Gnathostomiasis: Report of a case and brief review

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S HOUSTON. Gnathostomiasis: Report of a case and brief review. Can J Infect Dis 1994;5(3):125-129. Nematodes of the genus Gnathostoma can cause a spectrum of disease in humans. This includes a distinctive syndrome of intermittent migratory subcutaneous swellings, central nervous system involvement with high morbidity and mortality and occasionally, involvement of other organs. Gnathostomiasis is endemic in southern and eastern Asia, particularly Thailand, but has recently been reported from Ecuador and Mexico. Diagnosis is usually based on clinical, epidemiological and serological findings. A recent study suggests that treatment with albendazole is effective. This report describes a patient diagnosed in Canada in whom evidence of pericarditis was associated with gnathostomiasis and in whom treatment with albendazole appears to have been effective.

Key Words: Albendazole, Drug therapy, Eosinophilic meningitis, Gnathostoma spinigerum, Gnathostomiasis, Pericarditis

La gnathostomiase : Étude de cas et revue brève


Gnathostoma spinigerum was first identified in the gastric tumour of a tiger in 1836 (1). Human gnathostomiasis was first reported from Thailand in 1890 in a woman presenting with a breast mass (2). The disease is being recognized with increasing frequency due to better appreciation of the clinical features and improved diagnostic techniques. It is sufficiently common in endemic regions that occasional cases can be expected in North America, yet North American physicians are often not familiar with the disorder. Two recent developments make awareness of this infection more important: the recognition of endemic foci in the western hemisphere and evidence that medical therapy is effective.

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This report describes a patient whose gnathostomiasis was diagnosed in a very nontropical Canadian city and who had clinical evidence of pericardial involvement which has been reported only once previously.

**CASE PRESENTATION**

A 31-year-old woman of Thai origin was referred because of eosinophilia and recurrent superficial swellings.

She first noticed the swelling about six years previously and described almost identical recurrences approximately once a year since then. Each episode was characterized by the rapid appearance of a mildly painful swelling over the right lower forearm lasting about two weeks. Episodes were not accompanied by fever or other systemic symptoms nor by swelling or pruritis at any other site. She specifically denied any history of serious neurological disease or chest pain. She was entirely well between episodes. She had presented with similar symptoms to a physician in Canada two years previously. A peripheral eosinophilia was present and strongyloides larvae were found on stool examination at that time, but the eosinophilia resolved after treatment with thiabendazole. Her past history was otherwise negative.

The patient had spent most of her life in a small village in northern Thailand. Although now married to a Canadian, she continues to spend about six months each year in Thailand. She admitted eating a variety of local foods in Thailand including uncooked fish.

On examination she was afebrile and looked well. There was a subcutaneous swelling 4 to 5 cm in diameter with poorly defined margins over the distal right forearm. It was mildly tender but neither warm nor red. The only other abnormality on physical examination was a loud pericardial rub. This was documented by two physicians at the first visit and was noted at a subsequent visit two weeks later in spite of having been found to be absent on one examination in the interval. The cardiovascular system was otherwise completely normal with no evidence of tamponade.

The absolute eosinophil count was 5.9x10^9/L. Chest x-ray and electrocardiogram were normal; echocardiography revealed a trace of pericardial fluid. Western blot examination of serum using a somatic extract of Strongyloides larvae was found on stool examination at that time, but the eosinophilia resolved after treatment with thiabendazole. Her past history was otherwise negative.

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The absolute eosinophil count was 5.9x10^9/L. Chest x-ray and electrocardiogram were normal; echocardiography revealed a trace of pericardial fluid. Western blot examination of serum using a somatic extract of the third stage larva of Gnathostoma spinigerum (3) supported the clinical diagnosis of gnathostomiasis.

The swelling resolved completely without specific treatment. Albendazole 400 mg twice daily was given by mouth for 14 days starting several weeks after resolution of the swelling. No adverse effects were associated with the treatment. The pericardial rub was absent on each of several examinations performed more than two weeks after initial presentation. No recurrent swelling had been noted when the patient was contacted 30 months after completion of therapy and the absolute eosinophil count at 21 months was 0.2x10^9/L.

