Long-term changes in forest floor processes in southern Appalachian forests

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Abstract

Soil nutrient concentrations decreased in an aging southern Appalachian forest over a 20-year period. Construction of nutrient budgets showed significant nutrient sequestration aboveground including increased forest floor mass. We hypothesized that the changes in forest floor mass resulted from decreased litter decomposition rates because of decreased litter quality. In 1992 and 1993, we repeated a litter decomposition experiment conducted in 1969 and 1970 to test this hypothesis. In addition, we examined microarthropod populations and functional groups as litter decomposed. For four of the five species tested, first-year decomposition rates were about the same in both experiments. Initial litter nutrient concentrations of P were lower in all tree species in the most recent sampling. N, Ca, and Mg concentrations also declined in some species. These declines often resulted in decreased nutrient release rates during decomposition. Microarthropod populations differed significantly among litter species, as well as between years (probably resulting from differences in growing-season rainfall). For some litter species we found significant relationships between microarthropod populations and nutrient concentration (primarily C and N); however, most r²-values were low. Data suggest that changes in forest floor mass probably resulted from decreased litter quality and that those changes may have an effect on microarthropod populations.

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1. Introduction

Forests rely largely on internal nutrient cycling processes to maintain plant growth (Cole, 1995; Perala and Alban, 1982). The forest floor, with nutrient additions from foliage, fine wood, and coarse woody debris, plays a significant role in returning nutrients to the soil through decomposition. Federer (1984) and Covington (1981) have shown that forest floor mass increases during forest stand development. Prescott et al. (2000) suggested that forest floor accumulates with time, especially in boreal forests, and represents a
large pool of nutrients. Its decomposition is essential for nutrient availability in the short-term, but it may immobilize nutrients in the long-term.

Forest floor type has an influence on litter decomposition rates, and forest floor chemistry differs significantly among sites (Grigal and Ohmann, 1989). Litter decomposition rates involve the interaction of vegetation, soil nutrient availability, micro- and macro-fauna, microbial populations, and environmental conditions. Decomposition rates can provide an accurate prediction of soil and site quality or productivity (Johansson, 1994). Increasing rates of litter decomposition suggest accelerated nutrient cycling rates within the site. On a global scale, rates of litter decomposition are regulated by climate (Johansson et al., 1995). Hornsby et al. (1995) used microcosms to alter temperature and moisture in a loblolly pine plantation. He found that increased moisture and temperature increased decomposition rates. Murphy et al. (1998) studied decomposition of several native species along an elevation gradient in Arizona. They concluded that decomposition was largely regulated by moisture. Upon examination of the relationship between nutrient concentrations and decomposition rates, their data suggested that decomposition is limited by C availability, not nutrients. Melillo et al. (1989) found that in the initial phase of leaf litter decomposition (44 months in their study) cellulose is removed but the later phase focuses on lignin breakdown. Berg et al. (Berg and Meentemeyer, 2002; Berg et al., 2000) examined litter quality and decomposition rates across a transect in northern Europe. They found that substrate quality, specifically Mn concentration, explained 27% of the decomposition pattern, because of its influence on lignin degradation. In a later study they used the final stage of litter decomposition, the limit value, as an indicator of how climatic changes would alter the forest floor (Berg and Meentemeyer, 2002). The limit value of litter decay was unaffected by climate but was negatively related to initial litter N concentration and N additions caused decomposition to stop entirely.

Other research studies have shown that litter type controls decomposition rates. Keenan and others (1996) compared decomposition rates in coniferous forests of British Columbia. They examined western redcedar, western hemlock, and lodgepole pine and found that litter species, not site microclimate differences, regulated decomposition rates. They also concluded that litter type controlled site differences in N availability. Taylor and others (1989) improved overall litter quality by mixing slow and rapidly decomposing litter types. They found that decay rates were most similar to the rapidly decomposing species. Their data suggested that litter quality controls rates of decomposition, regardless of the environmental condition (Taylor et al., 1991). This was also the conclusion of an experiment examining nutrient cycling rates across a climatic and vegetation gradient in mixed temperate forests of the southern Appalachians (Knoepp et al., 2000). At the Coveeta Hydrologic Lab in western NC, litter decomposition rates were determined on five sites representing a gradient in vegetation type. Sites included a xeric mixed oak-pine community with low nutrient availability, two mixed oak sites (860 and 1090 m elevation), a nutrient rich cove hardwood forest, and a high N northern hardwood forest. Litter decomposition rates were determined for three species: two overstory species, Liriodendron tulipifera and Quercus prinus; and one evergreen understory species, Rhododendron maximum. On average, the two mixed oak sites, MO-H and MO-L, had the highest decay constant; followed by the northern hardwood, the mixed-oak pine, and the cove hardwood communities. Rates were not directly influenced by climate (high temperature or moisture) or nutrient availability (Knoepp et al., 2000) suggesting that additional or a combination of factors regulate litter decomposition.

