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Structure of a Microbial Community in Soil after Prolonged Addition of Low Levels of Simulated Acid Rain

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Humus samples were collected 12 growing seasons after the start of a simulated acid rain experiment situated in the subarctic environment. The acid rain was simulated with H2SO4, a combination of H2SO4 and HNO3, at two levels of moderate acidic loads close to the natural anthropogenic pollution levels of southern Scandinavia. The higher levels of acid applications resulted in acidification, as defined by humus chemistry. The concentrations of base cations decreased, while the concentrations of exchangeable H+, Al, and Fe increased. Humus pH decreased from 3.83 to 3.65. Basal respiration decreased with increasing humus pH, and total microbial biomass, measured by substrate-induced respiration and total amount of phospholipid fatty acids (PLFA), decreased slightly. An altered PLFA pattern indicated a change in the microbial community structure at the higher levels of acid applications. In general, branched fatty acids, typical of gram-positive bacteria, increased in the acid plots. PLFA analysis performed on the bacterial community growing on agar plates also showed that the relative amount of PLFA specific for gram-positive bacteria increased due to the acidification. The changed bacterial community was adapted to the more acidic environment in the acid-treated plots, even though bacterial growth rates, estimated by thymidine and leucine incorporation, decreased with pH. Fungal activity (measured as acetate incorporation into ergosterol) was not affected. This result indicates that bacteria were more affected than fungi by the acidification. The capacity of the bacterial community to utilize 95 different carbon sources was variable and only showed weak correlations to pH. Differences in the toxicities of H2SO4 and HNO3 for the microbial community were not found.

During the past few decades, the impact of acid deposition on the environment has been under intensive study. Changes in the soil chemical status and the function of the decomposer community may lead to imbalances in nutrient cycling and productivity of the ecosystem. Both direct and indirect soil-mediated effects of an increased acid load on the size, composition, and activity of the soil microbes have been reported (e.g., 19, 29), as have microbe-mediated changes in soil processes, such as litter decomposition (34).

Strong reductions in microbial activity will easily be detected if unrealistically high doses of acids are applied. The soil ecosystem may also be influenced by a subtle but permanent increase in the acidity of rain. However, to our knowledge no such studies exist. This acidification experiment in northern Finland is an attempt to study such subtle effects, since the annual sulfur and nitrogen loads applied are within the range of deposition over large areas in central Europe (25). The 12-year duration of the experiment is also important, since the impact of artificial acidification on soil microbes is suggested to depend on the duration of exposure to acid (26, 39) as well as on the rate of acid application (36).

Our main aim was to characterize the impact of artificial acidification on the humus microbial community structure by applying total-community-level techniques available now in microbial ecology. We used different fungus- and bacterium-specific measurements to study if acidification causes shifts in these main decomposer groups. We also used methods to determine microbial community structure in more detail without determining the exact species composition. One of these methods, the analysis of phospholipid fatty acids (PLFA), was shown earlier to be efficient in detecting the effects of increasing pH on the microbial community (9, 10, 22). Since the cultivable part of the bacterial community has been reported several times to be affected by acidification (e.g., 8, 38), we also analyzed the community structure of the cultivable bacteria by plate counting and subsequent PLFA analysis. Biolog GN microtiter plate incubation of the total humus suspension was performed to detect changes in the substrate utilization potential of the microbial community (24). The possible adaptation of the bacteria to the decreased humus pH was determined by the thymidine incorporation technique (6). We also compared bacterial activities with fungal activities to elucidate which of these organism groups might be mainly responsible for the previously detected decrease in soil respiration rate due to acidification in the study plots (39).

MATERIALS AND METHODS

Study area and sampling. The study area is situated near Kevo Subarctic Research Station in northern Finland, where the growing season is 110 to 125 days. The area is a dry, nutrient-poor, mixed pine-mountain birch woodland (Pinus sylvestris, Betula pubescens subsp. czerepanovii). The ground floor forms patches of dwarf shrub and lichen.

