IgG4, IgM   | 30               | 97                                                                       |
| Reilly et al. (2008)* | S.h              | WWA          | IgG1, IgG3                               | 42               | 89                                                                       |
| Satti et al. (2004)*  | S.m              | WWA          | IgE, IgG4                                | 21               | 28                                                                       |
| Satti et al. (1996)   | S.m              | SEA/WWA      | IgE, IgG1, IgG2, IgG3, IgG4, IgG4, IgM   | 90               | 17                                                                       |
| Satti et al. (1996b)* | S.m              | WWA          | IgA                                      | 90               | 25                                                                       |
| Snyman et al. (1997)* | S.h              | WWA          | IgE, IgG                                 | 21               | 14                                                                       |
| Snyman et al. (1998)* | S.h              | WWA          | IgE, IgG                                 | 90               | 10, 9                                                                    |
| Tweyongye et al. (2009)* | S.m          | SEA/WWA      | IgE, IgG1, IgG2, IgG3, IgG4              | 42               | 89, 124                                                                  |
| van Lieshout et al. (1999)* | S.m     | WWA          | IgE, IgG1, IgG3, IgG4, IgG, IgM          | 63               | 30                                                                       |
| Vereecken et al. (2007)* | S.m         | SEA/WWA      | IgA, IgE, IgG1, IgG2, IgG3, IgG4, IgG4, IgM | 42               | 21, 24, 61, 143                                                         |
| Walter et al. (2006)* | S.m              | SEA/WWA      | IgA, IgE, IgG1, IgG2, IgG3, IgG4, IgM    | 35               | 24, 28 (SEA)                                                             |
| Wilson et al. (2013)* | S.m, S.h         | WWA          | IgE                                      |                   |                                                                           |
| Zinyowera et al. (2006)* | S.h            | SEA/WWA      | IgA, IgE, IgG                           | 42               | 28                                                                       |

S.h, *S. haematobium*; S.m, *S. mansoni*; SEA, schistosome soluble egg antigen; WWA, whole worm antigen.

*The numbers represent the number of participants for each observation of each antigen time (SEA/WWA) and antibody isotypes (IgA, IgE, IgG1, IgG2, IgG3, IgG4, IgM).

*These references are given in supporting information Appendix S1.
intensity as shown in Table 2. All of the observations were categorized according to age/infection intensity categories. All predictors used for CART analysis are listed in Table 2.

2.6 Classification and Regression Tree models

Classification and Regression Tree (CART) models were used to identify influential predictors of the direction of change of schistosome specific antibodies after PZQ treatment. Briefly, CART models are non-parametric and nonlinear analyses which allow us to investigate the pattern of data without making data distribution assumptions of both outcome and predictors. CART models do not make assumptions about the type of association between dependent variable and influential predictors. In addition, CART models do not have mandatory minimal sample size that allows us to apply this model to study with a small sample sizes. Confidence intervals or standard errors are the most common weighting methods for meta-regression and for meta-CART. However, in this analysis the measures of antibody levels were ELISA OD values, which cannot be compared directly when coming from different research groups. Therefore, in this analysis, the sample size (the number of participants) was used for weighting. The number of participants across studies varied from 7 to 148. The potential predictors used for the analysis were as follows: schistosome species (S.mansoni, S.haematobium, or co-infection), days between treatment and follow-up, and disease prevalence (low/moderate or high), and age/infection intensity (child/low, adolescent/low, adult/low, child/high, adolescent/high, adult/high)(Table 2).