Research on the relationship between nutrients and litter decomposition has generated variable results. Foliar nutrient content is often representative of soil nutrient content or availability (Stump and Binkley, 1993; Wang and Klinka, 1997). Monleion and Cromack (1996) found that nutrient release and immobilization patterns during the initial phases of decomposition suggest which nutrients are limiting on a site. However, Prescott et al. (1993) found that while foliar N and P concentration influenced immobilization of these nutrients during decomposition, it was not related to soil nutrient availability. Aerts and De Caluwe (1997) used field grown, low N and high N litter to determine effects of nutrient concentration on decay rates and nutrient release. They found that high N species did not always have the greatest decomposition rates.
Soil and forest floor microarthropods and microbes are important as primary decomposers and litter transformers. Their activity changes the physical and chemical characteristics of the litter layer and is thus intimately connected with decomposition processes (Wardle, 2002). Microarthropods differ in their functional role of decomposition both among and within groups. For example, Addison et al. (2003) found that all adult Collembola examined fed on fungal hyphae; some also feed on decomposing plant material, organic matter, and invertebrate fecal material.

In a previous study on a north-facing hardwood (WS 18) and a white pine plantation (WS 17) watershed in the Coweeta Hydrologic Laboratory basin, we found a significant decrease in surface soil extractable cations (Ca 68%; K 44%; Mg 77%) and total N (43%) between 1970 and 1990 (Knoepp and Swank, 1994, 1997). Construction of nutrient budgets showed that these aggrading forests sequestered up to 50% of the nutrients aboveground, the rest was lost from the site via leaching to stream water. Along with the decrease in soil nutrients we measured an increase in forest floor mass of 2100 kg ha⁻¹ in the hardwood watershed and 5600 kg ha⁻¹ in the white pine plantation. However, because forest floor nutrient concentration decreased, there was no change in the total nutrient content in the hardwood watershed, 99 versus 93 kg Ca ha⁻¹. Forest floor nutrients decreased in the white pine watershed from 92 to 81 kg Ca ha⁻¹ in 1970 and 1990, respectively. We hypothesized that the forest floor mass increase resulted from decreased rates of litter decomposition because of decreased litter quality, the result of decreased soil nutrient availability.

At a similar time to the initial soil sampling of the Knoepp and Swank (1994) study (1969 and 1970) Kermit Cromack measured decomposition rates of foliage from five tree species common to the Coweeta basin (Cromack and Monk, 1975). They used four hardwood species in WS 18, Quercus prinus L., Acer rubrum L., Quercus alba L., and Cornus florida L., and one in WS 17, Pinus strobus L. Our objective was to repeat Cromack’s experiment in these watersheds using the same methods to examine long-term changes in decomposition and nutrient release rates comparing 1969 and 1970 to 1992 and 1993. We hypothesized that decomposition rates decreased over this time period, resulting in the increased forest floor mass measured in the soil resampling study (Knoepp and Swank, 1994). In addition, we examined microarthropod populations associated with the different litter species to determine potential relationships between decay rates and forest floor microarthropods. This information will be useful in predicting the effects of changes in nutrient pools and overstory species composition on microarthropod biodiversity.

2. Methods

2.1. Site description

This study was conducted at the Coweeta Hydrologic Laboratory, a 2180-ha USDA Forest Service facility in the southern Appalachians of western North Carolina, USA. Annual precipitation is 1900 mm with most months receiving at least 100 mm. The growing season extends from early May to early October. Mean monthly temperatures are highest in June through August (20 °C) and lowest in December through January (5 °C). Our study was within two Coweeta basin watersheds. Watershed 18 (WS 18) is a 13-ha north-facing reference watershed dominated by mixed oak hardwood vegetation; no forest management has been conducted there since the late 1920s. Watershed 17 (WS 17), a 13-ha north-facing watershed, was part of a water yield experiment and was clearcut repeatedly from 1940 to 1955; no materials were removed. In 1956, a white pine (Pinus strobus) plantation was established on WS 17 (Swank and Crossley, 1988).

2.2. Litter decomposition measurements

We repeated Cromack’s 1969 and 1970 (70s) litter decomposition study (Cromack and Monk, 1975) in 1992 and 1993 (90s) using the same tree species; Quercus prinus L. (chestnut oak, CO), Acer rubrum L. (red maple, RM), Quercus alba L. (white oak, WO), Cornus florida L. (dogwood, DW), and Pinus strobus L. (white pine, WP). We used the same plot locations and methodology. From September through November 1992 and 1993 we collected freshly fallen leaf litter by species from several large areas of WS 18 and WS 17. Litter was air dried and mixed thoroughly to ensure minimal variability among subsamples. We put
approximately 2.5 g of hardwood litter and 5.0 g of white pine litter (exact weight recorded) in 10 cm²; 1-
mnm mesh fiberglass screen litterbags (Crossley and
Hoglund, 1962) and tagged each with an identifying
number. We randomly selected four subsamples, and
dried them at 50 °C to determine the ratio of air-dried
to oven-dried weight, and for initial chemical analysis.

