REVIEW

Early lessons from schistosomiasis mass drug administration programs [version 1; peer review: 3 approved]

W. Evan Secor
Parasitic Diseases Branch, Division of Parasitic Diseases and Malaria, Center for Global Health, Centers for Disease Control and Prevention, Atlanta, GA, 30329, USA

Abstract
Mass drug administration using praziquantel is the backbone of the current strategy for the control of schistosomiasis. As the theoretical plans have moved into practical application, certain challenges with this approach have surfaced, and it is likely that annual mass drug administration alone may not be sufficient to achieve program goals. However, mass drug administration is still the only available intervention that can be readily used in the wide variety of settings where schistosomiasis is endemic. The task then becomes how to improve this approach and identify what adjuncts to mass drug administration are effective, as programs move from morbidity control to elimination goals. Other aspects worthy of consideration include how best to employ new diagnostic tools to more easily identify where treatment is needed, and new formulations of praziquantel to extend the availability of treatment to all age groups. The aim of this review is to highlight both areas of challenge and of opportunity to improve the public health impact of schistosomiasis control programs.

Keywords
schistosomiasis, mass drug administration, praziquantel, schistosome
A new focus on schistosomiasis control and elimination

Over the last decade, there has been an increased emphasis on schistosomiasis control, especially with respect to using mass drug administration (MDA) to reduce its prevalence and the intensity of infections. This shift has largely been driven by the introduction of the preventative chemotherapy approach for neglected tropical diseases and the passage of World Health Assembly (WHA) resolutions 54.19 (2001; http://www.who.int/neglected_diseases/mediacentre/WHA_54_19_Eng.pdf?ua=1) and 65.21 (2012; http://www.who.int/neglected_diseases/mediacentre/WHA_65_21_Eng.pdf) that set control and elimination goals for schistosomiasis, respectively. In addition, access to treatment for those persons in need has dramatically increased as a result of aid agency purchase and manufacturer donation of praziquantel, the only drug currently available for the treatment of schistosome infections, along with efforts of groups such as the Schistosomiasis Control Initiative (SCI, http://www3.imperial.ac.uk/scisto) to work with ministries of health to distribute praziquantel. Finally, increased support for operational research on how best to distribute treatment has facilitated the posing of questions that have direct public health impact in a more widespread approach than was previously possible. While there have been clear public health benefits associated with schistosomiasis MDA in certain settings1-3, it has not been an unqualified success4-6. This review will focus on some of the practical questions that have surfaced with the introduction of MDA for schistosomiasis and issues that have implications for the successful implementation of schistosomiasis control and elimination programs.

Challenges with providing MDA

Despite the dramatic expansion of praziquantel availability, data from 2013 indicate that of the more than 260 million people in need of treatment for schistosomiasis, less than 40 million received it. The shortfall can be attributed to a number of factors, including a remaining deficit (about 120 million treatments) in the amount of available praziquantel1. Furthermore, even with significant price reduction or donation of the drug, the costs associated with identifying where MDA is needed and the delivery of the drugs create barriers for many national control programs in the absence of external funding to support these activities. Other obstacles include the lack of compliance with treatment programs by persons needing to take the drug. While health education increases community cooperation with praziquantel delivery programs, the side effects (fever, nausea, abdominal pain, diarrhea, fatigue) associated with dying worms, especially the first time someone is treated when worm burdens tend to be the highest, can lead to wariness about receiving follow-up treatments4-11. Further research on how best to promote participation in treatment campaigns, some of which will need to be tailored to specific countries, languages, or ethnic groups, is essential. MDA strategies are geared towards school age children but adults and pre-school aged children also contribute to ongoing transmission, suggesting that achieving elimination will require treating these other age groups as well. There is also a growing recognition that even very young children can become infected with schistosomes and suffer health consequences. However, current formulations of praziquantel are not appropriate for MDA in this age group because of the size and taste of the tablets. The therapeutic dose younger children need may also differ from that required by older individuals, as suggested by recent studies in Uganda12.