The CART analysis was conducted to build a tree using the standard three steps: (a) growing a maximum-sized tree with largest number of subgroups, (b) pruning the tree to generate subtrees and (c) identifying the optimal sized tree with the minimal risk estimate following the methodology of Breiman. Initially, the maximum-sized complex trees were grown with data from all study variables for each antibody isotype. All potential predictors were compared using the Gini index to identify the optimum split of the dependent variable (increase or decrease in antibody level). Based on the Gini index, the strongest predictor variable and its splitting value, that is sub-groupings for categorical variables and cut-off values for continuous variables, were used to split the original data (ie, root node) into two subgroups (ie, daughter nodes). The subgroups were then divided repeatedly into smaller subgroups following the same procedure until they represented the most homogeneous subgroups achievable (ie, terminal node). In this study, terminal nodes of these maximum-sized trees were set to be pure (ie, the node contents only “increase” or “decrease” observations) or with only a single observation. Then a series of subtrees was generated by pruning the initial maximum-sized trees. To estimate the optimal subtree among the different sized sub-trees, 10-fold cross-validation analysis was conducted for each subtree followed by the selection based on the one standard error (SE) rule. Briefly, the cross-validation analysis is used to estimate the risk of misclassification using a randomly selected subset (ie, test samples) of the original dataset (ie, learning sample). The optimal tree is the one that yields the minimal risk estimate. However, the noisy nature of the data and the instability of the cross-validation procedure can lead to the selection of unstable and large trees. Therefore, following the one SE rule, the smallest tree that has a cross-validation risk estimate of less or equal to the minimal risk plus one SE was selected as the optimal tree (Figures S1 and S2).

### Table 2: List of potential predictors investigated and their measurement units/codes

| Potential predictors (units) | Codes |
|-----------------------------|-------|
| Schistosome species         | S.m, S.h, S haematobium. |
| Days after chemotherapy (days) | Continuous (14-180 days) |
| Agea/infection intensityb   | Child/low, Adolescent/low, Adult/low |
|                            | Child/high, Adolescent/high, Adult/high |

S.m, S mansoni; S.h, S haematobium.

aAge category: child (0-10 y), adolescent (11-20 y), adult (≥21).
bInfection intensity: S mansoni: Light 1-99 eggs/1 g faeces, Heavy ≥100 eggs/1 g faeces, S haematobium: Light 1-49 eggs/10 mL urine, Heavy ≥50 eggs/10 mL urine.

3 RESULTS

Following a systematic review, a total of 92 observations from 26 articles (published 1988-2013) met all inclusion criteria and were considered for the final meta-analysis. There was a high degree of heterogeneity in the direction of change of antibodies (increase/decrease) after PZQ treatment depending on population studied (Figure 2). The results showed a tendency towards an increase in anti-WWA antibodies, in contrast with no significant tendency or decrease in anti-SEA antibodies post-PZQ treatment (Figure 2). Five anti-WWA antibody isotypes (IgA, IgE, IgG1, IgG2, IgG4) showed a significant trend of increase ($X^2 = 12.25, P < 0.001$; $X^2 = 8.26, P = 0.004$; $X^2 = 6.55, P = 0.011$; $X^2 = 7.14, P = 0.008$; $X^2 = 10.71, P = 0.001$) (respectively) (Figure 2). In contrast, two anti-SEA antibody
isotypes (IgG, IgM) showed a significant trend of decrease after PZQ treatment ($\chi^2 = 8.07$, $P = 0.005$; $\chi^2 = 4.48$, $P = 0.034$, respectively) (Figure 2). The CART used to determine predictors of antibody level change identified age/infection intensity as the most common factor affecting post-treatment change. Thus, the analysis identified optimal trees for anti-SEA (IgA, IgE, IgG1, IgG2, IgM) and for anti-WWA (IgG, IgM) antibodies (panels A, C, D, E in Figures 3 and 4). Anti-SEA IgG1 showed a tendency to a decrease among adolescents and adults with high pre-treatment infection but an increase among all other age and infection intensity groups (panel B in Figure 5). Anti-WWA IgG showed the tendency to a decrease among children and adolescents with low pre-treatment infection intensity but an increase among all other age and infection intensity groups (panel D in Figure 5). The CART results showed the tendency of anti-SEA (IgA, IgM) and anti-WWA IgM to increase among children and adults, but decrease among adolescents regardless of their pre-treatment infection intensity (panel A in Figure 5). Similarly, the results showed a tendency of anti-SEA IgG2 to decrease among adolescents and increase among children regardless of their pre-treatment infection intensity. In addition, the results also showed that anti-SEA IgG2 tends to increase among adults with low pre-treatment infection intensity but decrease among adults with high pre-treatment infection intensity (panel C in Figure 2). Overall, changes in levels of anti-SEA IgG1, IgG2 and anti-WWA IgG after PZQ treatment were influenced by the combination of age and pre-treatment infection intensity. On the other hand, anti-SEA IgA, IgM and anti-WWA IgM were influenced by age only (Figure 5).

