**Echinococcus multilocularis** – first recorded case of Norway rat (*Rattus norvegicus*) in Poland

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Abstract

**Introduction.** *Echinococcus multilocularis* is a very dangerous zoonotic parasite threatening human health. The red fox is the main definitive host, and cats and dogs less commonly. Rats can be intermediate hosts.

**Objective.** The aim of the study was to determine the parasitofauna of Norway rats and some cats and dogs living on a farm near a forest.

**Materials and method.** A parasitological section on 15 Norway rats was conducted. The internal organs were examined by means of macroscopic and microscopic methods. For molecular examination, a QIAmp DNA Mini Kit (Qiagen) was used.

**Results.** Based on necropsy, parasitological and molecular examinations, of the 15 examined rats, 1 was found to have larvae of *E. multilocularis*, while 3 others had eggs of *Hymenolepis diminuta*, *H. nana* and *Syphacia obvelata*. The faeces of the pets did not contain any developmental forms of parasites.

**Conclusions.** This is the first case of *Echinococcus multilocularis* infestation in a rat in Poland.

**Key words**

*Echinococcus multilocularis*, rodent, *Rattus norvegicus*, intermediate host, Poland

INTRODUCTION

*Echinococcus multilocularis* is one of the most dangerous zoonotic parasites that pose a threat to human health [1, 2, 3]. The literature abounds in studies devoted to this invasion in animals in different areas of the world and the resultant threat to human health. In Poland, 121 cases of alveococcosis in humans have been described, of which 23 (19%) died [1].

*E. multilocularis* is an armed tapeworm belonging to the *Taenidae* family. It reaches 2–4 mm in length and its strobila consists of only 4–5 segments. It undergoes an indirect life cycle, and the definitive host is predominantly the red fox (*Vulpes vulpes*) [2]. The literature indicates that other animals are also definitive hosts, such as the raccoon dog [4, 5, 6], Arctic fox [7], golden jackal [8], and the wolf [9, 4] as well as two domestic species, dogs and cats [10, 11, 12]. Intermediate hosts are found mainly among rodents, such as muskrats, arvicolid voles, in which the larval form develops in the liver. Less frequently, *E. multilocularis* are found in murids and coypus [2]. In contrast, pigs and horses, similarly to humans, are not typical intermediate hosts, which rarely develop the invasive form [12, 14, 15]. Intermediate hosts, including humans acting as aberrant intermediate hosts, are infected through the ingestion of eggs excreted in the faeces of definitive hosts.

Studies on the prevalence of *E. multilocularis* in foxes have been conducted in Poland, the same as in other European countries, yielding a mean prevalence at 16.5%, with significant variation in different regions of the country (which in the case of the north-east and south-east reached 50%). In the area of the Lublin Province, where the studies presented in the article were carried out, the prevalence of foxes was 18.2 [16]. In Poland, an invasion by *E. multilocularis* was found in 2 dogs, which was confirmed by molecular examinations [10]. However, no tapeworms have been reported in cats. No studies into the presence of its larval stages in rodents have been performed. Earlier research by Malczewski et al. [17] did not yield positive outcomes.

**OBJECTIVE**

The aim of the study was to determine the prevalence of parasitofauna in the Norway rat (*Rattus norvegicus*), with particular regard to its role as an intermediate host for the *E. multilocularis* tapeworm, and also to determine the prevalence of parasitofauna in companion animals (cats and dogs) living on the farm where the examined rats were obtained.

**MATERIALS AND METHOD**

The research covered 15 Norway rats collected during routine rat extermination. All of them originated from one farm in the Lublin Province, located about 1 km from forest areas. Additionally, red foxes were very often seen on this farm.

A parasitological dissection of the rodents was conducted. The internal organs (liver, lungs, heart, kidney and gastrointestinal tract) were examined by means of parasitological macroscopic and microscopic methods. The macroscopic examination involved a visual inspection and
incision of the internal organs. The microscopic examinations of intestinal content were carried out using a flotation technique involving a saturated solution of salt and sucrose (specific gravity 1.28–1.30), and a decantation method to evaluate qualitatively the composition of parasitofauna [18]. Particular species were identified morphologically with cell light microscope system software (Olympus) [19]. The mucous membrane and scraped fragments of the bowels were inspected using a biological microscope.