**DISCUSSION**

Human gnathostomiasis is usually caused by the nematode G. spinigerum. Cases of human disease due to Gnathostoma hispidum (4), Gnathostoma nipponicum (5) and Gnathostoma doloresi (6) have been described.

**Life cycle (Figure 1):** First-stage larvae in water are ingested by a tiny crustacean, cyclops. Second-stage larvae in infected cyclops are in turn ingested by a second intermediate host, frequently one of several species of fresh water fish. The parasite penetrates the gastric wall and the third-stage larva encysts in the muscle of the second intermediate host. 'Secondary' infection with third-stage larvae, of another second intermediate host, can occur if it ingests the 'primary' second intermediate host. This has been demonstrated experimentally in at least 33 animals including crustaceans, fish, amphibians, reptiles, birds and mammals, and appears to occur commonly in nature (7). The infection is productive only in the definitive hosts - dogs, cats and some wild animals - where larvae mature in the wall of the stomach, mate there and release eggs into the feces. The eggs hatch in water producing first-stage larvae.

The main source of human infection is thought to be ingestion of inadequately cooked flesh of either the 'primary' or 'secondary' second intermediate hosts harbouring third-stage larvae. Fresh water fish and domestic poultry may be the most important sources (8). Based on animal models, it is thought that ingestion of
water containing infected cyclops may be an occasional mechanism for human infection (9). Experimentally, the parasite can penetrate the intact skin of cats, indicating that handling the flesh of infected fish could also be a source of human infection (10). Three cases of apparent perinatal transmission have been reported (11,12). Human infection involves third-stage larvae of either sex and of varying degrees of maturity, ranging from 2.2 to 16.25 mm in length (11).

**Geographic distribution:** Most cases of human gnathostomiasis have been reported from Asia, particularly Thailand but including China, Japan and several countries of southeast Asia from Bangladesh to the Philippines. One case was reported in a woman who had lived only in Europe and Israel (13). Recently there has been a number of reports of human gnathostomiasis from Latin America (14-16) and infested fresh water fish have been documented there (17).

**Clinical features:** It is thought that symptoms and organ damage in gnathostomiasis are due to mechanical tissue injury produced by this macroscopic nematode in its migrations.

The incubation period between ingestion of gnathostoma larvae and the first development of cutaneous symptoms is three to four weeks (7). Abdominal pain and vomiting occurred within 24 to 48 h, followed by pulmonary, then cutaneous symptoms in a common source outbreak involving nine individuals who consumed raw fish at a party (18).

The clinical features of reported cases of gnathostomiasis are documented in two comprehensive reviews (9,19). The most common presentation is that of recurrent subcutaneous swellings which are usually painless but sometimes associated with pain sufficient to result in confusion with bacterial cellulitis. These swellings reflect the migration of the parasite. They may last one to two weeks and recur at intervals of weeks to months at the same or different sites. Systemic symptoms are not usually noted. A peripheral eosinophilia is usual but not invariable. A superficial creeping eruption has also been described (20). There is a tendency for recurrences to become less frequent and less severe over time but they may persist for over 10 years indicating that the parasite may live that long in the human host. A syndrome of migratory subcutaneous nodules may also be seen in paragonimiasis infections (62).

The most serious manifestation of gnathostomiasis is central nervous system (CNS) involvement, first proven only in 1967 (21) although it had been previously suspected (22,23). Recognized presentations include encephalitis, painful radiculitis, myelitis, cranial nerve palsies, subarachnoid hemorrhage (24) and eosinophilic meningitis or combinations of these. Gnathostoma, unlike angiostrongylus, does not seem to be neurotropic but to involve the CNS only by chance. Brain involvement and subarachnoid hemorrhage have also been described in infections with *Paragonimus westermani var szechuanensis* (62). CNS gnathostomiasis may occur in any age group. Only 7% of 162 patients in one report had a history of intermittent migratory cutaneous swellings while 4% had unilateral eyelid swelling (25).

