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TEMPORAL OCCURRENCE OF THIRD-STAGE LARVAE OF ELAEOPHORA SCHNEIDERI IN TABANUS LINEOLA HINELLUS ON SOUTH ISLAND, SOUTH CAROLINA

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INTRODUCTION

Records of intermediate hosts of the arterial worm in the southeastern United States are restricted to a single report. Couvillion et al. (1984) found two species of horse flies, Tabanus lineola hinellus and T. nigrovittatus, infected with third-stage larvae of E. schneideri; and T. l. hinellus was considered the primary intermediate host.

The transmission of arterial worms is seasonal since it depends on the patterns of activity of horse flies, as well as seasonal aspects of the life history of the parasite in horse flies. Clark (1972) and Clark and Hibler (1973) provided data on the sea-

sonal occurrence of horse flies and larval *E. schneideri* in the western United States. The objective of this paper was to describe the temporal patterns of occurrence of *E. schneideri* in *T. l. hinellus* on South Island, South Carolina.

MATERIALS AND METHODS

The study was conducted on the Tom Yaw-key Wildlife Center, an island complex comprised of South, Cat, and North islands in Georgetown County, South Carolina. It was selected as the study site because white-tailed deer (Odocoileus virginianus) infected with E. schneideri have been found on the South Island portion of the area (Hibler and Prestwood, 1981) and also because potential intermediate hosts were abundant. This study was conducted on a 48-ha hammock on the northern section of South Island. Detailed description of the area has been given by Epstein (1983).

In 1981, female T. l. hinellus were trapped for consecutive 3-day periods at monthly intervals from July through September. In 1982, trapping was done for consecutive 2-day periods every 2 wk from mid-April through mid-October. Six canopy traps and two malaise traps (Roberts, 1976) were placed in openings adjacent to woodlands alongside a north-south road. Traps were set and baited with dry ice at daylight. Horse flies were removed alive at midday and again when trapping was terminated at dusk. Following transport to the laboratory, live horse flies were dissected in search of third-stage larvae of E. schneideri (Couvillion et al., 1984) and larvae were counted.

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TABLE 1. Numbers of female *Tabanus lineola hinellus* collected, examined, and found positive for third-stage larvae of *Elaeophora schneideri* and intensity of infection.

Date	Number of horse flies			Number of larvae in infected horse flies	
	Collected	Examined	Positive (%)	Average	Total
1981		-			-
6 Jul⁴	130	130	0 (0)	0	0
4Aug	69	66	1 (1.50)	15	15
8 Sep	745	517	0 (0)	0	0
29 Sep	1,293	1,276	4 (0.31)	24	95
Totals/means	2,237	1,989	5 (0.25)	22	110
1982					
13 Apr ^b	0	0	0 (0)	0	0
21 Apr	0	0	0 (0)	0	0
30 Apr	0	0	0 (0)	0	0
4 May	61	51	0 (0)	0	0
13 May	170	163	2 (1.23)	5	10
22 May	321	314	0 (0)	0	0
8 Jun	1,970	1,940	13 (0.67)	4	55
22 Jun	477	456	1 (0.22)	1	1
6 Jul	173	158	0 (0)	0	0
20 Jul	525	518	0 (0)	0	0
3 Aug	86	81	0 (0)	0	0
17 Aug	253	246	3 (1.22)	15	45
31 Aug	2,086	2,072	2 (0.10)	3	6
14 Sep	1,492	1,473	4 (0.27)	38	152
28 Sep	648	637	1 (0.16)	51	51
12 Oct	452	442	0 (0)	0	0
Totals/means	8,714	8,551	26 (0.30)	12	320
Overall totals/means	10,951	10,540	31 (0.29)	14	430

^{*} First day of consecutive 3-day collection period.

To confirm the presence of infected deer on the study area, 12 deer were examined at necropsy in search of *E. schneideri*. Samples of forehead skin were examined histologically for microfilariae. Representative specimens of adult *E. schneideri* and histologic sections of skin with microfilariae were deposited in the U.S. National Parasite Collection, Beltsville, Maryland (Accession Nos. 78487, adults; 78488, histologic sections).

RESULTS

Data on T. l. hinellus collected and examined for third-stage larvae of E. schneideri are summarized in Table 1. Overall, 10,951 T. l. hinellus were captured, and 10,540 were dissected for third-stage larvae (Couvillion et al., 1984). Sim-

ilar proportions of infected horse flies were captured at different (P > 0.05) traps.

During 1982, T. l. hinellus were present during all trapping periods over 22 wk from May through mid-October (Table 1; Fig. 1). Following the first appearance of T. l. hinellus on 4 May, the population increased rapidly, reaching a peak on 8 June (Fig. 1). Abundance then decreased until late August/early September, when there was another pronounced increase in numbers of horse flies captured. Horse flies were not trapped during May and June in 1981; therefore a complete assessment of temporal changes in abundance cannot be made for that year.