Hardwood leaf litterbags were placed in a grid
pattern (0.5 m apart, randomized by species and
collection date) in three locations (lower, middle, and
upper portion) within WS 18. Similarly, we placed WP
leaf litterbags within WS 17. Litterbag installation
occurred on December 17, 1992; and December 14,
1993, placing 15 and 19 litterbags of each species in
each location in 1992 and 1993, respectively. One
litterbag of each species was randomly collected from
each grid location, lower, middle, and upper, every 27–
35 days in 1992 (13 total collections) and every 25–56
days in 1993 (18 total collections)—shorter intervals in
the summer and longer intervals in the dormant season.
We collected all remaining litterbags at the final
collection date. Total length of the decomposition study
was 394 days in 1992 (final collection January 1994)
and 710 days in 1993 (final collection November 1995).

2.3. Collection and chemical analyses

We collected litterbags from the field, returned them
to the laboratory, and refrigerated at 4 °C. Following
microarthropod extraction (described below), litterbags
were oven dried at 50 °C, removed from the mesh bags,
and weighed. Oven-dried samples were ground using a
Wiley mill with a 2-mm mesh for chemical analysis.
Material was muffled (480 °C) and digested in a
solution of 20% nitric acid. Both 70s and 90s sample
analyses were conducted using the same nitric acid
digestion procedure. In 70s nitric acid-digested samples
were analyzed using direct-reading spark-emission
spectroscopy for Ca, K, and Mg; P was determined
colorimetrically on an autoanalyzer. Total N was
determined using micro-Kjeldahl digestion technique
(Cromack and Monk, 1975). In 90s, nitric acid
digests were analyzed for Ca, K, and Mg using atomic
absorption spectroscopy (Perkin-Elmer 300), P was
determined colorimetrically, and total N and C were
determined by combustion using a Perkin-Elmer 2400
CHN analyzer. In 90s ashed samples were weighed after
muffling to determine ash free dry weight (AFDW).

We applied an exponential decay equation (ln(final
weight/initial weight) = -kt) (Olson, 1963) to data
from all plots and both years to determine a single
decay constant (k) for 1 year, for each species using
the regression procedure of SAS (SAS, 2000). Comparisons between 70s and 90s data were based
on 1 year of litter decay 365 ± 13 days; actual portion
of a year was used for calculations. White pine decay
rates were based on 10 months of litter decay due to
missing data points in the archived 70s dataset. We
also used this approach with nutrient concentrations
(μg g⁻¹) to determine nutrient release rates during
decomposition in the two studies. We calculated decay
rates for each litter species for 90s only using
AFDW.

2.4. Microarthropods

At each collection date litterbags were collected,
returned to the laboratory, and refrigerated at 4 °C. The
next day, we placed the litterbags on Berlese funnels for
microarthropod extraction. Microarthropod samples
were stored in 100% EtOH until processed. Microar-
thropod populations were determined in litterbags
collected after 9–12 months of decomposition for the
1992 (n = 19) and 1993 (n = 7) decomposition study.
Microarthropods were separated into the following
groups: Collembola, mature oribatid, immature oriba-
tid, mesostigmatid, and prostigmatid mites. Total
microarthropod population in each litterbag also was
determined. Microarthropod abundances were deter-
mined as the number of animals per gram dry litter. We
calculated populations per gram litter AFDW for
analysis of microarthropods relationship with litter
nutrient concentrations.

2.5. Statistical analysis

Significant differences between calculated decay
rates and nutrient release rates for 70s and 90s data sets
were determined using a paired t-test comparison. We
computed the standard deviation of the difference
(S.D. diff) for 70s and 90s decay rates using the standard
error (S.E.) for k-values of each species computed by
SAS (S.E. = sqrt(S.E. ² k0 + S.E. ² k0)); setting t = 1.671
for probability < 0.05 with 62 d.f. (S.D. diff =
S.E. * 1.671). Differences between the 70s and 90s
decay rates and nutrient release rates greater than
S.D.diff are significant. We used the same method to determine significant differences among decay rates on an AFDW basis of litter species in 90s.

The archived 70s data set did not allow us to compute standard errors for initial nutrient concentrations. We estimated the S.E.70s using the proportion of S.E.90s to mean initial nutrient concentrations. We computed S.D.diff, using S.E. * 2.131 (probability < 0.05, d.f. = 4) for a t-test as described above. Differences in nutrient concentrations between 70s and 90s greater than t-value are significant.

Microarthropod abundance values were not normally distributed, we used Generalized Linear Models (GLIM) in the Genmod Procedure of SAS (SAS Systems for Windows, 8e (SAS, 2000)), using Poisson distributions and log link functions (Littell et al., 2002). We examined the relationship between litter nutrient concentration and microarthropod populations per gram AFDW data with the Stepwise Procedure of SAS (SAS, 1985). Default settings of Stepwise were used; the final model is presented for relationships identified with a probability of a value greater than F ≤ 0.05.