Peripheral eosinophilia is usual in CNS gnathostomiasis - an eosinophilia of greater than 10% was found in 57% in one series (25). Cerebrospinal fluid (CSF) pleocytosis with at least 10% eosinophils is present at some point in the course of nearly all patients with CNS gnathostomiasis (25,26). The other important cause of eosinophilic meningitis, *Angiostrongylus cantonensis*, is also endemic in parts of southeast Asia but the clinical syndrome is usually one of meningeval involvement only and is self-limiting (27). Among 162 patients with CNS gnathostomiasis, bloody or xanthochromic CSF was found in 64%, CSF pressure was elevated in more than half, CSF protein exceeded 50 mg/100 mL in 72% but CSF glucose was depressed in only 10% (25).

Case fatality was at least 12% among 162 patients with probable CNS gnathostomiasis in the largest series (25). The authors believed the true case fatality rate was considerably higher since 38 patients were taken home unimproved, many critically ill. Sixty-five per cent were considered ‘improved’, many with persisting neurological deficits. Autopsies of patients with CNS gnathostomiasis have revealed grossly visible worm tracks with accompanying necrosis and hemorrhage in the brain or spinal cord (28).

Gnathostomiasis has also been reported to involve the eye (13,29-41), the urinary (42,43) and genital tracts (44,45), the ear (46,47), the abdomen (48,49), the breast (2,50), and the lung or pleura (51-55). Ocular involvement seems to be the least uncommon of these; even with prompt surgical removal of the worm, permanent injury is common.

There is one report in the literature of a patient with a pericardial effusion, documented by echocardiogram, associated with gnathostomiasis (56). The patient, of Nepalese origin, also had a pleural effusion, peripheral eosinophilia and intermittent migratory swellings over the chest wall from one of which a gnathostome was removed. Following removal of the worm (and treatment with diethylcarbamazine) the patient experienced prompt clinical and radiological improvement without relapse.

**Diagnosis:** The diagnosis of gnathostomiasis is usually made in the presence of a distinctive clinical syndrome, most often intermittent migratory swellings or compatible CNS disease, in a person from an endemic area. Retrieval and identification of the worm confirm the diagnosis with certainty when this is possible. Even when the worm is missed, histology of the panniculitis lesions is said by one author to be characteristic (14). The presence of a marked peripheral blood eosinophilia is typical but not invariable (57). Skin testing for gnathostomiasis has been used, but recent develop-
ments in serological testing appear to be very promising (3,58).

**Treatment:** Until recently the only treatment known to be effective was removal of the worm. Corticosteroids may speed resolution of cutaneous swellings (56) but have not been proven to have any benefit in the most serious form of the disease – in the CNS (25). Ivermectin and albendazole appear to have a therapeutic effect in animal models (59,60). One recent report describes the elimination of episodes of intermittent migratory swelling at six months follow-up in 89 of 100 patients treated orally with either 400 mg daily or 400 mg bid of albendazole for 21 days (61), while recurrences were noted in all 12 placebo recipients. Eosinophilia resolved in more than 90% of those treated but in only one of 12 placebo recipients, and serological titres declined in the treatment but not the placebo group. Interestingly, treatment appeared to provoke migration of the larvae to the skin where 22 were removed from 21 patients. It is to be hoped that longer follow-up and further experience will confirm these very encouraging results. There are no reports of albendazole treatment of CNS disease.

In the present patient the typical history and appearance of intermittent migratory swellings, the epidemiological history, and the positive serological test clearly indicate a diagnosis of gnathostomiasis. A pericardial rub is an objective clinical finding which is rarely if ever ‘incidental’. The striking temporal association of a pericardial rub with an episode of intermittent migratory swelling, and the absence of an alternative cause, strongly suggest that it was related to gnathostomiasis.

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**REFERENCES**