The study plots belong to a large-scale investigation in which the effects of acid rain on the subarctic ecosystem have been monitored since 1985 (30). Each plot (5 by 5 m) supported at least one pine and one mountain birch. The total number of plots used in the present study was 60, situated in three adjacent subareas. The treatments were dry control (D), irrigated control (W) treated with spring water (pH 6), and two levels of simulated acid rain, medium (m) and high (h). Originally (1985 to 1988), simulated acid rain was prepared by adding both H2SO4 and HNO3 (1.9:1 [wt/wt]) (SN subarea). In 1989, the treatments were modified as follows. In the SN subarea, the treatments continued unchanged (SN-m, pH 3.8; SN-h, pH 2.9), but in the S subarea only H2SO4 (S-m, pH 4.1; S-h, pH 3.1)
was applied and in the N subarea only HNO₃, (N-m, pH 4.7; N-h, pH 3.4) was applied. Plots were treated two or three times per week during June, July, and August. The cumulative S and N loads are shown in Table 1.

Humus samples were collected with a soil corer (2.8-cm diameter, 30 cores from each plot) in July 1995. Samples were sieved (2.8-mm mesh) and stored at 4°C for 2 weeks before basal respiration, substrate-induced respiration (SIR), Biolog GN microtiter plate (Biolog Inc., Hayward, Calif.), or PLFA analyses were performed. Subsamples for determination of the incorporation of [³H]thymidine, [¹⁴C]leucine, and [¹⁴C]acetate into ergosterol measurement, and plate counts were frozen (–18°C) for 5 months before the analyses. The samples were thawed and kept at room temperature for 3 to 7 days before the analyses.

Chemical analyses. Total organic carbon and nitrogen were determined by dry combustion (Leco CHN-600). For nutrient analyses, air-dried humus was extracted with 0.1 M BaCl₂ and the suspension was analyzed with an inductively coupled plasma emission spectrometer (ICP-AES, ARL 3580). The same suspension was also titrated with NaOH to measure exchangeable acidity. Effective cation-exchange capacity (CEC) was expressed as centimoles kilogram⁻¹, and base saturation (BS) was expressed as a percentage of Ca, Mg, K, and Na of CEC. Humus pH was measured in a water/1:1.7 [vol/vol] suspension. Dry weight was determined by drying duplicate subsamples at 105°C overnight, and organic matter content was obtained by heating at 550°C for 4 h. Detailed descriptions of the chemical extractions are given by Tamminen and Starr (37).

Microbiological analyses. The water content of fresh humus was adjusted to 60% of the water-holding capacity before respiration determinations. Basal respiration was then measured as CO₂ evolved in 39 h. SIR (2), which measures the potential biomass C of active microbes (Cmic) in soil, was determined as described by Priha and Smolander (35). Fresh humus, equaling 2 g (dry weight), with two replicates, was used in both analyses.

Phospholipid extraction and analysis of PLFA were done as previously described by Frostgård et al. (23). The total amount of PLFA was used to indicate the total microbial biomass. The sum of the PLFA considered to be predominantly of bacterial origin (i15:0, a15:0, 15:0, i16:0, 16:1ω9c, 16:1ω7t, 17:0, 17:0, 18:0ω7t, 18:0ω9t, and cy19:0) was chosen as an index of the bacterial biomass (21). The quantity of 18:ω6t,ω9t was used as an indicator of fungal biomass since 18:ω6t,ω9t has been suggested to be mainly of fungal origin in soil (18). It was found to correlate with the amount of ergosterol (21), a sterol found only in fungi.

Biolog GN microtiter plates containing 95 different carbon sources and a redox dye were incubated with the homogenized 10⁻²-diluted (0.9% NaCl) humus suspension. The plates were incubated at 20°C and read after 20, 46, 72, 92, 122, 140, and 164 h.

Bacterial growth rates were estimated by thymidine and leucine incorporation with the bacterial community extracted by homogenization-centrifugation as described by Blåth (4, 5). Two replicates were used for each humus sample.

The pH response of the bacterial community was measured, with some modifications, as described by Blåth (6). The pH of the extracted bacterial suspension was adjusted to approximately 3.8 with citrate-potassium phosphate buffer (final concentrations, 0.33 mM citric acid and 0.066 mM phosphate) or to 7.2 with potassium phosphate buffer (final concentration, 6.6 mM phosphate). The buffer concentrations were chosen in order to produce minimal inhibition of incorporation. The thymidine incorporation procedure was then performed as described above. The logarithm of the ratio of incorporation at pH 3.8 to that at pH 7.2 was used as an index of the bacterial community adaptation to pH, where a higher value indicates an increased tolerance for a lower pH (see reference 6 for further explanation).