3. Results and discussion
3.1. Litter decomposition rates

One-year decay rates for chestnut oak (CO), dogwood (DW), red maple (RM), and white pine (WP) did not differ significantly for measurements conducted in 70s and in 90s (Table 1). White oak litter decay rates decreased significantly from k = 0.68 in 70s to k = 0.54 in 90s. At both measurement periods DW exhibited the greatest decay rate, k = 1.26 in both 70s and 90s. Lowest decay rates occurred in WP, k = 0.55 in 70s and k = 0.49 in 90s. For all species, we examined variation in decay rates between years within a study period. We found decay rates in the 70s study were very similar; rates varied between 8 and 19% for all species and differences were greatest for the species with the lowest decay rate. Variation between years was greater for the 90s study period, ranging from 8 to 32%. Again, differences between years were least for rapidly decomposing species (8% for D) and greatest for the slow decomposers (25, 25, and 32% for WO, WP, and CO).

<table>
<thead>
<tr>
<th>Leaf species</th>
<th>CO</th>
<th>D</th>
<th>RM</th>
<th>WO</th>
<th>WP</th>
</tr>
</thead>
<tbody>
<tr>
<td>CROMACK (69 and 70)</td>
<td>0.58 (0.03)</td>
<td>1.26 (0.03)</td>
<td>0.84 (0.03)</td>
<td>0.68 (0.03)</td>
<td>0.55 (0.04)</td>
</tr>
<tr>
<td>KNOEPP (92 and 93)</td>
<td>0.53 (0.03)</td>
<td>1.26 (0.03)</td>
<td>0.87 (0.03)</td>
<td>0.54 (0.03)</td>
<td>0.49 (0.02)</td>
</tr>
</tbody>
</table>

k values where ln(initial weight/final weight) = kr, r = time in portion of year. Decay rates based on 1 year (365 ± 13 days). WP decay rate based on 300 days of decomposition. Values in parentheses are standard error of the regression equation.

* Significant difference between 69 and 70 and 92 and 93 based on a paired t-test comparison (p < 0.05, d.f. = 62).

We examined climatic differences between 70s and 90s and also between years at each period for insights into regulation of decay rates during these two studies. There were no differences in mean annual temperature between the two study periods. In both 70s and 90s studies, average annual air temperature was within 0.3 °C of the 69-year average temperature at the main climate station of Coweeta Hydrologic Lab (Fig. 1). Average growing season temperatures for 70s and 90s were within 0.8 °C of the long-term average. In contrast, precipitation patterns varied both between and within the studies (Fig. 2). Total rainfall varied in summer and fall, the two periods with highest temperatures and maximum decomposition rates. Summer

![Graph of monthly temperature](image)

Fig. 1. Mean monthly temperature (°C) at Coweeta Hydrologic Laboratory main climatic station during litter decomposition studies initiated in 1969, 1970, 1992, and 1993; also shown is the 69-year average monthly temperature.
rainfall at Coweeta averages 41 cm. During the two summers in the 70s litterbag study, total rainfall was 6 and 3 cm greater than average. The 90s summers were more extreme; total summer rainfall was 21 cm below average 1 year and 26 cm above average the next. The differences in rainfall, and resulting differences in soil moisture availability, between the 2 years may help explain differences in decay rates. Fall rainfall showed less variation; 70s study years had a 26% increase and 14% decrease in rainfall; in 90s there was a 15% increase and a 2% decrease. This variation did not appear to affect decomposition patterns.

Some research has suggested that litterbag studies are useful as indices but are not absolute decay rates. Blair and others (1990) examined the litterbag method, comparing single species and mixed species litterbags to determine if measured decomposition rates accurately estimate nutrient release or accumulation from forest litter. They found that single species litterbags may underestimate decay rates and nutrient release or accumulation from decomposing litter. Pillers and Stuart (1993) compared litterbag decay and a mass balance approach to calculate decomposition rates in interior and coastal redwood stands in California. They found that each method identified a different site as having the maximum rates of forest floor turnover. The mass balance method was significantly correlated with environmental conditions (summer relative humidity, temperature, vapor-pressure deficit, and litter moisture); the litter bag method was not.

3.2. Nutrient concentration and release

We measured significant changes in nutrient release patterns from litter during the first year of decomposition between 70s and 90s (Table 2). Phosphorus release decreased significantly in all five species measured (Table 2 and Fig. 3). White oak and

Table 2
Rate of nutrient release (positive value) or accumulation (negative value) during litterbag decomposition for four deciduous (chestnut oak, CO; dogwood, D; red maple, RM; white oak, WO) and one conifer (white pine, WP) species

