The Effect of Phytophagous Nematode Grazing on Blue Grama Die-off

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Abstract

Nematode populations were sampled in ungrazed and heavily grazed areas in northeastern Colorado under patches of healthy, senescing, and dead blue grama to test the hypothesis that phytophagous nematodes may cause the senescence. Densities of plant parasites were significantly different under the 3 plant types. Live blue grama supported the highest numbers (1.2 X 10⁵/m²) and dead blue grama, the lowest (2 X 10⁴/m²). Bacterial feeding nematodes also varied significantly with plant type. Highest densities were found under senescing plants (2.4 X 10⁵/m²) and lowest densities were under dead plants (7 X 10⁴/m²). Total densities were slightly but insignificantly lower in the heavily grazed area. Scarlet globemallow and fringed sagewort supported lower populations of both plant parasites and bacterial feeders than did live blue grama. The densities under live blue grama were not unusually high and well within the values reported in the literature for arid grasslands.

Methods

Study Site

The study was conducted on the Central Plains Experimental Range (CPER) operated by the USDA Service, Agricultural Research, 18 km NE of Nunn, Colo. The vegetation is shortgrass prairie dominated by blue grama with other grasses, sedges, forbs, and pricklypear cactus (Opuntia polyacantha Haw.) also present. The mean annual precipitation at CPER is 31 cm, based on 30 years of data. May through August are the wettest months, accounting for more than 50% of the annual precipitation. Monthly mean temperatures vary from a high of 22°C in July to a low of -2°C in January.

Sampling Areas

Two areas were selected for sampling. One was in the Environmental Stress Plot (ESA) initially established by the Grassland IBP. The site we selected was adjacent to areas previously manipulated but it had not received any treatment other than relief from ungruelate grazing pressure. The other site had been heavily grazed by cattle during summers since 1939. In both areas, soil is an Ascalon sandy clay loam and blue grama was the dominant species.

Both areas could be characterized by numerous patches of senescing and dead blue grama. The most prevalent pattern was apparently healthy blue grama surrounded by senescing plants that could be identified by the mixture of dead and live leaves. In the center of these patches were dead blue grama plants. Most dead plants were widely spaced. Thus it appeared that the die-off began in the center and moved outward in all directions.

A large representative patch in each area was selected for sampling and was divided into 4 quarter circle quadrants. Two cores, randomly selected, were taken from each quadrant within each of the 3 plant types for a total of 8 replicates taken each time of sampling. Cores were 4.8 cm in diameter and 10 cm deep. In 1978 the ungrazed plot was sampled on June 5, June 27, and August 4, and the grazed plot was sampled on June 27 and August 4.

Vegetation was not sampled on the plots where the soil samples were taken; but adjacent dead patches, senescing areas, and healthy areas were sampled on .09 m² quadrats. Weight of individual species was estimated and converted to oven-dry weights based on clipped samples.

The dead patches were dominated by annual forbs in both areas (Table 1). Pricklypear cactus and scarlet globemallow (Sphaeralcea coccinea (Pursh) Rydb.) were the main perennial species occurring in the dead patches in both areas. Fringed sagewort (Artemisia frigida Wild.) was also abundant within the dead patches in the ungrazed area. Additional cores for nematode extraction (8 replicates in each area) were taken within the quadrants of the dead patches only to include the roots of scarlet globemallow in the grazed area and the roots of fringed sagewort in the ungrazed area. Each soil core was returned to the laboratory, suspended in cold water, and the suspension sieved (No. 325 sieve) to remove nema-
todes. The material retained by the sieve was put on a Baerman funnel for 48 hours (Christie and Perry 1951). The number of nematodes in each sample (and their trophic category) was determined by counting individuals present in each of 3-1 ml subsamples of a 50-ml suspension as described by Smolik (1974). All counts were corrected for extraction efficiency, which averaged 60% for bacterial feeding nematodes, 46% for Dorylaimida, and 59% for the plant parasites. These were the 3 dominant trophic groups. The Dorylaimida consists of species which are plant parasites, predators and omnivores. But they were not assigned to trophic groups since species determination was not possible within the scope of this study.

