Establishment of Biomphalaria tenagophila Snails in South Africa

To the Editor:

Schistosomiasis, known since ancient times, is caused by blood flukes (Trematoda: Schistosomatidae). It is a major communicable disease with public health and socioeconomic effects in the developing world (1). Among parasitic diseases, schistosomiasis ranks second only to malaria with regard to the number of persons infected and at risk. The life cycle of schistosomes is complex, requiring specific freshwater snails as intermediate hosts for larval development and multiplication. Among Schistosoma species that affect humans, Schistosoma mansoni is the most likely to invade new areas mainly because of the adaptability and invasiveness of its intermediate host, Biomphalaria snails. Natural populations of these snails are usually found in tropical standing water or freshwater in South America and Africa, but they also reach 30° latitude in subtropical areas (1,2). Many species of these red-blooded planorbid snails (Gastropoda: Planorbidae) are able to survive a long time when removed from their freshwater habitat (1). Of the 34 Biomphalaria species, 4 (B. glabrata, B. pfeifferi, B. straminea, and B. tenagophila) have recently expanded their native ranges (3). They have been introduced to areas where other Biomphalaria species are endemic (e.g., Congo and Egypt) or to subtropical zones that have no frost period (Texas, Louisiana, Florida, Hong Kong) (3,4). None of the known invasions, whether peripheral range expansion or long distance dispersal, reached the temperate zone. Spreading of the blood-fluke snails to schistosome-free areas may enable the parasite to colonize new habitats concurrently, expanding the potential area of clinical schistosomiasis.

We collected these snails in spring 2005, autumn 2006, and autumn 2007, near Răbăgâni, Romania, Eastern Europe (46°45′1.3″N, 22°12′44.8″E) in a hypothermal spring. Water temperature was 25°C in the spring and 16°C–25°C, gradually decreasing, along the brook course. In and beside an abandoned concrete pool next to the spring, we collected 100 shells and 34 living specimens that macroscopically resembled Biomphalaria spp. snails. All 16 dissected animals proved to be fully developed adults, according to the maturity of their genital organs (Figure). Using available identification keys (5), we tentatively identified these snails as B. tenagophila. Voucher specimens have been deposited in the Hungarian Natural History Museum (accession nos. HNHM96857 and HNHM95433).

DNA was extracted from the foot muscles of 3 specimens by using QIAamp DNA Mini Kit (QIAGEN, Hilden, Germany). For amplification of the partial mitochondrial 16S ribosomal RNA gene, we used a PCR with primers 16Sar and 16Sbr (6). Nucleotide sequences were determined in both directions. PCR products of ≈430 bp were detected from all 3 samples. Automatic cycle sequencing of the randomly selected amplicon (GenBank accession no. EU069412) showed 99.74% similarity to B. tenagophila (AF449615, Brazil).
Our morphologic, anatomic, and molecular data unambiguously prove the occurrence of *B. tenagophila* snails in Romania. *B. tenagophila* snails had been found earlier (in 2004) at this location but had presumably been misidentified as dwarf specimens of a common European species, *Planorbarius corneus* (7). Consequently, *B. tenagophila* snails have been not only introduced, but also established in Râbâganí, representing the furthest self-sustaining population of this species from the equator.

*B. tenagophila* is a new species for the European fauna. It could represent a founder population of unknown origin for further spread into Europe, which might easily be accomplished by migrating birds or more likely by plants used in aquariums (3). Although no trematode larvae were detected in the observed specimens, clinical schistosomiasis can be imported by immigrants or tourists into Europe, as has been reported in Romania and neighboring Hungary (8, 9). If eggs were released in feces of humans infected with the blood flukes, they could hatch in the environment and the larvae could develop to an infective stage in these snails. The observed local cultural and social factors involving natural water (washing clothes, bathing) in Râbâganí where *B. tenagophila* have been found may also increase the chance of human infection.