<table>
<thead>
<tr>
<th>Leaf species</th>
<th>P 69 and 70</th>
<th>92 and 93</th>
<th>N 69 and 70</th>
<th>92 and 93</th>
<th>Ca 69 and 70</th>
<th>92 and 93</th>
<th>Mg 69 and 70</th>
<th>92 and 93</th>
<th>K 69 and 70</th>
<th>92 and 93</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO</td>
<td>0.47 (0.05)</td>
<td>−0.14</td>
<td>0.06 (0.07)</td>
<td>−0.00 (0.03)</td>
<td>0.20 (0.07)</td>
<td>0.15 (0.05)</td>
<td>0.91 (0.12)</td>
<td>0.74 (0.07)</td>
<td>1.65 (0.13)</td>
<td>1.36 (0.09)</td>
</tr>
<tr>
<td>D</td>
<td>0.82 (0.07)</td>
<td>0.47 (0.05)</td>
<td>0.64 (0.09)</td>
<td>0.36 (0.03)</td>
<td>1.00 (0.04)</td>
<td>0.71 (0.06)</td>
<td>2.51 (0.15)</td>
<td>1.63 (0.08)</td>
<td>3.92 (0.22)</td>
<td>4.05 (0.14)</td>
</tr>
<tr>
<td>RM</td>
<td>0.65 (0.04)</td>
<td>0.00 (0.05)</td>
<td>0.06 (0.04)</td>
<td>0.11 (0.04)</td>
<td>0.38 (0.04)</td>
<td>0.49 (0.05)</td>
<td>1.73 (0.13)</td>
<td>1.19 (0.08)</td>
<td>2.35 (0.13)</td>
<td>2.35 (0.14)</td>
</tr>
<tr>
<td>WO</td>
<td>0.57 (0.02)</td>
<td>0.13 (0.06)</td>
<td>0.02 (0.03)</td>
<td>0.07 (0.02)</td>
<td>0.32 (0.03)</td>
<td>0.11 (0.05)</td>
<td>0.70 (0.09)</td>
<td>0.36 (0.07)</td>
<td>2.19 (0.19)</td>
<td>1.05 (0.08)</td>
</tr>
<tr>
<td>WP</td>
<td>0.71 (0.08)</td>
<td>0.15 (0.04)</td>
<td>0.09 (0.12)</td>
<td>0.44 (0.04)</td>
<td>0.42 (0.04)</td>
<td>0.32 (0.06)</td>
<td>1.66 (0.13)</td>
<td>0.97 (0.04)</td>
<td>2.06 (0.24)</td>
<td>2.11 (0.18)</td>
</tr>
</tbody>
</table>

Rates shown are k values where \( \ln(\text{initial nutrient content/final nutrient content}) = kt \), \( t \) = time in portion of year. Decay rates based on 1 year (365 ± 13 days). WP decay rate is based on 300 days of decomposition. Values in parentheses are standard error of the regression equation.

* Significant difference (d.f. = 62; \( p < 0.05 \)) between 69 and 70 and 92 and 93 based on a paired t-test comparison.
WP released P at very low rates in 90s compared to 70s. However, RO and CO shifted from releasing P in 70s to either retaining or accumulating P in 90s. Release of Ca changed significantly between 70s and 90s, decreasing for D, and WO but increasing for RM. Patterns of N release changed significantly for DW and WP; DW released significantly less N in 70s compared to 90s, release rates increased for WP.

Nutrient release from decomposing litter may indicate nutrient limitations in either the decomposing material or the site. Laskowski et al. (1995) examined changes in nutrient content during decomposition and found nutrient release and retention were driven by biological, physical, and chemical processes. Of the nutrients we measured, they identified changes in N, Ca, and Mg, as being regulated by biological processes whereas K release is a physical process. Blair (1988a) measured the release of cations from decomposing litter on a south-facing hardwood watershed in the Coweeta basin using three of the species we used; D, RM, and CO. His results were similar to ours, with the greatest Ca, Mg, and K release from DW followed by RM then CO (Table 2). In another Coweeta study, Heneghan et al. (1999) compared decomposition rates of CO in the tropics and in the temperate Coweeta basin. He found that although decomposition rates were greater for CO placed in the tropical location, litter at both sites had the same N concentration when 50% of the initial litter remained.

Comparison of litter nutrient concentrations before placement in the field showed that in many instances, initial nutrient concentrations were significantly lower in 90s compared to the 70s sample period (Table 3). In most cases these lower initial concentrations coincide with the reduction in nutrient release rates presented above. Initial P concentrations were significantly lower for all five-leaf species in 90s compared to 70s. Dogwood and WP litter also had lower concentrations of N and Ca. White oak litter also decreased in Ca concentrations. The decrease in litter nutrient concentrations coincided with a decrease in soil nutrient content on WS 18 and WS 17 during a similar 20-year period, 1970 to 1990 (Knoepp and Swank, 1994; Knoepp and Swank, 1997). The cations Ca, K, and Mg, as well as total N decreased significantly in both the A and AB horizons in WS 18. Concentrations also decreased in the A horizon of WS 17, but not in the AB horizon. Initial soil concentrations in WS 17 were much greater in 1970, because of the watershed level treatment during a water use experiment in which all vegetation was cut, and left in place in most years between 1942 and 1955 (Swank and Crossley, 1988).