**Laboratory Experiments**

To determine nematode response to different moisture levels, wet or dry, 16 senescent and 16 live blue grama plants were dug up from the ungrazed area and potted in 10 by 10 cm plastic pots in the field on August 4, 1978. They were returned to the laboratory and placed in an environmental chamber which was set to duplicate both summer temperature and light regimes of the Central Plains Experimental Range. Half of the senescing plants and half of the live plants were watered every 2 days (wet treatment) and the remaining half were watered every 4 days (dry treatment) to field capacity. On November 4, the nematodes were extracted and counted as described above.

**Analysis**

Logarithmic transformations of nematode counts were accomplished to provide normally distributed data for analysis of variance. The model for the field study was a stratified random design with 2 factors: date and plant type. There were 8 replicates for each of 4 plant types on each of 2 or 3 sampling dates. Differences between means were compared by Least Significant Range (LSR, Sokal and Rohlf 1969).

The laboratory data were also log transformed for ANOVA. The model is a completely randomized design with 2 factors: plant type and moisture regime.

**Results**

**Field Study**

On the ungrazed plot there was both a significant date (α ≤ 0.001) and plant (α ≤ 0.01) effect on plant parasites (Fig. 1). On all sampling dates, densities were highest under live blue grama, intermediate under senescing blue grama, and lowest under fringed sagewort and dead blue grama. Densities of plant parasites were high in the spring, low in late June, and intermediate in August under live and senescing blue grama and under the fringed sagewort. Plant parasites under dead blue grama declined in mid-season but did not increase in August.

Densities of bacterial feeders (microbivores) averaged about twice those of the plant feeders and there was also a significant date (α ≤ 0.005), plant (α ≤ 0.001) and interaction effect (α ≤ 0.01). As with the plant parasites, under the live and senescing blue grama and fringed sagewort, an early season high was followed by a significant mid-season decline (α ≤ 0.05) with a subsequent increase in August (Fig. 1). The significant date-treatment interaction was caused by this mid-season high sandwiched between early and a late season low densities under dead blue grama. It is also interesting to see that on all dates there were more microbivores under senescing than under live blue grama.

The Dorylaimida (Fig. 1) which includes omnivores, plant parasites and predators experienced a significant treatment effect (α ≤ 0.001) and a significant date-treatment interaction (α ≤ 0.05). Generally, the changes reflect those of the microbial feeders except that the August densities for Dorylaimida were higher or equal to the spring densities. The Dorylaimida population under dead blue grama also appeared to have a mid-season high, but...
there was no significant difference in density among the 3 sampling dates.

The heavily grazed plot was sampled only in late June and early August. For the plant parasites (Fig. 2) there were significant date (α≤0.01) and treatment effects (α≤0.05). As in the ungrazed plots, a late June low was followed by a significant increase in August under the live and senescing plants.

A similar pattern was exhibited by some of the microbivores (Fig. 2) with a significant date (α≤0.001) and interaction effect (α≤0.001). The most notable change was an increase in densities in the live and senescing plots from June 27 to August 4. The other treatments did not change significantly.

For the Dorylaimida the date (α≤0.01), treatment (α≤0.001) and interaction (α≤0.01) effects were all significant. Again, there was an increase under live and senescing plants from June to August. The scarlet globemallow population declined and the populations under dead blue grama increased but neither change was significant.

Although the ANOVA was significant in all cases, a comparison of individual mean densities at each date, usually revealed a significant difference only between the extreme values; e.g., in Figure 1 on August 4, the plant parasites were significantly different only under live blue grama (the highest density) and dead blue grama (the lowest density). Further individual comparisons may be made by the LSR values indicated in each Figure.