Overall, 0.29% of T. l. hinellus har-

First day of consecutive 2-day collection period except for 21 April through 22 May which were 1-day collection periods.

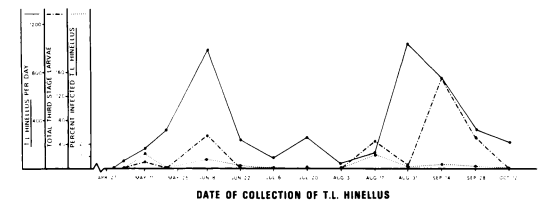


FIGURE 1. Temporal distribution of the abundance of T. l. hinellus, percent infected with third-stage larvae, and total number of third-stage larvae.

bored third-stage larvae (Couvillion et al., 1984). Average prevalence of infected T. l. hinellus was 0.25% for 1981 and 0.30% for 1982. Prevalence of infected T. l. hinellus varied among collection periods and ranged from 0 to 1.5% and 0 to 1.23% during 1981 and 1982, respectively (Fig. 1). Prevalence of infected T. l. hinellus was similar among collection periods for 1981 (P > 0.05, $\chi^2 = 6.028$); however, highly significant differences (P < 0.01, $\chi^2 = 22.69$) were noted during 1982. In 1982, prevalence of infected horse flies, like horse fly abundance, was biphasic with peaks occurring in mid-May (1.23%) and mid-August (1.22%) immediately preceding peaks in T. l. hinellus populations.

The mean number of third-stage larvae in infected T. l. hinellus was 14 for the two seasons combined, ranging from one to 47 ($\bar{x}=22$) in 1981 and from one to 64 ($\bar{x}=12$) in 1982 (Couvillion et al., 1984) (Table 1). During 1982 the intensity of infection of T. l. hinellus increased as time progressed (r=0.644). A quadratic regression model fit the data more precisely (r=0.728) and indicated that the intensity of infection varied non-linearly with time and increased to the highest level in late summer. This increase resulted in greater recovery of infective larvae during August-September (mean intensi-

ty = 25) than in May-June (mean intensity = 4) (Fig. 1).

Examinations for adult E. schneideri were done on 12 white-tailed deer over the 2-yr period. Of 10 deer collected in May and June of 1981, eight harbored adult E. schneideri and two were positive for microfilariae on histologic examination. In 1982, two deer were examined for adult E. schneideri in June and October. Microfilariae were found only in the deer examined in June; however, only a single dead adult worm was found in the carotid artery of the deer examined in October. Overall, infected deer harbored one to 31 $(\bar{x} = 6.5)$ adult nematodes. The intensities of infection with adult E. schneideri in deer with and without microfilariae were nine to 31 ($\bar{x} = 18$) and one to three ($\bar{x} = 18$) 1.5), respectively. Few microfilariae were seen in deer that were positive. Four deer without microfilariae harbored one nematode each.

DISCUSSION

The production of two generations of adult T. l. hinellus on South Island in 1982 was similar to the situation for T. l. hinellus in coastal Texas (Thompson, 1973) and T. l. lineola in Alabama (Burnett and Hays, 1977) and South Carolina (Sheppard, 1972). This is in contrast to most

temperate climate species of horse flies that produce only one generation per year (Harwood and James, 1979).

Differences among trapping periods in prevalence of infected horse flies were attributed in part to fluctuations in the ratio of blood-fed (potentially infected) to unfed horse flies. For instance, during peaks in T. l. hinellus populations, the ratio of blood-fed to unfed horse flies would be small because large numbers of newly emerged T. l. hinellus were attracted to traps while seeking their first blood meal.

Although deer were not examined to determine whether there was a seasonal change in numbers of microfilariae in forehead skin, the temporal change in the intensity of infection of horse flies raises the question of a circannual rhythm of microfilariae in deer, with highest levels in late summer. Circannual rhythms of microfilariae have been reported for *Onchocerca gutturosa*, a filarial parasite of cattle (*Bos taurus*) in which microfilariae reside in skin (Eichler, 1973).

Compared to primary intermediate hosts of the arterial worm in the western United States, the prevalence and intensity of infection of T. l. hinellus on South Island were low. The average prevalence of infection for Hybomitra spp. was as high as 19.1% in one area of New Mexico (Hibler et al., 1971). Average prevalences of H. laticornis and H. aatos with thirdstage larvae in New Mexico were 8.8% and about 2.0%, respectively (Clark and Hibler, 1973; Davies, 1979). The mean intensity of infection of T. l. hinellus with third-stage larvae was less than half of that reported for infected horse flies in the western United States by Clark and Hibler (1973), and only about one-fourth of that of H. aatos (Davies, 1979). The low prevalence and intensity of infection of T. l. hinellus probably were due to the low prevalence and few microfilariae of patently infected deer and/or to biological and behavioral characteristics of horse flies

that affect uptake and development of microfilariae.

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