Aerts and De Caluwe (1997) examined the effects of variation in leaf litter chemistry on decomposition and nutrient release using field-grown litter from low N and high N sites. Decomposition rates were not always greater for the high N species of litter. However, increased nutrient content of a single litter species decreased the immobilization of N and P, increasing nutrient release rates. Combining the 70s and 90s data also shows this pattern. The decrease in initial litter nutrient concentration resulted in a lower rate of nutrient release from each species. This pattern was observed among species within the 90s studies;
Table 3
Initial nutrient concentration (µg g⁻¹) of leaf litter, before placement in the field

<table>
<thead>
<tr>
<th>Leaf species</th>
<th>P (µg g⁻¹)</th>
<th>N (µg g⁻¹)</th>
<th>Ca (µg g⁻¹)</th>
<th>Mg (µg g⁻¹)</th>
<th>K (µg g⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>69 and 70</td>
<td>92 and 93</td>
<td>69 and 70</td>
<td>92 and 93</td>
<td>69 and 70</td>
</tr>
<tr>
<td>CO</td>
<td>1269</td>
<td>247*</td>
<td>7561</td>
<td>6366</td>
<td>10263</td>
</tr>
<tr>
<td></td>
<td>(26)</td>
<td>(427)</td>
<td>(438)</td>
<td>(1421)</td>
<td>(120)</td>
</tr>
<tr>
<td>D</td>
<td>1185</td>
<td>574*</td>
<td>10106</td>
<td>8431*</td>
<td>23800</td>
</tr>
<tr>
<td></td>
<td>(44)</td>
<td>(438)</td>
<td>(523)</td>
<td>(518)</td>
<td>(1421)</td>
</tr>
<tr>
<td>RM</td>
<td>1200</td>
<td>350*</td>
<td>6500</td>
<td>6086</td>
<td>11450</td>
</tr>
<tr>
<td></td>
<td>(27)</td>
<td>(523)</td>
<td>(523)</td>
<td>(518)</td>
<td>(1421)</td>
</tr>
<tr>
<td>WO</td>
<td>1179</td>
<td>337*</td>
<td>8049</td>
<td>7505</td>
<td>11683</td>
</tr>
<tr>
<td></td>
<td>(40)</td>
<td>(317)</td>
<td>(317)</td>
<td>(317)</td>
<td>(224)</td>
</tr>
<tr>
<td>WP</td>
<td>1572</td>
<td>216*</td>
<td>8727</td>
<td>6123*</td>
<td>6376</td>
</tr>
<tr>
<td></td>
<td>(5)</td>
<td>(143)</td>
<td>(143)</td>
<td>(143)</td>
<td>(72)</td>
</tr>
</tbody>
</table>

Values shown are 69 and 70 means of calculated values for all litterbags collected and 92 and 93 means of four initial litterbag samples selected at random. Values in parentheses are standard errors of the mean.

* Significant difference between 69 and 70 and 92 and 93 means based on a t-test (d.f. = 4; t = 2.132; p < 0.05).

DW had the greatest N and P concentration and the greatest rate of N and P release during decomposition. However, in 70s data the pattern did not hold. WP had the greatest and WO the least initial P concentration, yet WO had one of the greatest and WP the least P release rates during decomposition. Bockheim et al. (1991) examined release of 12 elements during decomposition of litter from one conifer and three deciduous species in northern Wisconsin. They found that all species tested showed increases in N, Ca, S, Zn, Mn, Fe, Cu, and A1 during the first year of decomposition; P, K, Mg, and B decreased. This differs from our 70s data, in which RM and WP retained N and only WP retained P. All other litter species released Ca, P, N, Mg, and K (Table 2). However, the 90s results were similar to Bockheim’s study, only DW released both N and P and WP had a N release rate of 0.11. Calcium, Mg, and K were released during the first year of decomposition for all species with few significant differences from 70s. Blair (1988b) studied changes in nutrient concentrations of D, RM, and CO during 2 years of decomposition on a south-facing reference watershed at Coweeta. Similarly, he found that only dogwood released P by the end of the study.

3.3. Microarthropods

Comparison of total microarthropod populations in 1992 showed no significant differences among litter species, but numbers differed significantly among collection months. In contrast, the 1993 data showed that total populations differed among both litter species and collection month. Mesostigmatid, mature oribatid, and immature oribatid mites differed significantly among litter species in 1992 (Table 4). There were also significant differences in mesostigmatid and immature oribatid mite abundances among

Table 4
Likelihood of significant differences in microarthropod populations among litter species

