Commentary

Population dynamics of *Taenia polyacantha* metacestodes in the bank vole *Clethrionomys glareolus*

Voitto Haukisalmi & Heikki Henttonen

*Haukisalmi, V., Department of Zoology, Division of Ecology, P.O. Box 17, FIN-00014 University of Helsinki, Finland*

*Henttonen, H., Department of Forest Ecology, Finnish Forest Research Institute, P.O. Box 18, FIN-01301, Vantaa, Finland*

Received 1 February 1993, accepted 16 February 1993

*Taenia polyacantha* is a holarctic tapeworm of canids, especially the foxes, *Vulpes* spp. The larval stage of *Taenia polyacantha* is a relatively rare parasite of rodents, the highest prevalence reported being 12% (Wiger et al. 1974). Adult tapeworms inhabit the small intestine of the final host, producing eggs to be expelled in faeces. When the intermediate rodent hosts ingest food contaminated with tapeworm eggs, embryos migrate into the body cavity and develop there into larvae, or metacestodes (Rausch 1985). A metacestode of *T. polyacantha* is readily identified by the number of rostellar hooks (52–62) and by the distinctive “tail” (Verster 1969, Wiger et al. 1974).

We have continuously collected data on metacestodes of microtine rodents at Pallasjärvi, western Finnish Lapland, since 1970. Despite the examination of thousands of voles belonging to six species (*Clethrionomys glareolus, C. rutilus, C. rufocanus, Microtus agrestis, M. oeconomus* and *Lemmus lemmus*), *T. polyacantha* remained absent until 1987, when a single infected vole was found. Thereafter it has been regularly present at Pallasjärvi. We here analyse the age and sex relations and population dynamics of *T. polyacantha* in the most abundant species of the microtine community, the bank vole *Clethrionomys glareolus*, in 1988–1992.

**Materials and study sites**

The voles (*n = 1229*) were obtained from three main study sites: 1) a trapping site on the upper part of a forested mountain slope (*n = 825*), 2) several permanent study grids at lower elevations (*n = 274*), and 3) three low elevation live trapping grids (*n = 55*), where vole densities have constantly been higher than in the surrounding areas due to the provision of supplemental food (Henttonen et al. 1987 and Henttonen unpubl.). In addition, we have included data (*n = 75*) from several other forest sites nearby. The habitat of all study sites is fairly similar, i.e. spruce-(*Picea abies*) dominated taiga forest with a thick moss layer; blueberry (*Vaccinium myrtillus*) is the dominant forb species.
Infection levels in relation to sex and functional groups of voles

The best log-linear model (Fienberg 1970) for association between vole sex (S), functional group (F), and presence/absence of metacestodes (M) was SF,MF (fit to the observed data: $G^2 = 1.52$, $df = 3$, $P = 0.68$). This model indicates that the sample sex ratio and prevalence of metacestodes depend on the functional group, but there is no association between the prevalence and sex of the vole (Table 1).

The prevalence of *Taenia polycantha* was very low in immature, subadult voles (age 1–2.5 months), but clearly higher in overwintered voles (age 8–12 months), which represent the same cohort (Table 1). Data from site 1 in 1988 show that the prevalence in overwintered voles continues to increase during the summer (Fig. 1). The mature summer-born voles (those already breeding during the summer of their birth and disappearing before the winter, age 2–3 months), show intermediate prevalence compared to that of subadult and overwintered voles. The number of metacestodes per infected host did not vary markedly between host sexes (Mann-Whitney test for overwintered voles: $U = 856.0$, $P = 0.43$) and functional groups (Table 1).

These patterns suggest that the probability of a vole’s acquiring a *T. polycantha* infection increases with increasing exposure time, i.e., host age. There is no indication, however, that infections by *T. polycantha* metacestodes would increase vole mortality, e.g., by increasing vulnerability to (fox) predation. Mortality due to heavy infections is expected to lead to a decrease in prevalence (Lester 1984) and mean abundance (Anderson & Gordon 1982) of parasites in the older age groups of hosts; these phenomena were not observed in *T. polycantha*.

Long-term population dynamics

Data for male and female voles have been combined in the following description of long-term population changes, because host sex does not affect occurrence of *T. polycantha* (Table 1).

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Table 1. Percent of bank voles *Clethrionomys glareolus* infected by *Taenia polycantha* metacestodes by sex and functional group at Pallasjärvi (1987–1992), and median number and range of metacestodes in infected hosts.

<table>
<thead>
<tr>
<th>Functional groups</th>
<th>n</th>
<th>%</th>
<th>Males median</th>
<th>range</th>
<th>n</th>
<th>%</th>
<th>Females median</th>
<th>range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Immature</td>
<td>351</td>
<td>1</td>
<td>20.0</td>
<td>20–27</td>
<td>312</td>
<td>2</td>
<td>24.0</td>
<td>5–24</td>
</tr>
<tr>
<td>Mature, summer-born</td>
<td>31</td>
<td>6</td>
<td>20.5</td>
<td>5–36</td>
<td>55</td>
<td>5</td>
<td>36.0</td>
<td>25–36</td>
</tr>
<tr>
<td>Mature, overwintered</td>
<td>479</td>
<td>11</td>
<td>22.0</td>
<td>1–121</td>
<td>359</td>
<td>13</td>
<td>21.0</td>
<td>1–60</td>
</tr>
</tbody>
</table>
sites 1 and 2, the prevalence of *T. polyacantha* remained at a low level throughout the study period (Fig. 1). The apparent infection peak in overwintered voles later in the summer of 1988 at site 1 is due to the fact that these voles were older (age 10–12 months) than the overwintered voles in the spring samples (late May, age 8–10 months). In 1989, the prevalence of overwintered voles in the extra food grids (site 3) was low, comparable to that at other sites, but thereafter their prevalence reached a high level (75–90%). The high frequency of infected bank voles in the food grids indicates that foxes foraged intensively on these high-density prey patches.

The colonization and persistence of *T. polyacantha* at Pallasjärvi suggest that vole or fox densities have changed leading to an enhancement in the transmission of cestodes. We used game inquiries from Lapland as a source for a fox (*Vulpes vulpes*) abundance index; these inquiries are made each year in March (Finnish Game and Fisheries Research Institute; for details, see Helle & Kauhala 1991). The autumnal density index of all voles was based on regular, extensive snap-trappings in diverse habitats at Pallasjärvi (Henttonen et al. 1987, Henttonen unpubl.).

During 1987–1991, the average abundance indices of voles (18.5) and foxes (2.29) were higher than during 1970–1986 (15.8 and 1.86, respectively), and the variability of vole populations was lower than during the three earlier five-year periods (Fig. 2). Vole dynamics at Pallasjärvi, and more generally in Lapland, have changed since the mid-eighties, resulting in more stable dynamics in forest rodents (Henttonen
1987, Henttonen et al. 1987). The increase in density and stability of host populations has evidently been the key factor promoting the colonization of *T. polyacantha* at Pallasjärvi. This empirical observation supports the prediction of the theory that efficient transmission and persistence of parasites is possible only if the host populations exceed a certain threshold density (Anderson 1982).

References