Acetate incorporation into ergosterol (31) was used to estimate fungal growth rates. Humus suspensions (0.25 g [fresh weight] of humus and 1.5 ml of distilled water) were incubated for 18 h with labeled acetate (50 μl of [³H]acetate [2.11 GBq mmol⁻¹] and 450 μl of 1 mM nonradioactive acetate) at 20°C. Incubation was stopped with 2 ml of 5% formalin. The humus slurry was centrifuged, and the supernatant was discarded. Ergosterol was then extracted and hydrolyzed according to Ek et al. (16), except that cyclohexane was used and the hydrolysate time was 1 h. Ergosterol was analyzed by high-performance liquid chromatography. Ergosterol peaks were collected into scintillation vials. Relative (¹⁴C)acetate incorporation into ergosterol was expressed as disintegrations per minute per microgram of ergosterol.

The same homogenized bacterial suspension that was used for thymidine incorporation was used for plate counts. Suspensions were diluted with 0.1% peptone and spread on tryptone soy agar plates. Plates were incubated at room temperature for 4 weeks, and the number of CFU, both total and yellow colonies, was calculated from plates with less than 80 CFU. Agar plates on which the number of CFU was high (about 100 to 300) were then flooded with citrate buffer, and a 1.5-ml portion of the bacterial suspension was taken for further analysis of PLFA as described above (CFU-PLFA).

Statistical analyses. The results are expressed per organic matter content, since Vanhala et al. (39) found the organic C content of humus to be an important source of variation in the subarea plots. SN subarea N had a lower cumulative acid load than subareas SN and S (Table 1), and the density of the natural vegetation was higher in subarea N than in the other two subareas. Therefore, the different treatments (D, W, m, and h) were not quite equivalent in the different subareas, and a two-factor analysis of variance (ANCOVA) (including treatment and subarea) was not used. Instead, all of the measured variables, including the scores in the multivariate analysis (PLFA, CFU-PLFA, and Biolog measurements), were subjected to an analysis of covariance (ANCOVA). The analysis was performed with treatment and subarea as classifying variables and humus pH as the covariant. The interaction between subarea and treatment was also tested. The Newman-Keuls comparison of means was then performed if those sources of variation were found to be significant. A significant covariant revealed if the variable was related to pH and thus to acidification. A significant treatment effect indicated impacts which were due to something other than pH, e.g., the effect of watering. A significant subarea effect indicated a different response due to the composition of the acid. This kind of model thus enabled the testing of all 60 plots with the same analysis and allowed a comparison of the S and N subarea acids as well. Most of the results are presented by plotting all 60 study plots against humus pH. Variables which showed significant treatment effects are presented as an average for the five replicates of the treatments with standard error bars.

The moles percent of the individual PLFA were standardized by dividing them with their standard deviations before being subjected to principal-component analyses (PCA). Partial least-squares (PLS) regression analysis (Unscrambler II program) was used to compare the total humus PLFA and plated-community PLFA patterns. Calculation and standardization of the Biolog data on the basis of the reading time of the plates were performed following Garland (24). Reading times at which average well color development was between 0.7 and 1.0 were then selected for the statistical analysis. Absorptions of the separate wells were divided by the mean absorbance of the plate, and the results were subjected to PCA.

The study plots, although situated in a dry forest site, could be divided according to their original ground vegetation and moisture into three classes: dry, medium, and moist plots. The effect of vegetation was not included in the ANCOVA because of the uneven distribution of the vegetation within the treatment and also because it was already partly included in the model since it affected the soil pH. A separate ANOVA with vegetation, treatment, and subarea as sources of variation was, however, used to determine the reasons for the formation of principal components of the PLFA data.

### RESULTS

Chemical analyses. The results of the chemical analyses are presented in Table 2. At both the SN-h and the S-h plots, the pH of the humus layer decreased from 3.83 (mean for W plots) to 3.65 (mean for h plots). The N-h treatment, which had a considerably smaller proton load than the SN-h or S-h treatments and therefore a higher pH of the irrigation water, did not affect the humus pH. The concentrations of Ca, K, and Mg decreased with decreasing pH, leading to decreased BS and slightly lower CEC. The exchangeable acidity and the concentrations of Al and Fe increased due to acid treatments, but the trend was not significant in an ANCOVA model when pH was used as a covariant.