Discussion

Both field and laboratory studies have shown that control of nematodes from shortgrass and mixed grass prairies could significantly increase net primary production (NPP) over the short term. Smolik (1977) found that nematicide application to mixed grass prairie vegetation increased harvestable herbage by 29–59%. Stanton et al. (1981) applied nematicide to plots on the Central Plains Experimental Range and root production increased about 25%. In additional laboratory experiments to measure blue grama response to both above- and below-ground grazing, Stanton (1983) indicated that populations of one species of plant parasite feeding on blue grama for 140 days reduced NPP about 13%. The experimental densities were lower than field densities by at least one-half and were far less species rich (one blue grama plant in the field may host several species of plant parasites). Thus, even at normal field densities, nematodes may exert strong grazing pressure on blue grama roots.

Phytophagous nematodes probably influence plant growth; conversely, changes in root biomass may also affect nematode density. The seasonal pattern in density (early high, mid-season low, late season high) is probably a response to new roots and changes in soil moisture. Field observations, through windows placed on excavations at the Central Plains Experimental Range (Ares 1976), revealed that root growth began a short time before leaf growth in the spring. However, mid-season soil desiccation resulted in the death of 30–60% of the new roots. Late in the season with high water potential, roots again increased in length. However, it is uncertain whether the nematodes die with root die-off or enter anhydrobiosis. Anhydrobiosis is a quiescent state induced by...
lowering of soil moisture. This phenomenon has been described in a number of species (see review by Demeure and Freckman 1981) and is probably one of the reasons nematodes are so abundant in arid grasslands. They can essentially wait through periods of low soil moisture with little or no metabolic activity. When conditions become favorable, the nematodes again become active. Plant parasites were most abundant under live, healthy plants because they prefer to feed on new roots. Surprisingly, plant parasite densities under dead blue grama and fringed sagewort did not differ. Fringed sagewort generally supports fewer plant parasites than blue grama (Stanton et al. 1981). Volatile compounds such as terpenes are present in the root exudates of fringed sagewort which may deter belowground herbivores. Scarlet globemallow, also with low nematode densities, has a long, deep tap root so it probably provides fewer feeding sites than the shallow fibrous roots of blue grama. Thus, these 2 plant species do not support as large a population of nematodes as blue grama and might be at a selective advantage under stress conditions when nematode populations are high.

Microbivores were the most abundant trophic class, varying from 4 to 2.2 \times 10^5 \text{m}^{-2} and populations were highest under senescing plants. Microbivores feed on bacteria and the dying roots probably provided more resources for the microbial populations. Another major carbon source for bacteria is root exudates. High exudation rate probably correlates with new root growth. Thus, with favorable moisture regimes in the spring and high root exudation, microbial populations should be high.

In the ungrazed plot the microbivore densities under dead blue grama reached a maximum in mid-season probably because the dead roots provided a large source of carbon. This is consistent with findings from our laboratory experiment in which the highest microbivore densities were from drought-stressed senescing plants.

The Dorylaimida densities were within the range of the plant parasites, and the seasonal trends in their density almost paralleled those of the microbivores. Since many of the Dorylaimida species are predators, they may use the very abundant bacterial-feeding nematodes as a major prey item.

In summary, plant parasitic densities on both the grazed and ungrazed plots under live blue grama varied from $1 \times 10^5$ to $1 \times 10^8 \text{m}^{-2}$, well within the range reported from other areas of the Central Plains Experimental Range (R. Anderson personal communication, Stanton et al. 1981). Since these densities are not abnormally high, it is unlikely that nematodes were the cause of the die-off, although they certainly may have been a contributing factor. It is also possible that plant species invading the die-off patches (e.g. fringed sagewort and scarlet globemallow) are successful in part because of chemical or morphological traits that make them resistant to nematode and white grub feeding. Since nematodes may remain viable but inactive in the soil for many years, plant species which provide a root system that is palatable and accessible to nematodes may be difficult to establish even with proper climatological conditions. This idea is speculative but certainly should be investigated further.

References