Fortunately, efforts are currently underway to develop a pediatric formulation of praziquantel and to define the dosing regimen. As this formulation becomes available, operational research on how to carry out treatments targeted to young children will be needed, in parallel with the development of treatment delivery approaches to improve compliance among school children and adults.

Advances in determining where to treat

The initial requirement for any control program is to accurately determine the areas where treatment is needed. For schistosomiasis, this has traditionally been done by parasitologic assessment of stool or urine samples, depending on the schistosome species endemic in the area. Although parasitologic methods have recognized limitations in sensitivity, they were (until very recently) the only feasible option for estimating the prevalence of intestinal schistosomiasis and the only way to monitor the intensity of human infection for any of the species. As a result, the current World Health Organization (WHO) guidelines for schistosomiasis control are heavily dependent on detection of eggs in stool for Schistosoma mansoni and S. japonicum or urine for S. haematobium13. Recently, a point of contact (POC) test that detects a S. mansoni carbohydrate antigen in the urine of infected individuals has become commercially available. This circulating cathodic antigen (CCA) POC test can indicate a relative intensity of infection and distinguish active infection, or reinfection, from cure following treatment. A large number of studies have evaluated the POC-CCA in comparison to stool examination by the Kato-Katz method and found that it is at least as good as traditional stool examination for mapping areas in need of MDA14-19. In general, the POC-CCA appears to be more sensitive than traditional stool examination methods but questions remain about whether disparities in results obtained when comparing the two methods are attributable to the known insensitivity of the Kato-Katz method or imperfect specificity of the POC-CCA test20. However, when considering all the expenses associated with laboratory testing and treatment-associated expenditures, the costs of using either test are comparable10,21. Because the POC-CCA does not require equipment, it should be easier to deploy than the Kato-Katz method in areas that need mapping for S. mansoni prevalence. Nevertheless, training for POC-CCA use and interpretation will be needed and there is a distinct need to develop bench aids for this test. It is also not possible to simply apply the WHO guidelines that were written for morbidity control based on stool exam prevalence levels to the POC-CCA, which consistently detects higher prevalence levels. Thus, while the introduction of the POC-CCA is perhaps the most important technical advance for S. mansoni control since the release of praziquantel, more operational research is needed before it can achieve its full potential and be incorporated into WHO recommendations. The POC-CCA may also be useful for detecting S. japonicum infections22.

The paradox of the POC-CCA test is that, even though it uses a urine sample for the assay, it is not a reliable test for S. haematobium infections. Fortunately, another carbohydrate, the circulating anodic antigen (CAA), is effective for detecting both urogenital and intestinal schistosomiasis23-25. It is also considered more sensitive and specific for S. mansoni than the POC-CCA. The drawback of the CAA test is that it involves equipment-requiring processing of samples prior to testing and the output is a non-visual signal
that requires an automated reader. However, research is ongoing to develop the CAA into a more field applicable test that would have the advantage of detecting both intestinal and urinary schistosomiasis.

The detection of specific antibodies may also become important for control and elimination programs for schistosomiasis, although they are likely to be employed in different settings, or at different phases of the program than egg or antigen detection tools. An advantage of antibody assays is the ability to directly observe the collection of finger stick blood from the population being surveyed. Although urine collection is easier than stool collection, it is not feasible or culturally acceptable to directly observe the collection of either and therefore both present an increased risk of sample sharing. In addition, small quantities of blood can be used in multiplex assays that may simultaneously test for a variety of neglected tropical diseases, other infectious agents, and monitor vaccine coverage. Thus, a single sample can be used for several public health programs, thereby providing cost savings, compared to performing an independent survey for each infection or vaccine response of interest. Many low-cost rapid diagnostic tests are based on antibody detection and could be adapted for schistosomiasis, provided the proper antigen was selected. It is also theoretically possible to develop pan-schistosome or species-specific antigens depending on the intended use of the assay. A major drawback with using antibody-based assays is that the current antigens that are used for immunodiagnosis are recognized by host sera even after successful cure. Thus, it is not possible to distinguish former infections from active infections with great certainty, and therefore not possible to monitor decreases in prevalence levels as a control program progresses. Antibodies are also a less reliable indicator of intensity of infection than egg or antigen detection methods. However, antibody detection will likely be very useful for schistosomiasis elimination programs; children born after cessation of transmission would not be exposed to infection and thereby become very sensitive sentinels to confirm that elimination has been achieved. Further, as immunodiagnostic tests are developed that use individual antigens rather than antigen mixtures, immunoaassays that are positive during active infection but become negative shortly after treatment may become available. One antigen that shows promise in this regard is recombinant SP-13 from S. japonicum. Detection of parasite DNA in stool or urine could also be a sensitive method for specifically identifying active infections.