Microbial biomass and activity. The total biomass of active microorganisms measured by SIR (Fig. 1a) and the total amount of microbe-derived PLFA (data not shown) were found to be dependent on the pH (P levels of the covariant pH were <0.001 and <0.01, respectively). SIR and total PLFA were positively correlated (r, 0.621). Actual reduction in the biomass due to the irrigation treatments was very small; e.g., the mean SIR biomass was 14.1 mg of Corg g of organic matter⁻¹ and the total PLFA biomass was 2.6 μmol g of organic matter⁻¹ at SN-W plots, while the same values at SN-h plots

<table>
<thead>
<tr>
<th>TABLE 1. Cumulative acid loads of the Kevo study area</th>
<th>Treatment</th>
<th>Cumulative acid load, 1985–1995 (g/m²)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>S</td>
</tr>
<tr>
<td>W</td>
<td>0.93</td>
<td>0.17</td>
</tr>
<tr>
<td>SN-m</td>
<td>4.40</td>
<td>1.38</td>
</tr>
<tr>
<td>SN-h</td>
<td>22.1</td>
<td>7.59</td>
</tr>
<tr>
<td>S-m</td>
<td>4.40</td>
<td>0.60</td>
</tr>
<tr>
<td>S-h</td>
<td>22.1</td>
<td>2.79</td>
</tr>
<tr>
<td>N-m</td>
<td>2.13</td>
<td>1.32</td>
</tr>
<tr>
<td>N-h</td>
<td>8.38</td>
<td>7.25</td>
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were 13.1 mg of C\textsubscript{mic} g of organic matter\textsuperscript{−1} and 2.4 \textmu mol g of organic matter\textsuperscript{−1}, respectively. Basal respiration rate was affected by both the humus pH and watering treatment (Fig. 1b). Respiration rate decreased (\(P < 0.001\)) with decreasing pH but increased (\(P < 0.05\) for the treatment effect) with watering treatment. The changes in respiration rate were larger than the changes in biomass values, being 17.5, 12.7, and 14.5 \textmu g of CO\textsubscript{2}-C g of organic matter\textsuperscript{−1} at SN-W, SN-D, and SN-h plots, respectively. The lower respiration rate of D plots (situated in dry and medium vegetation types) is also shown in Fig. 1c, representing the driest vegetation type had relatively low humus pH and low respiration rate levels.

Microbial community structure measured by PLFA. PCA of PLFA showed separation of high-acidity-level irrigated samples from the SN and S plots along the second principal component (PC 2) (Fig. 2a). When the scores of that component were subjected to ANCOVA, they were significantly affected by humus pH (\(P < 0.001\)). The individual PLFA responsible for the separation were 10Me16, i16:1, i16:0, and 18:1\textsubscript{m}, respectively. Basal respiration rate was affected by both the humus pH and watering treatment (Fig. 1b). Respiration rate decreased (\(P < 0.001\)) with decreasing pH but increased (\(P < 0.05\) for the treatment effect) with watering treatment. The changes in respiration rate were larger than the changes in biomass values, being 17.5, 12.7, and 14.5 \textmu g of CO\textsubscript{2}-C g of organic matter\textsuperscript{−1} at SN-W, SN-D, and SN-h plots, respectively. The lower respiration rate of D plots (situated in dry and medium vegetation types) is also shown in Fig. 1c, representing the driest vegetation type had relatively low humus pH and low respiration rate levels.

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matrix and plated-community PLFA data (CFU-PLFA) were used as the y matrix. As a result of the PLS regression, the score plot separated the acidified plots along the first PLS regression component (data not shown). PLS regression also created loading values for individual PLFA. The loading values for the first PLS regression component of the humus PLFA matrix are shown in the x axis, and the loading values for the first PLS regression component of the CFU-PLFA matrix are shown in the y axis (Fig. 2d). Especially i16:0, i16:1, and 10Me16 increased in both analyses, while cy17:0, cy19:0, 16:1v7c, and 18:1v7 decreased in both analyses. The changes in 15:0, 16:0, 17:0, 18:0, i14:0, and i17:0 did not follow this pattern.