4 | DISCUSSION

Praziquantel (PZQ) is currently the recommended drug for treatment of schistosomiasis. Immunology field studies have reported that PZQ treatment can enhance the development of host protective immunity against future re-infection. Schistosome parasite-specific antibodies are thought to play an important role in this protective immunity. As schistosomiasis endemic areas see increasing PZQ treatment in Mass Drug Administration (MDA) programs, it is important to determine any patterns in population-wide changes in immune responses and their drivers. Here, we conducted
a systematic review and meta-analysis to identify predictors that influence the direction of change in schistosome-specific antibody isotype levels after PZQ treatment in humans.

Our analyses showed that the antibodies' direction of change after chemotherapy is highly variable among different populations. There was only single antibody isotype: anti-WWA IgA that have been reported to increase after PZQ treatment in all the human populations studied. Our analyses also showed that the antibodies to whole worm antigens (WWA) being reported to increase in the majority of studies. In contrast, studies of anti-SEA antibody isotypes showed a weak tendency towards decreased levels after PZQ treatment. PZQ treatment has been reported to damage adult worm tegument, therefore allowing the host immune system to detect schistosome worm antigens that would otherwise not be accessible until those worms die naturally.

This theory is in line with a previous report that PZQ treatment enhances the host immunological recognition of *S. haematobium* specific proteins. The results of the present study are consistent with this hypothesis and that the elevation of some anti-WWA antibodies (IgA, IgE, IgG1, IgG2, IgG4) is less likely to be affected by characteristics of the populations.

There are multiple schistosome-specific antibodies whose association with protection for schistosomiasis has been reported. In particular, schistosome specific IgE and IgA are commonly associated with protection against re-infection after PZQ treatment in humans. Ideally, vaccination programs would be the most effective control strategy for schistosomiasis. Nevertheless, although a number of studies have been conducted to develop an

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**FIGURE 3** Classification and Regression Tree Models identifying profiles of observations that had higher (increase) or lower (decrease/no change) anti-SEA antibody isotype levels after praziquantel treatment for (A) anti-SEA IgA, (B) anti-SEA IgE, (C) anti-SEA IgG1, (D) anti-SEA IgG2 or (E) anti-SEA IgM. No tree was obtained for the remaining anti-SEA antibody isotypes. The hierarchy of the Classification and Regression Tree Model starts from the terminal nodes at the top. Abbreviations for age/infection intensity groups are listed in the text box and described in Table 2.
effective vaccine for schistosomiasis, we still do not have any licensed schistosome vaccine. A review of animal studies has shown that attenuated schistosome parasite vaccines are more efficient than recombinant purified protein vaccines.\textsuperscript{43} One potential reason for this improved protection is that attenuated parasite vaccines expose hosts to a large variety of schistosome parasite antigens with a single vaccination and this strategy may provoke an overwhelming response. Similarly, PZQ treatment and damage it does to the worm may release an equivalently large number of antigens that boost acquired protective immunity. Therefore, although the main purpose of PZQ treatment is to cure infected people, it is likely that a beneficial side effect is an increased protection against future re-infection. In line with this theory, our meta-analysis shows that indeed, both anti-WWA IgE and anti-WWA IgA are elevated after PZQ treatment in the majority (but not all) of study populations. Those reports together with our anti-WWA IgE and IgA results confirm the “booster” effect of PZQ treatment on a proposed correlate of acquired protective immunity among schistosome affected populations.

There are a number of studies that have reported the potential immunizing effect of PZQ treatment among affected populations. Nevertheless, the length of protection that could be induced by PZQ treatment remains unclear. Our analysis demonstrated the tendency for both anti-WWA IgE and anti-WWA IgA levels to be elevated during a 180-day period after treatment which suggests at least a 6 month protective window. Naus et al. (1998) conducted an immunological study among Cameroonian children and found that anti-WWA IgE levels were elevated 1 month after the PZQ treatment but then became lower than pre-treatment baseline level 12 months after the treatment.\textsuperscript{44} That report together with our results indicates that the immunizing effect induced by PZQ treatment can last for as long as 6 months post-treatment, but may not last longer than 12 months. Further epidemiological studies are therefore required to confirm the dynamics of anti-WWA IgE and IgA after PZQ treatment to estimate the length of this induced protective immunity.

