Ultrasound Monitoring of Structural Urinary Tract Disease in *Schistosoma haematobium* Infection

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A major advance in our understanding of the natural history of *Schistosoma haematobium*-related morbidity has come through the introduction of the portable ultrasound machines for non-invasive examination of the kidneys and bladder. With the use of generators or battery packs to supply power in non-clinical field settings, and with the use of instant photography or miniaturized thermal printers to record permanent images, it is possible to examine scores of individuals in endemic communities every day. Broad-based ultrasound screening has allowed better definition of age-specific disease risks in urinary schistosomiasis. Results indicate that urinary tract abnormalities are common (18% overall prevalence) in *S. haematobium* transmission areas, with a 2-4% risk of either severe bladder abnormality or advanced ureteral obstruction. In longitudinal surveys, ultrasound studies have shown that praziquantel and metrifonate therapy are rapidly effective in reversing urinary tract abnormalities among children. The benefits of treating adults are less well known, but research in progress should help to define this issue. Similarly, the prognosis of specific ultrasound findings needs to be clarified, and the ease of sonographic examination will make such long-term follow-up studies feasible. In summary, the painless, quick, and reproducible ultrasound examination has become an essential tool in the study of urinary schistosomiasis.

Key words: ultrasonography - diagnostic imaging - schistosomiasis haematobia - hydronephrosis - bladder diseases

Urinary schistosomiasis afflicts an estimated 111 million persons in Africa and the Middle East (van der Werf & de Vlas 2001). Although *Schistosoma haematobium* infection may sometimes result in only minimal pathology, there are a significant proportion of infected patients (25-50%) who experience moderate to severe morbidity (King et al. 1988). Population surveys in endemic areas indicate that hematuria, dysuria, anemia, and inflammatory urinary tract pathology are the most common forms of *S. haematobium*-associated disease (Chen & Mott 1989). Urinary schistosomiasis typically manifests as a lifelong process of tissue injury because, although individual *S. haematobium* worms survive only 4-5 years, multiple worm infections (and reinfection) are common in high-risk areas, so that parasite-related inflammation and fibrosis continue for decades. After 20 to 30 years of infection, the late complications of *S. haematobium* infection begin to emerge clinically. These more life-threatening, late disorders include kidney dysfunction, obstruction of the ureters or bladder outflow, and/or urothelial metaplasia and cancer formation (King 2001).

With the advent of safe and effective oral therapy, repeated drug treatment has become the favored approach to controlling disease due to urinary schistosomiasis (WHO 1993). The aim of this therapy is to reduce infection intensity and thereby minimize (or even prevent) tissue damage and risk for disease. Given the expense of drug delivery on a population basis, certain operational questions arise: i.e., What is the optimal timing of drug treatment to prevent or regress structural morbidity? and Are there persons at special risk for morbidity, for whom more intensive therapy should be provided?

Because structural morbidity caused by *S. haematobium* is readily demonstrated on ultrasound examination, the sonogram has become a very useful tool for defining the population prevalence of infection-associated disease. Ultrasound-based population surveys have permitted an effective assessment of the reversibility of structural disorders among different age groups, and they are beginning to provide the essential data for decision analysis on the choice of treatment regimens for long-term population-based drug treatment programs.

**MATERIALS AND METHODS**

The findings summarized in this paper are derived from cross-sectional and longitudinal morbidity surveys performed in Coast Province, Kenya, primarily in the Msambweni area of Kwale District. This area is highly endemic for *S. haematobium*, but does not support transmission of other forms of schistosomiasis, i.e., *S. mansoni* or *S. intercalatum*. Details of these studies, performed during the years 1984-1992, 1997, and 2000-present, have been previously published (King et al. 1988, 1991, 1992, Olds et al. 1999, Subramanian et al. 1999, Hodder et al. 2000). Target study populations have comprised: (a) school age children from area villages and (b) total populations of selected villages within the study area. All participants in these studies gave informed consent and were studied under protocols approved by the Human...
Ultrasoundographic examinations were performed in the field with the use of portable ultrasound machines powered by generator (initially Toshiba Sonolayer SL, later Shimadzu model SDU-350A) or a rechargeable battery pack (Sonosite 180). Urinary tract pathology was detected by sector scanning and scored according to standardized protocols (Subramanian et al. 1999, Richter et al. 2000). Urinary tract imaging was performed after adequate hydration with the bladder full. Post-voiding images were also obtained for those subjects demonstrating kidney, ureteral, or bladder abnormalities. Images were stored for later review, either with the use of Polaroid cameras, portable thermal printers, or by image capture in electronic files.