**Substrate utilization potential.** Study plots treated with high-level acid load were slightly separated along PC 2 (Fig. 2c). Scores of PC 1 and PC 2 were tested with ANCOVA, and the results showed them both to be slightly affected by soil pH ($P < 0.05$), indicating that there were some changes in substrate utilization due to humus pH. However, PC 1 mainly separated the D plots ($P < 0.01$ for the treatment effect). This finding, together with the fact that the two principal components explained only 20% of the variation in the Biolog GN data, makes the interpretation of the results difficult.

**Fungal biomass and growth rate.** Fungal biomass, measured both with fungal PLFA 18:2v6,9 and with ergosterol (data not shown), remained relatively unchanged during the experiment. The fungal growth rate was estimated by $[^{14}C]$acetate incorporation into ergosterol (Fig. 1d). The changing pH did not have a significant impact on the incorporation rate, although the highest values were usually found at lower pHs.

### DISCUSSION

According to the soil acidification hypothesis, which includes the loss of base cations and increases in exchangeable H$^+$, Al, and Fe, possibly resulting in a lowered pH (37), the humus was acidified as a result of the acid treatments (Table 2). A small decrease in the humus pH at the higher acid load was also seen even though the natural variation in pH was large compared to the effect of the treatments.
Anderson and Domsch (3) found the total microbial biomass and respiration rate to be lower in naturally acidic soils than in soils with a neutral pH. In our study, the basal respiration rate (Fig. 1b) and to some extent the total biomass (Fig. 1a) were also dependent on pH and therefore on acidification, even though the pH range studied by us was much smaller. Respiration rate was also affected by irrigation (see results from the ANCOVA) and vegetation type (Fig. 1c). It is likely that the lack of watering was the reason for the lower respiration rate in the D plots, since increased moisture has been reported several times to raise microbial activity (40, 42). The lowest respiration rates were found at the plots situated in the dry vegetation type, irrespective of the treatment (Fig. 1c). Humus pH at the dry vegetation type tended to be lower than that at the other vegetation types, indicating that the lower respiration rate at the dry vegetation type might have been due to an originally lower pH at those areas. This lower pH might also be the reason why the dry vegetation type was more sensitive to the effects of acidification (30).

The analysis of humus PLFA revealed changes in microbial community structure due to pH (Fig. 2a). The individual PLFA more typical of the acid-treated plots separated along PC 2 and were considered to be of bacterial origin. Most of them, e.g., branched PLFA i16:0, i16:1, 10Me16, and 10Me17, are common in gram-positive bacteria (32). The differences in moles percent were, however, small. PLFA of eucaryotic origin, e.g., 18:2ν6,9 and 20:4 (1, 21), did not respond to pH but separated to the left along PC 1, the vegetation axis, indicating the connection of the fungal biomass to different vegetation types. An altered humus bacterial community composition due to pH was also supported by the PLFA pattern of the cultivable bacterial population (Fig. 2b), since the S or SN plots treated with higher acid levels (in addition to some W plots) separated in the PCA. The increase in relative moles percent of branched PLFA in low-pH humus was larger but was much more variable in this analysis than in the humus PLFA analysis. PLS regression of the plated and direct humus PLFA analyses showed similarities between PLFA affected by pH (Fig. 2d).
Most of the PLFA increased (i16:0, i16:1, and 10Me16) or decreased (cy17:0, cy19:0, 16:1ω7c, and 18:1ω7) with decreasing humus pH in both analyses. Some differences were also seen; for example, PLFA i14:0 and i17:0 increased in acidified plots when the cultivable community was analyzed and decreased in the same plots in the humus PLFA analysis. This result might indicate that some parts of the bacterial population are excluded or enriched on the agar plates. The differences in saturated straight-chain PLFA were probably due to these PLFA being present in eucaryotes in soil. As a whole, it appears that plate counts were not solely a result of selection by the plating technique, since the bacteria responsible for the greatest changes in bacterial PLFA due to acidification were probably able to grow on agar media.