\textbf{FIGURE 4} Classification and Regression Tree Models identifying profiles of observations that had higher (increase) or lower (decrease/no change) anti-WWA antibody isotype levels after praziquantel treatment for (A) anti-WWA IgG or (B) anti-WWA IgM. No tree was obtained for the remaining anti-WWA antibody isotypes. The hierarchy of the Classification and Regression Tree Model starts from the terminal nodes at the top. Abbreviations for age/infection intensity groups are as listed in the text box in Fig 3, also described in Table 2.
The analysis revealed a significant proportion of studies showing an increase rather than decrease in anti-WWA IgG4 (86% of observations reported increase) after treatment. Longitudinal and cross-sectional population studies have demonstrated the association of both anti-SEA IgG4 and anti-WWA IgG4 with human susceptibility to re-infection after treatment in schistosomiasis endemic areas.45,46 In particular, IgG4 has been suggested as a possible blocking antibody that inhibits the action of protective IgE in both *S. haematobium* and *S. mansoni* infections.39,47,48 Field studies have reported that the ratio of IgE to IgG4 has a positive influence on resistance to future *S. mansoni* re-infection.49,50 There were multiple observations that reported the direction of change for both IgE and IgG4 after praziquantel treatment. The half of those anti-SEA studies (nine observations) have reported the post-treatment increase of IgE and decrease of IgG4, supporting the immunizing effect of praziquantel treatment. On the other hand, the majority of anti-WWA studies reported the post-treatment increase of both IgE and IgG4. More research is required to determine how the ratio of these two antibodies changes after chemotherapy, and how they relate to protection against re-infection.

The CART analysis results showed that for several antibody isotypes, the direction of change after chemotherapy can be partially explained by a combination of the population’s age and pre-treatment infection intensity. Pre-treatment infection intensity determines the quantity of schistosome parasites’ antigens that participants are exposed to following PZQ treatment. On the other hand, in endemic areas, participant age correlates with the cumulative history of exposure to schistosome parasites.51 A study in baboons reported that attenuated schistosome parasite vaccines were effective regardless of the different schistosome infection history (naïve or infected then PZQ-treated).52 Supporting this, our results suggest that some isotypes that offer putative protection against re-infection (anti-WWA IgA and IgE) are boosted by praziquantel treatment for a minimum of 6 months post treatment regardless of age. However, for other putatively protective isotypes (anti-SEA IgA) our results do indicate an effect of age. Overall, our result suggests that when schistosome parasite vaccine for human use becomes available, the effectiveness of the vaccine might be affected by not only vaccination dose but also by the age/infection history of participants. Effective vaccine development would benefit from investigating the immunological variability of the target human population.

There are limitations of the current study. First of all, CART analysis does not allow the random effects to be taken into account. Therefore, although multiple observations were extracted from a single article, random effect was not considered in the current study. A preliminary analysis was conducted using mixed effects logistic regression models, which allowed random effect in the model. However, these models were not considered further. This was mainly because the majority of the models had very high information criterion values, indicating instability (results are not shown). Additionally, there are some previously identified candidate predictors such as sex21 and co-infection with human immunodeficiency virus (HIV),22 which could not be considered in the current study due to the fact that only a limited number of reports were available.

This meta-analysis revealed that literature studies, on the whole, tended to report an increase of anti-WWA antibodies isotypes after PZQ treatment. Our analyses also show a considerable variability in change among different antigens, antibody isotypes and populations following treatment with PZQ, confirming the work of Mutapi (2001).18 Although the combination of age and infection intensity, alongside the number of days after treatment, were identified as influential predictors for some antibody isotypes, there is no single predictor that consistently affects all antibody isotypes in the same way. Our results also demonstrated that antibody isotypes that have been reported to have a protective effect against future re-infection (anti-WWA IgA, IgE) can be stimulated by PZQ treatment in the majority of cases for a minimum of 6 months after treatment. These results, therefore, reinforce previously reported protection enhancement following PZQ treatment.

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**DISCLOSURES**

None.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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