MDA plus what?

Current schistosomiasis control guidelines are based on different frequencies and target populations of MDA, with the most intensive intervention being annual MDA of all community members. This approach is predicated in large part on the benefits of annual MDA for reducing prevalence of lymphatic filariasis, onchocerciasis, and blindness, and in the near term it is likely that other interventions involving water are the most likely adjuncts to MDA for reducing infection. An unfortunate side effect of the introduction and initial success of praziquantel was the assumption that treatment alone would be adequate to reach program goals. This belief contributed to reduced support for research into other control approaches. Because of the expense of water and sanitation systems, efforts to reduce urine or fecal contamination of water, as well as limiting exposure of people to contaminated water, will rely heavily on health education and behavioral modification. These approaches may need to be specifically tailored to individual communities and are therefore difficult to readily apply across endemic areas. The control of intermediate host snails can be highly effective but most interventions that have demonstrated success involve the introduction of molluscs and other aquatic species or entail the introduction of non-native snail predators, both of which have limited acceptability for local populations or groups with environmental concerns. One exciting idea that does not suffer from these limitations is represented by The Upstream Alliance project, which will reintroduce native Macrobrachium vollenhoveni prawns in the Senegal River upstream of the Daima Dam. Following completion of this dam in 1986, there was an outbreak of new schistosome infections that has in part been attributed to the interruption of the prawn’s ability to move up the river from its breeding grounds in brackish water. M. vollenhoveni are voracious predators of the schistosome intermediate host snails. The restriction of their migration led to an expansion of snail populations, which in turn led to increased transmission of schistosomiasis. The Upstream Alliance hopes to couple prawn aquaculture with the reintroduction of prawns above the dam to create an economically self-sustaining intervention to reduce schistosome infection prevalence. If successful, it will be a robust model for snail control programs in other areas with native Macrobrachium spp. populations.

Adapting programs as they progress

One of the "staged control" strategy for schistosomiasis, program objectives change from morbidity control to reduction of infection to elimination of transmission to post transmission surveillance, depending on the prevalence levels of the population. One of the biggest unknowns for schistosomiasis control is what prevalence cutoffs merit changes in treatment strategies and, in fact, what those different strategies should be as the goals of the program change. As mentioned above, the current WHO guidelines for schistosomiasis were developed at a time when praziquantel was less abundant and more expensive, and therefore the primary goal was the reduction of severe hepatosplenic disease for intestinal schistosomiasis and the prevention of bladder and kidney complications for urogenital schistosomiasis. Now that praziquantel is more readily available, it is recognized that treatment should be extended more generally, as it is not only those with the most severe manifestations of schistosomiasis that suffer morbidity. In fact, a strong case has been made that there is no such thing as an asymptomatic schistosome infection. Thus, there may really be no practical differences between reducing morbidity and reducing infection.
Monitoring of control program progress, like mapping, has traditionally been performed by measuring egg prevalence and the intensity of schistosome infections in school age children because they provide a useful barometer of the level of infection in other age groups in the community. This age group also tends to have the highest intensities of infection, so decreases in their egg output would result in fewer eggs that could contaminate fresh water and infect snails. Fewer infected snails would lead to the release of fewer infectious cercariae and a theoretical reduction in the “force of transmission”. A test to measure force of transmission in areas where people come into contact with water would be a more timely way to assess the impact of control efforts, rather than having to rely solely on measuring changes in infection levels in people. Unfortunately, previous attempts to measure the number of cercariae in water have not been successful. Sentinel mice and snail sampling, followed by cercarial shedding or PCR, have been somewhat informative but have not yet been incorporated as ways to monitor the impact of control programs. A method that has been somewhat successful in assessing water bodies for cercariae of avian schistosomes, ultrafiltration followed by real-time PCR, can detect as few as 5 cercariae in 100 liters of water. If this method will also work for detecting cercariae of schistosomes infectious for humans, control programs may have a technique to more directly assess the impact of different control efforts.