In addition to the changes in humus and plated-community PLFA patterns, the proportion of yellow colonies among total CFU increased with decreasing humus pH (Fig. 3d), indicating an altered bacterial community composition. The percentage of yellow colonies has been reported to decrease with lime and ash treatments of coniferous forest soil resulting in a higher pH (7). An increase in coniferous forest humus pH has been reported to change the microbial community measured by PLFA analysis toward more gram-negative and fewer gram-positive bacteria (9, 10, 22). The abundance of PLFA common in gram-negative bacteria (cy17:0, 18:1ω7, 16:1ω5, and 16:1ω7c) (41) increased with increasing pH (22), while in our study these PLFA were more typical in the control areas with a higher pH. On the other hand, the levels of most of the fatty acids more abundant at the control areas of the liming treatments (i15:0, i16:0, and 10Me16) were found to be increased due to acidification. Consequently, coniferous forest humus seems to contain a bacterial group, consisting mainly of gram-positive bacteria, which easily adapts to an acid environment, and a group of bacteria, mainly gram-negative ones, which more easily adapts to humus with a more neutral pH. However, whether this situation represents a direct pH effect or an effect of pH altering carbon availability and thus selecting for more active bacteria, as suggested earlier (10), cannot be elucidated from the present results.

The potential of the bacterial community to degrade different carbon sources appeared to be only slightly affected by pH (Fig. 2c). However, the suitability or sensitivity of the Biolog method for bulk humus samples might be questioned, since reductions in the number of bacteria utilizing starch, protein,
pectin, xylan, and cellulose after 4 years of irrigation in these same study plots were found (27).

When the bacterial community incorporated thymidine in a solution in which pH was decreased or increased with buffers, the community of the plots having a lower humus pH showed greater adaptation to the more acidic environment than did the community of the plots having a higher humus pH (Fig. 3c). However, the bacterial growth rate in general, as indicated by the thymidine incorporation rate (Fig. 3b), decreased due to acidification in spite of the fact that at least part of the bacterial community was adapted to the altered environment. Since changes in the PLFA pattern and percentage of yellow colonies indicated a change in bacterial community structure, it is likely that the increased acid tolerance of the bacterial community resulted at least partly from a shift in species composition.

Fungal biomass appeared not to suffer from acidification to the same extent as bacterial biomass, since the amounts of both fungal PLFA 18:2ω6,9 and ergosterol were unaffected by the treatments. Thus, it is likely that the decrease in total activity (basal respiration) was due to pH affecting mainly the bacterial community. These results are therefore in accordance with several previous reports about unchanged (11) or even increased (28) fungal biomass or a higher fungal/bacterial biomass ratio (13) caused by acidification. On the contrary, results showing decreased fungal biomass (17, 20) or decreased fluorescein diacetate-active fungi (8, 33) also have been reported, but these results are mainly from experiments with high acid treatments. Thus, it is likely that the decrease in total activity (basal respiration) was due to pH affecting mainly the bacterial community structure, it is likely that the increased acid tolerance of the bacterial community resulted at least partly from a shift in species composition.

Instead of direct toxicity of protons or anions, reduction in the availability of carbon for microbes has been suggested to be the main reason for the adverse effects of acidification (15, 33). Reduced substrate availability of heterotrophs may thus determine microbial activity; this situation may be evidenced as lowered respiration rate or bacterial activity, as shown in the present study. Another reason for the changes in the microbial community may be the effects of acid on the plants. The same factors that influence plant growth, e.g., environmental stress, may affect plant root exudation (14). Bäck et al. (12) reported reduced branch and needle growth of the pines of these Kevo study plots as a consequence of a decline in photosynthetic activity caused by irrigation of S and SN plots. However, since we did not find any differences among S, SN, and N plots that could not be attributed to an altered pH, we suggest that the acidification effect on the microorganisms was not mediated through the plants.

The present study showed that even though the northern European woodlands are naturally very acidic, the present low-level but prolonged acid rain resulted in clear signs of acidification, as defined by soil chemical variables. Litter decomposition rate, the ultimate sign of disturbances in the decomposition process, had earlier been shown to be decreased in these study plots (30), yet only marginal effects on total microbial biomass and basal respiration rate were found. The community-level techniques, however, were sensitive enough to reveal disturbances in the structure and general activity of the bacterial community, even though at least some of the bacteria were found to be adapted to an altered humus pH. Fungal biomass and activity were not affected or were affected only to a minor extent. This result could, of course, be due to the more specific character of bacterial methods (PLFA, thymidine incorporation, and Biolog analyses), while fungal community structure was not analyzed to the same extent. However, at least the total biomass and the growth rate of the fungi were unchanged, while those of the bacteria were reduced.

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