<table>
<thead>
<tr>
<th>Organism</th>
<th>Year</th>
<th>Log-likelihood</th>
<th>$\chi^2$</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prostigmata</td>
<td>1992a</td>
<td>123.62</td>
<td>22.16</td>
<td>0.0358</td>
</tr>
<tr>
<td>Prostigmata</td>
<td>1993</td>
<td>77.94</td>
<td>4.94</td>
<td>0.2933</td>
</tr>
<tr>
<td>Mesostigmata</td>
<td>1992</td>
<td>121.45</td>
<td>10.34</td>
<td>0.0376</td>
</tr>
<tr>
<td>Mesostigmata</td>
<td>1993</td>
<td>172.19</td>
<td>21.04</td>
<td>0.0003</td>
</tr>
<tr>
<td>Mature oribatids</td>
<td>1992</td>
<td>222.23</td>
<td>21.93</td>
<td>0.0002</td>
</tr>
<tr>
<td>Mature oribatids</td>
<td>1993</td>
<td>137.02</td>
<td>2.57</td>
<td>0.6317</td>
</tr>
<tr>
<td>Immature oribatids</td>
<td>1992</td>
<td>239.9</td>
<td>17.27</td>
<td>0.0017</td>
</tr>
<tr>
<td>Immature oribatids</td>
<td>1993</td>
<td>246.66</td>
<td>12.25</td>
<td>0.0156</td>
</tr>
<tr>
<td>Collembola</td>
<td>1992b</td>
<td>54.79</td>
<td>15.02</td>
<td>0.0018</td>
</tr>
<tr>
<td>Collembola</td>
<td>1993</td>
<td>200.52</td>
<td>4.19</td>
<td>0.3815</td>
</tr>
</tbody>
</table>

Data analyzed (using Proc GENMOD) were total numbers of microarthropods per three replicates for each litter species per collection date. Dates collected were September, October, November, and December in 1993 and September, October, and December in 1994. Litter species were red maple, dogwood, chestnut oak, white oak, and white pine (d.f. = 4).

* Not significant, because date × species was significant.

b Convergence not attained for at least one side of profile likelihood; many samples in October had no collembola in them.
litter species in the 1993 collections (Table 4). Populations of prostigmatid mites and Collembola showed no significant differences among litter species in either 1992 or 1993 (Table 4). On the other hand, Collembola was the only population of microarthropod that differed significantly between the 1992 and 1993 sample years (Fig. 4A). In 1992, greater numbers of mesostigmatid mites were found in RM and DW litter than in other species (Fig. 4B). Mature oribatid mite numbers were greatest in WP (Fig. 4D) and immature oribatids were most abundant in RM (Fig. 4C). In 1993, mesostigmatids were found in low numbers in WO and WP (Fig. 4B). Mature oribatids were most abundant in DW and WP litter, and immature oribatids were again most abundant in RM. Although these associations are interesting, their significance in relation to forest floor decomposition rates is tenuous. Forest floor litter in these mixed hardwood ecosystems is typically representative of species. Research has shown that microarthropods more effectively utilize mixed litter (Kaneko and Salamanca, 1999; Seastedt, 1984), probably because of increased habitat heterogeneity. This question deserves further study, as most quantitative litterbag decomposition studies continue to use a single species of litter with a few exceptions (Blair et al., 1990; Kaneko and Salamanca, 1999; Reynolds et al., 2003). We found no differences in Collembola populations among different litter species (Fig. 4A). This contradicts results reported by Reynolds (1976), who found Collembola abundances varied with litter species at Coweeta. However, those data were not collected over a full year and thus did not include fall and winter. Average numbers of Collembola per litter species in
1992 were less than half those of the 1993 collections (Fig. 4a), and many October 1992 samples had no Collombola. This may reflect the lack of moisture; summer 1992 had a 20-cm rainfall deficit (Fig. 2), and historically October has the lowest average rainfall of any month at Coveeta. Reynolds (1976) found a high correlation between precipitation in the 36-h period preceding litterbag collections and numbers of Collombola. Pflag and Wolters (2001) also reported decreased Collombola abundance in drought-stressed litter. Although Collombola numbers increased for our 1993 collections, we still had no significant differences in Collombola numbers among litter species. However, inspection of the data certainly indicate a trend similar to that reported by Reynolds (1976), with greater Collombola numbers occurring in more rapidly decomposing litter—RM and D—and fewest numbers in WO.

We explored the relationships between litter nutrient concentrations (total N, C:N ratios, P, K, Mg, and Ca) and microarthropod populations, using a stepwise regression analysis (Table 5). Generally, there were more significant relationships between microarthropod populations and litter nutrients in 1992, when moisture was more limiting than in 1993. Most correlations were negative, suggesting either that populations increased as decay proceeded, or that fungal populations, that support most microarthropods, decreased as nutrient concentrations increase. Collombola populations were correlated with nutrient concentrations of WO and WP only in 1992, while nutrient concentrations and oribatid mite populations were related in all litter species in 1992 (Table 5). For litter types most resistant to decomposition, CO, WO, and WP, populations were often related to C:N ratio and N. In the more readily decomposing DW and RM litters we found oribatid mites were related to P and Mg, respectively. The immature oribatids showed few significant relationships in either 1992 or 1993. The response difference between the two age classes may be related to avoidance of competition within the same taxon. The prostigmatids show nutrient relationships only in the litter species most resistant to decay—CO, WO, and WP. Relationships for other mite suborders and litter nutrients are difficult to interpret.