**RESULTS AND DISCUSSION**

In *S. haematobium* infection, the adult schistosomes preferentially localize to the veins of the kidneys, ureters and bladder, although other organs of the pelvic area are often involved (Smith & Christie 1986, Poggensee et al. 1999). Egg deposition by mature female worms induces local inflammation, with granulomas, polyps, ulcers, sandy patches, and calcification of the walls of the ureters, bladder and reproductive organs (Fig. 1). Hydronephrosis frequently occurs, either because of acute inflammation localized in the ureteral wall, or as a late consequence of cumulative fibrotic injury to the ureters (Smith & Christie 1986). In studies of *S. haematobium*-related morbidity from different geographic areas, results of ultrasound studies have been generally well correlated with findings from IVP and cystoscopic examination. However, some radiologists feel that there are significant differences in the information provided by IVP versus ultrasound examination: tissue calcification and ureteral dysfunction are likely to be more apparent on dye-contrast roentgenographic studies, while hydronephrosis and deformity of poorly-function kidneys are better assessed on ultrasound examination.

Cross-sectional surveys utilizing ultrasonography in endemic villages in coastal Kenya indicate a 15-22% overall prevalence of bladder thickening and granulomata, with peak prevalence (36%) between the ages of 16 and 20 yrs, one to five years after the peak of infection intensity (Fig. 2). Hydronephrosis is found in 5% of residents, with peak prevalence of 10% among adolescents aged 12-15, with a second peak (6%) among subjects older than 49 yrs. It appears from longitudinal treatment studies that “acute” polypoid-granulomatous lesions, more common in younger individuals, are first to regress after therapy, and may disappear within weeks after treatment (King et al. 1992, Hatz et al. 1998). “Late” complications, such as hydronephrosis, may take a longer time (1-2 yrs) to show improvement, particularly in males (King et al. 1992). Elimination of infection does not terminate ongoing immune response to eggs trapped within tissues, and it is possible that fibrosis progresses for some time after successful cure of infection, and this may result in persistent dysfunction of the urinary tract.

In high transmission areas, reinfection with *S. haematobium* may occur in 1-2 yrs, with resulting recrudescence of urinary tract abnormalities detected by ultrasound (Hatz et al. 1998). This observation indicates the need for repeated antiparasitic therapy during the highest risk period of life, i.e., childhood and early adolescence, in order to minimize risk for late complications of urinary schistosomiasis. Analysis of compliance and drug efficacy of an eight year control project in coastal Kenya

![Fig. 1: probable disease causation pathway for urinary schistosomiasis due to *Schistosoma haematobium* infection. Solid lines indicate links based on experimental evidence. Dotted lines indicate associations where there is suggestive but not conclusive evidence of causation.](image1.png)

![Fig. 2: contrasting findings for age-specific infection and disease in urinary schistosomiasis in the Msambweni area of Coast Province, Kenya. The dark line indicates geometric mean intensity of infection prior to first treatment for the age groups indicated. Note peak intensity of infection in the 12-15 yrs age group. Shaded and open bars indicate group-specific prevalence of infection-associated morbidity in terms of hematuria, proteinuria, or structural abnormalities of the bladder or kidneys as detected on ultrasound. Although peak prevalence of hematuria and proteinuria mirrors the infection intensity profile, ultrasound abnormalities become more common only in older age groups (> 15 yrs), 1-5 yrs after the greatest period of *Schistosoma haematobium* infection intensity.](image2.png)
indicates that regular treatment of school age children can result in a >50% reduction in an individual’s cumulative experience with parasite infection and his or her exposure to egg deposition and consequent disease (Fig. 3, see also Subramanian et al. 1999).

Does treatment during childhood have prolonged effects on the risk for S. haematobium-related morbidity? When young adults are reexamined 7-13 yrs after an antischistosomal treatment campaign, they have fewer bladder abnormalities and a reduced number of cases of hydroureter when compared to age- and gender-matched controls from untreated villages (Subramanian et al. 1999). This is despite of the fact that, at the time of their follow up examinations, the previously treated subjects have about the same prevalence and intensity of S. haematobium infection as untreated controls.

There remain, however, unanswered questions about the clinical implications and prognosis of urinary schistosomiasis-related abnormalities detected on ultrasound. Are the abnormalities observed among older adults reversible with therapy? What is the association between structural urinary tract abnormalities and renal insufficiency, risk for bacterial superinfection, and cancer formation? and finally, How do sonographic findings predict disability levels for currently- or previously-infected individuals? Answers to some of these questions are expected from longitudinal studies now in progress, but more work will be needed to fully understand the natural history of treated and untreated schistosomiasis haematobia.

In summary, ultrasound examination provides a painless, quick, and reproducible measure of the structural urinary tract morbidity associated with S. haematobium infection. Its introduction permitted, for the first time, an evaluation of S. haematobium-associated disease on a population basis, and it is clear that the portable ultrasound has now become an essential tool in field studies of urinary schistosomiasis.

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REFERENCES