For identification, the isolated parasites were immersed in 70% ethanol with 5% glycerin [20, 21]. The tissues and internal organs in which lesions had been identified were placed into 70% ethanol for molecular examination. The diaphragm muscles and other striated muscles were examined using the digestion method in order to detect Trichinella spiralis larvae [18].

For molecular examination, the DNA of the tissues with lesions was isolated using a QIAmp DNA Mini Kit (Qiagen) in accordance with the manufacturer’s instructions. A nested PCR method was used, as described by Dinkel et al. [22], with some modifications concerning the reaction mixture and time conditions of amplification [14]. The sequence for amplification was a section of the E. multilocularis mitochondrial 12S rRNA gene. The fragment specific to E. multilocularis (250 bp) was amplified. The E. multilocularis positive products of nested PCR were sequenced. The samples for sequencing were purified in Sephadex G-50 columns. Sequencing was performed using a BigDyeTM Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems) on an ABI3730xl Genetic Analyzer (Applied Biosystems). The sequenced data were analyzed and compared in the GenBank using BLAST searchers.

Additionally, the parasitological examination covered faeces from 2 dogs and 7 cats living on the farm. These animals live mainly outdoors and can freely access the area surrounding the farm. Their faeces was examined 3 times at 24-hour intervals using a flotation technique involving a saturated solution of salt and sucrose (specific gravity 1.28–1.30) and a decantation method to evaluate qualitatively the composition of parasitofauna [18]. Particular species were identified morphologically with cell light microscope system software (Olympus) [23].

RESULTS

In the liver of 1 rat, the macroscopic examination revealed cystic, soft, 2–4 mm large follicles, attached to another and filled with fluid. The lesions were extensive and encompassed over half the liver (Fig. 1,2). The whole isolated liver was placed in 70% alcohol and sent to the State Veterinary Institute in Pulawy, Poland, for further molecular examination, which found the follicles to be E. multilocularis larvae. The specific PCR product (250 bp) was found in nested PCR. A comparison of the sequencing results of the obtained amplicons with the GenBank database confirmed that the causative agent, in all 3 cases, was E. multilocularis (sequences corresponding to E. multilocularis mitochondrial 12S rRNA gene).

Microscopic examination of the intestinal content in 3 other rats, employing both flotation and decantation, revealed the presence of parasitic eggs. In 2 rats, eggs of the Hymenolepis diminuta tapeworm were detected, and were single invasions (with 1 species). In contrast, a mixed invasion was detected in 1 rat, with eggs of H. nana and eggs of the Syphacia obvelata nematode. For the rat which was found to have E. multilocularis larvae; microscopic examination was negative. No Trichinella spp. larvae were found in the muscles of the examined rats.

The faeces obtained from the companion animals (cats and dogs) did not contain any developmental forms of parasites.

DISCUSSION

Rodents, including rats, are often subject to other pathogens while living in various environments (silvatic or synanthropic), including those parasites which threaten human health. One of them is E. multilocularis, with humans serving as aberrant, non-specific intermediate hosts. In this role, humans are infected through ingestion of eggs excreted in the faeces of a definitive host [2]. As definitive hosts, carnivores living in the wild (mainly red fox) and domesticated animals, such as cats and dogs, become infested by ingesting intermediate hosts. Various authors have confirmed the presence of the larval forms in small rodents, e.g. the common vole (Microtus arvalis), water vole (Arvicola amphibius), bank vole (Myodes glareolus), field vole (Microtus agrestis), and various Apodemus species [24]. However, there are few reports concerning Norway rats as intermediate hosts for E. multilocularis. In Europe, the larvae of this tapeworm were found in Rattus norvegicus in Romania [25]. Outside Europe, its presence has been demonstrated in rat liver in Japan. Of the 42 examined animals, only 1 exhibited protozoalceces in large liver cysts [26].

CONCLUSIONS

The discovery of the larval forms of the E. multilocularis tapeworm in a rat is the first recorded case in Poland. This proves that rats can indeed be an important link in the spread of this invasion on farms and in the immediate vicinity of humans. Companion animals (cats, dogs) which hunt and reside on farms can become infested, and consequently pose a significant danger to human health. Although the
result of the parasitological examination of the companion animals was negative, the risk of them becoming infested cannot be excluded. It should be remembered that these are hunting animals which live mainly outdoors and are in contact with the natural environment surrounding the farms. The detection of the larval form in a rat confirms the fact that foxes living in the vicinity of the farm are infested. The farm has been monitored and the companion animals periodically examined for the presence of parasites.

REFERENCES


