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Schistosome Materials for Vaccine Development

by

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DISTRIBUTION STATEMENT A
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Temperature and food being controlled at optimal levels, the presence of rotifers in the snail colonies was the most important factor in parasite maintenance. Rotifer infestation of schistosome-infected snails was highly inimical to parasite development. With rotifer control, large numbers of schistosomes and products of host-schistosome interaction were produced on a weekly schedule. Limits of variability in production of parasites were set for the laboratory conditions.
BACKGROUND

The Immunoparasitology Department at the Naval Medical Research Institute (NMRI) is involved in research centered primarily on the development of effective vaccines against several parasitic diseases. Immunological research in one such disease, schistosomiasis, is particularly difficult due to the limited quantity of schistosomal materials available to most laboratories. It has been the objective of this contract to supply large quantities of schistosomal materials to investigators at NMRI and the Biomedical Research Institute (BRI) to help realize the goal for the development of an effective vaccine against schistosomiasis. The various materials provided include adult schistosomes, eggs, cercariae, schistosomules, cercarial penetration enzymes, and vertebrate and invertebrate host serum and tissues.
METHODOLOGY

A Puerto Rican strain of *Schistosoma mansoni* was maintained in *Biomphalaria glabrata* snails and Swiss albino mice. Six 20-gallon aquaria, supporting approximately 1000 snails each, provided uninfected snails. Approximately 300 snails (5-7mm dia.) were collected each week and exposed individually with 6-8 miracidia. When requested, snails were exposed to 1 miracididium each for development of single-sex schistosomal infections. Miracidia were derived from livers of 8-week infected mice. A constant supply of approximately 1000 infected snails were maintained for production of cercariae.

From these infected snails, from 1 to 3 million cercariae were collected 4 days per week and processed as needed or used for experimental work.

For the production of adult parasites and eggs two hundred and fifty Swiss mice were exposed percutaneously weekly to 250-270 cercariae each. Adult worms were perfused from these mice at 7 weeks post-infection, and eggs isolated from the livers. Schistosomules, the postpenetration stage of cercariae, were recovered from ear skin of mice exposed *in vivo* to cercariae. Schistosomules were also collected after cercarial penetration of dried rat abdominal epidermis, or by the Colley syringe shear technique.

Fifty to 100 ml of secreted enzyme solution were harvested two to four days each week from cercariae stimulated to secrete in a temperature gradient over skin surface lipid or the active fraction, linolenic acid. Total protein (Lowry method) and enzyme activity against azocoll (dye-coupled collagen) were established spectrophotometrically for each collection.
RESULTS AND DISCUSSION

The list of investigators at NMRI and BRI for whom schistosomal materials were supplied by the contract is given in Appendix I. Each week the materials supplied consisted of: 8 to 12 million cercariae; 20 to 25 thousand adult worms; 600,000 to 900,000 schistosomules, the numbers adjusted to demand; frozen irradiated schistosomules; about 200ml of preacetabular gland enzyme secretion; and excretions and secretions of cercariae as requested.

Maintenance of this large-scale life cycle of *S. mansoni* allowed investigations into the nature of the observed natural fluctuations encountered in cercarial harvests and infectivity. We noticed that cercarial production decreased during infestation of the snail colony with rotifers of the genera *Philodina* and *Rotifer*. A dramatic decrease in cercarial production per snail (Figure 1) was noted, as well as an inhibition of normal cercarial activity. The cercariae which did emerge from the affected snails were lethargic, accumulated on the bottom surface of the dish, and had tightly curled tail furci.

In summary, the provision of schistosomal materials on a large scale requires close control of the conditions known to be important for optimum parasite development. These conditions include maintaining constant temperature of infected snails, systematic feeding, reduction of rotifer infestation as much as practical, etc. Fluctuations in cercarial and adult worm production persist even when controlling these known optimal conditions. It is important therefore to continue investigating details of the schistosome life cycle in order to supply materials in quantity for vaccine development.
SIGNIFICANT ACCOMPLISHMENTS

1) Established natural fluctuations of *S. mansoni* life cycle under our conditions.

2) Compared the results of 1 miracidium vs 8-10 miracidia snail exposures in terms of:
   (a) patency of infection in snails
   (b) deaths of snails
   (c) cercarial production
   (d) cercarial infectivity

3) Demonstrated that rotifer-infestation of schistosome-infected snails reduced cercarial production.
Appendix 1.

List of Investigators Supplied with Schistosomal Materials on Contract ONR N00014-76-C-0146.

NAVAL MEDICAL RESEARCH INSTITUTE

Dr. W. E. Vannier
Dr. K. D. Murrell
Dr. D. A. Dean
Dr. A. H. Smith
Dr. P. Minard
Dr. D. W. Taylor
Dr. P. Coulis
Dr. V. Schinski
Dr. M. Stek
Dr. C. H. Dorsey
Dr. A. Attallah

BIOMEDICAL RESEARCH INSTITUTE

Dr. M. A. Stirewalt
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Dr. E. Hayunga
Dr. C. Cousin
Figure 1. Histogram comparing the average number of cercariae per snail from Rotifer (R) and Non-Rotifer (C) Biomphalaria glabrata on nine consecutive Collecting Days.
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ABSTRACTS OF PRESENTATIONS

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