Seastedt (1984), surveyed microarthropod impacts on litter nutrient concentrations and found highly variable effects. Relationships depended on such

<table>
<thead>
<tr>
<th>Organism</th>
<th>Year</th>
<th>Leaf species</th>
<th>CO</th>
<th>D</th>
<th>RM</th>
<th>WO</th>
<th>WP</th>
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<tr>
<td>Collembola</td>
<td>1992</td>
<td></td>
<td></td>
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<td></td>
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<tr>
<td></td>
<td>1993</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oribatid mites</td>
<td>1992</td>
<td>C:N, 0.30 (0.01)</td>
<td>P, 0.36 (0.01)</td>
<td>Mg, 0.30 (0.02)</td>
<td>N (+), C:N (+), 0.90 (&lt;0.01)</td>
<td>C:N, N, 0.38 (0.01)</td>
<td>P (+), C:N, 0.94 (&lt;0.01)</td>
</tr>
<tr>
<td></td>
<td>1993</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Immature oribatids</td>
<td>1992</td>
<td>Ca, 0.21 (0.05)</td>
<td>Mg, 0.22 (0.04)</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>1993</td>
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</tr>
<tr>
<td>Prostigmatids</td>
<td>1992</td>
<td>CN (+), 0.21 (0.04)</td>
<td></td>
<td>K (+), N, C:N, 0.57 (&lt;0.01)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1993</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mesostigmatids</td>
<td>1992</td>
<td>P (+), C:N, Mg, N, 0.99 (&lt;0.01)</td>
<td></td>
<td>Mg (+), Ca, K (+), 0.95 (0.02)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1993</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total microarthropods</td>
<td>1992</td>
<td></td>
<td>Mg, 0.26 (0.03)</td>
<td>C:N, N, 0.37 (0.02)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1993</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.68 (0.02)</td>
</tr>
</tbody>
</table>

All data available were included in the models with n = 19 or 21 in 1992 and n = 7 for 1993. Shown are the nutrients included in the final stepwise model (variables with a positive relationship are designated (+) all others are negative), the final model $r^2$ and the probability of a value greater than $F$. Data shown are those in which a relationship was identified and the probability of a value greater than $F$ ≤ 0.05; other data are omitted.
factors as microbial affinities for particular nutrients and the form and amount of nutrients entering the litter from outside sources, such as throughfall. Given the wide variation in precipitation between the 2 years of our study, it is not surprising that the relationships between microarthropods and litter nutrients differed between the 2-years. Microbial biomass, including fungi, has been shown to significantly decrease with drought (Pflug and Wollers, 2001). Hasegawa and Takeda (1996) also reported drought in the first year of their study, with corresponding decreases in microarthropod populations during dry months. Most microarthropods feed on fungi; however, different collembolan species, may prefer different fungal species (Addison et al., 2003; Klironomos et al., 1992; Lussenhop, 1992). At our level of taxonomic resolution, we cannot tease apart some of the specific relations between particular microarthropod species and litter nutrients. This suggests that our population data may reflect a drought response with low numbers of Collembola in 1992 compared to 1993.

We found negative relationships between C:N ratios and Collembola, prostigmatids, mesostigmatids, and mature oribatids in both years. This is similar to results reported by Hasegawa and Takeda (1996) for pine needles with high C:N ratio and low litter quality. The C:N ratios decline during decomposition due to microbial respiration and N becomes increasingly available to the microbial biomass necessary to support microarthropods. The negative relationship between N concentrations and microarthropods in some instances is surprising. It may reflect the shift from N immobilization in earlier stages of decomposition to N release which could be conflated because of our lumping of the fall and early winter population data. Also, as Filser (2002) noted, correlations between Collembola and total C and N are usually weak under field conditions. They suggest that Collembola (and other microarthropods) responses to litter nutrients are indirect, because they are mediated by microbial population responses to litter nutrient concentrations and other parameters, such as temperature, moisture, litter species, and interactions with other soil biota.

Relationships between microarthropods and other nutrients (P, K, Mg, and Ca) were variable making any general conclusions impossible. The role of nutrients in litter colonization by microarthropods was reviewed by Seastedt (1984). He reported significant positive relationships between P and microarthropod populations in some studies and no relationship in others. He concluded that population control by K are rare because significant amounts of K can enter the litter system from precipitation and throughfall (Seastedt, 1984).

4. Conclusions

Data show no significant changes in first-year decomposition rates between the 70s and 90s studies. However, significant changes in initial litter nutrient concentrations and nutrient release patterns in the 90s suggest that long-term decomposition rates may be limited by nutrient availability. The long-term forest floor accumulation measured in a previous study could be the result of nutrient limitations. Microarthropod populations measured in the 90s varied with leaf species, resource availability, and rainfall. There were significant negative relationships between microarthropods and C:N in some litter species. Relationships differed between years, probably because of differences in precipitation. Microarthropod responses to nutrients are difficult to interpret because they are mostly indirect based on relationships with microbes, other soil biota, and numerous abiotic factors. However, data suggest that changes in forest floor nutrient content and nutrient release patterns may impact populations.

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References