We believe that *B. tenagophila* in Europe, together with the global climate change and a possible encounter of these snails with schistosomes, could pose a public health risk. Measures must be taken to prevent the spread of this species into European freshwater. Chemical control is not possible in Râbâganí because it is an area where other rare and endangered snail species are protected (7). Therefore, the manual collection and removal of all the *B. tenagophila* specimens in the area seems to be the only possibility for eradication, which might remain in effect for years. To avoid similar establishments, we suggest regular malacologic and parasitologic surveillance of at least the thermal and hypothermal water bodies for these tropical invaders around European settlements.

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**References**


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LETTERS

Rickettsia aeschlimannii Infection, Algeria

To the Editor: Only 2 cases of Rickettsia aeschlimannii infection have been reported. We report 2 additional cases documented in Algeria by immunofluorescence (IF) assays and confirmed by Western blot (WB) assays and cross-adsorption studies.

Tick-borne rickettsioses are now recognized as emerging or reemerging human infections worldwide. These zoonoses, caused by intracellular bacteria within spotted fever group (SFG) Rickettsia spp., share characteristic clinical features including fever, rash, and sometimes inoculation eschar at the bite site (1). In North Africa, cases of rickettsioses are rarely documented (2). In Algeria, only Mediterranean spotted fever caused by R. conorii has been described (3).

From 2000 through 2006 in Algeria, all patients with suspected rickettsioses seen at the infectious diseases units of Constantine and Batna hospitals were included in a prospective study; clinical and epidemiologic data and acute-and convalescent-phase serum samples obtained 2–4 weeks later were collected. Serum samples were sent to Marseille, France, where they were analyzed by an IF assay, using 9 SFG rickettsial antigens (R. conorii conorii, R. conorii israelensis, R. africaca, R. sibirica mongolitimonae, R. aeschlimannii, R. massiliae, R. helvetica, R. slovaca, and R. felis) and a typhus group antigen (R. typhi) (3). The IF assay result was considered positive 1) if immunoglobulin ( Ig) G titers were ≥128 and/or IgM titers were ≥64 for R. conorii and 2) if IgG titers were ≥64 and/or IgM titers were ≥32 for other rickettsial antigens (3). When cross-reactions between several antigens were noted, rickettsial antigen was considered to represent the infectious agent if titers of IgG and/or IgM antibody against this antigen were at least 2-fold higher than titers of IgG and/or IgM antibody against other rickettsial antigens (3,4). When the difference in titers among several antigens was lower than 2-fold, WB assays and cross-adsorption studies were performed (4,5). A total of 135 patients were included in the study. We describe 2 cases of R. aeschlimannii infection. Cases caused by other SFG rickettsiae will be reported elsewhere.

An 80-year-old man who reported contact with dogs parasitized by ticks had a 7-day history of high fever, headache, myalgia, and vomiting. On physical examination, a generalized maculopapular rash, 2 eschars (right shoulder and knee), and bilateral hemorrhagic signs on the retina were noticed. Elevated levels of liver enzymes (aspartate aminotransferase 187 U/L, alanine aminotransferase 108 U/L), hyponatremia (sodium 120 mmol/L), and hypokalemia (potassium 2.9 mmol/L) were found. IF assay showed raised levels of IgG/IgM against R. aeschlimannii (512/64) and R. conorii (128/0).

The second patient, a 36-year-old man, reported a 15-day history of fever with headache and failure of amoxicillin and cotrimoxazole treatments. Oral aphthous, a maculopapular rash, and purpuric lesions on the arms were noticed. IF assay showed raised levels of IgG/IgM at the same titer (2,048/32) against R. conorii, R. aeschlimannii, and R. massiliae. WB assays and cross-adsorption studies confirmed that antibodies were directed against R. aeschlimannii (Figure). Both patients recovered after doxycycline treatment (1).

R. aeschlimannii was first characterized as a new SFG rickettsia after its isolation from Hyalomma marginatum marginatum ticks in Morocco in 1997 (6). Thereafter, R. aeschlimannii has been detected in this tick species in southern Europe and North Africa (7), as well as in H. m. rufipes in sub-Saharan Africa (1). Preliminary data have suggested that these Hyalomma

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Letters

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