First Molecular Identifications of *Necator americanus* and *Ancylostoma ceylanicum* Infecting Rural Communities in Lower Myanmar

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Abstract. Hookworms are enteric parasitic roundworms infecting an estimated 400 million persons worldwide. Herein, we provide the first molecular identifications of human hookworms from certain parts of rural Lower Myanmar. DNA was extracted from hookworm-positive stool samples, as determined by microscopy. DNA sequences of the partial internal transcribed spacer 1, full length 5.8S gene, and partial internal transcribed spacer 2 were determined and compared with available hookworm sequences from public databases. Of the 11 polymerase chain reaction–positive samples, eight (Bago Region, *N* = 4; Mon State, *N* = 4) yielded sequences with high similarity to those of *Necator americanus*. A further three sequences (Mon State, *N* = 2; Bago Region, *N* = 1) showed high similarity with those of *Ancylostoma ceylanicum*. The latter is primarily a parasite of dogs and represents a zoonosis. Given that different species of hookworms exhibit different epidemiological and biological characteristics, accurate identification is essential for the planning and execution of effective control programs for hookworm infections.

Human hookworm infection remains a main public health problem, particularly in developing countries in tropical and subtropical regions due to poor sanitation and hygiene practices.3 An estimated 438.9 million people are infected with hookworms globally, experiencing chronic blood loss and anemia, and often leading to retardation of physical and cognitive growth in children.2 Of the two major hookworm species that cause human infection, *Necator americanus* is worldwide in distribution with the highest infection rates in sub-Saharan Africa, the tropical sections of the Americas, south China, and southeast Asia. *Ancylostoma duodenale* is endemic in certain parts of India, China, sub-Saharan Africa, north Africa, and a few areas of the Americas.3 In recent years, zoonotic hookworm species such as *Ancylostoma ceylanicum* and *Ancylostoma caninum* have been reported as significant health problems in many areas.4,5 *Ancylostoma ceylanicum* is now the second most common hookworm species infecting humans in Asia and possible cause of cutaneous larvae migrans or abdominal symptoms in infected patients, with reported high egg-positivity rates in Thailand, Laos, Cambodia, and Malaysia.6–10 National surveys on the prevalence of soil-transmitted helminths (STHs) in Myanmar reported a reduction in hookworm prevalence in schoolchildren from 6.5% to 0.3% 7 years after a national deworming program.11,12 A diagnosis of hookworm infection was usually made by microscopic examination of eggs or larvae in the fecal samples. However, species identification of hookworms is difficult and cannot be done using eggs or larvae. Each genus and species of hookworm infecting humans has its own distinct biology, life cycle, pathophysiology, and epidemiology, knowledge of which therefore impacts on designing control measures.6 A molecular approach to identification is therefore imperative and copro-molecular methods for species identification of hookworms have been reported in neighboring countries (Thailand,8,13 Lao People's Democratic Republic [PDR],14 and Cambodia6,15). Molecular methods were used to identify as *A. ceylanicum* a case of hookworm infection in a French individual who had visited Myanmar. This is the only previous report on molecular characterization of hookworm species in Myanmar.16 In the present study, molecular identification of hookworms that infect humans in rural parts of Lower Myanmar is reported for the first time. This genetic data is of value in continuing explorations into epidemiology of hookworms in Myanmar populations.

This study protocol was approved by the Khon Kaen University Ethics Committee for Human Research (HE581396) and the Research and Ethical Committee of the University of Medicine 1, Yangon, Myanmar (5695/Research and Ethics 2015). Each participant was informed of study methods, risks, and benefits of the process. Before enrollment, written consent was obtained from all adult participants and from parents or legal guardians of minors. All infected persons were treated according to the World Health Organization guideline for STH control programs in Myanmar.17

A total of 364 fecal samples was collected from individuals between 5 and 60 years of age living in three different rural study areas of Lower Myanmar (Kyaik Hto Township, Mon State, *N* = 100; Shwe Kyin Township, Bago Region, *N* = 127; Thone Gwa Township, Yangon Region, *N* = 137) (Figure 1) during June and December 2015. Twenty-one samples (5.8%) were positive for hookworm eggs after stool examination by formalin ethyl acetate concentration technique. Of these, 11 (3.0%) were from the Bago Region, nine (2.5%) from Mon State, and one (0.3%) from the Yangon Region. DNA was extracted from all positive fecal samples (preserved in 95% ethanol and stored at −20°C) using a QIAamp® DNA stool mini kit (Qiagen, Hilden, Germany), according to the manufacturer’s instructions. The polymerase chain reaction (PCR) was carried out using a GeneAmp® PCR System 9700 (Applied Biosystems, Singapore City, Singapore). Primers used were specific for *Ancylostoma* as well as *Necator* species and amplified most of the ribosomal internal spacer region (partial internal transcribed spacer 1, 5.8S rRNA gene, and partial internal transcribed spacer 2). Primer sequences

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are RTHW1F (forward): 5′-GAT GAG CAT TGC WTG AAT GCC G-G-3′ and RTHW1R (reverse): 5′-GCA AGT RCC GTT CGA CAA ACA G-3′. The PCR reaction and conditions were as reported previously. PCR products were run on a 1% agarose gel to demonstrate amplicons of approximately 485 base pairs (bp) (typical for N. americanus) or 380 bp (typical for Ancylostoma spp.). Eleven samples yielded positive PCR results. DNA sequencing of these was performed using an Applied Biosystems 3730 + DNA Analyzer and ABI big dye Version 3.1 (Applied Biosystems, Foster City, CA). Sequencing was done in both directions, using the PCR primers as sequencing primers. The nucleotide sequences obtained were compared with those available in the GenBank database by BLAST-N search via NCBI. Sequences were obtained were compared with those available in the GenBank database by BLAST-N search via NCBI. Sequences were obtained whereFig. 1. Study areas in Lower Myanmar.

Prevalences of hookworms in Cambodia are reportedly very high,18 and are predominantly due to N. americanus and A. ceylanicum (51.6% equal prevalence); only 3.2% of people were found infected with A. duodenale.5 In Thailand, the principal hookworm species infecting humans was confirmed as N. americanus. However, A. duodenale, A. ceylanicum, and A. caninum were also detected in different parts of the country.8,13 The predominance of N. americanus in Lower Myanmar is similar to a report from Peninsular Malaysia where N. americanus was more common than A. ceylanicum, but A. duodenale infection was not discovered.13 In contrast, a report from Lao PDR people found that infections due to A. duodenale and the animal hookworms, A. caninum and A. ceylanicum, were slightly more prevalent than those due to N. americanus.14 Hookworm infections constitute a significant neglected tropical disease in the human populations of southeast Asia.2 Our study has been the first to use molecular techniques to identify the species of hookworms, including the zoonotic A. ceylanicum, infecting rural people in Lower Myanmar. Although animal hookworm infections of people have been regarded as rare and overlooked, A. ceylanicum has been reported from a French traveler who had visited Myanmar and presented with bloody diarrhea.16 The present study also identified A. ceylanicum from asymptomatic rural people from the study area in Myanmar. Zoonotic awareness should be a concern because domestic animals such as dogs and cats act as natural hosts.19 The geographical differences in species of hookworm causing infections in humans are probably related to climate, temperature, environmental factors, parasite behavior, and ethnicity.5 The extent of variation within the cox1 sequence of human hookworm still needs further investigation as reported previously16 to explore potential haplotype-linked differences in zoonotic, epidemiological, and pathological characteristics of human hookworm in different localities of Myanmar.

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REFERENCES