Is praziquantel sufficient?

Another potential concern for treatment programs is the incomplete efficacy of praziquantel. The introduction of MDA for schistosomiasis has, fortunately, not resulted in evidence of widespread clinical resistance to drugs. However, it has long been recognized that a single treatment, especially for persons with high intensities of infection, is not adequate to kill all the worms. This finding has been further highlighted in recent studies comparing parasitologic and antigen detection assays following treatment. Many individuals become egg negative but retain antigen positivity, suggesting that viable adult worms remain even if egg excretion has stopped. From a transmission perspective, and perhaps even from a morbidity standpoint, if egg laying stops and does not resume, the treatment has accomplished its goal. However, worms that are not killed may only be temporarily affected and subsequently resume egg laying. Interpretation of post treatment data is also complicated by the decreased efficacy of praziquantel against juvenile worms that can produce eggs once they mature. Provision of a second dose of praziquantel to target worms that may not have been killed by the first treatment produces increased cure rates and greater egg reduction than a single treatment. These studies highlight the need to continue research into improved formulations of praziquantel, if there are more effective dosing schedules than annual MDA depending on a location’s force of transmission or schistosomiasis species, as well as control versus elimination program goals. There is also concern that reliance on a single drug is risky, which argues for continued investigation into new or repurposed drugs for treatment.

Conclusions

The prospects for a global reduction in schistosomiasis prevalence and intensity are better now than ever before. Sub-Saharan Africa remains the biggest challenge, although even more developed countries like Brazil and China that have had control programs in place for many years still have much work remaining to achieve elimination. In addition to the WHA resolutions, the formation of groups such as SCI, the Schistosomiasis Consortium for Operational Research and Evaluation (SCORE), the Coalition for Operational Research on Neglected Tropical Diseases (COR-NTD) and, most recently, the Global Schistosomiasis Alliance (GSA) has provided better opportunities for researchers to interact with WHO, ministry of health officials, and schistosomiasis control program managers to identify and test practical solutions for challenges encountered where MDA has been used, and to begin to define and test the strategies for effecting and verifying elimination where appropriate. The pending completion of multi-year operational research studies should provide strong data for the development of updated evidenced-based guidelines for schistosomiasis in the near future. However, progress and answers will take time, requiring patience from donors and governmental aid agencies. Similarly, when prevalence levels decrease and schistosomiasis becomes a lower public health priority, ministries of health in endemic countries will need to maintain control activities amidst competing agendas to achieve elimination. Diagnostic tools with improved sensitivity and specificity, as well as operational research on how to employ them, are critical needs for elimination strategies in areas with decreasing prevalence and intensity of infection. Continued coordination of efforts, along with innovative thinking to identify effective interventions to complement MDA, will be necessary to reduce the public health burden and ultimately eliminate schistosomiasis.

Abbreviations

CAA, circulating anodic antigen; CCA, circulating cathodic antigen; MDA, mass drug administration; POC, point of contact; WHA, World Health Assembly; WHO, World Health Organization.

Disclaimer

The findings and conclusions in this report are those of the author and do not necessarily represent the official position of the Centers for Disease Control and Prevention.

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