A correlative study was performed to determine if variation in streambed microbial community structure in low-order forested streams can be directly or indirectly linked to the chemical nature of the parental bedrock of the environments through which the streams flow. Total microbial and photosynthetic biomass (phospholipid phosphate [PLP] and chlorophyll a), community structure (phospholipid fatty acid analysis), and physical and chemical parameters were measured in six streams, three located in sandstone and three in limestone regions of the Bankhead National Forest in northern Alabama. Although stream water flowing through the two different bedrock types differed significantly in chemical composition, there were no significant differences in total microbial and photosynthetic biomass in the sediments. In contrast, sedimentary microbial community structure differed between the bedrock types and was significantly correlated with stream water ion concentrations. A pattern of seasonal variation in microbial community structure was also observed. Further statistical analysis indicated dissolved organic matter (DOM) quality, which was previously shown to be influenced by geological variation, correlated with variation in bacterial community structure. These results indicate that the geology of underlying bedrock influences benthic microbial communities directly via changes in water chemistry and also indirectly via stream water DOM quality.

Microorganisms are one of the most important groups of living organisms. Their small size, ubiquitous distribution, metabolic diversity, and genetic plasticity cast microorganisms in the role of recycling agents for the biosphere, making life possible for more complex organisms (47). To better understand the ecology of microbial communities, it is important not only to describe the community composition but also to identify biological and/or environmental factors that regulate their diversity (34, 52).

Baas-Becking’s dictum that “everything is everywhere, but the environment selects” has inspired many studies and debates since its inception and is considered the precursor to the niche concept adopted by macroecologists (10). To better understand the ecology of microbial communities, it is important not only to identify community composition but also to identify biological and/or environmental factors that regulate their diversity (34, 52).

Our study focused on bedrock, a most basic component of the environment, and its influence on microbial community structure in stream sediments, as stream ecosystems provide critical links in global hydrological and biogeochemical cycles and are now viewed as transformative rather than transportive ecosystems (2).

The geochemical setting influences several key environmental determinants and determines the availability of resources that can be physiologically exploited by microorganisms. It is these interactions between microorganisms and their resources that most likely contribute to metabolic diversity (43). Several studies have identified environmental factors that structure sedimentary microbial communities in streams, including light, flow, water temperature, organic carbon concentrations, anthropogenic pollution, hydrology, pH, sediment grain size, inorganic nutrients, and dissolved oxygen (1, 6, 9, 19, 30, 55, 59, 63), but the structuring role of geological formation is unknown.

The linkage between geology and water chemistry in streams is due to minerals released by the weathering of parent bedrock. The input of dissolved chemicals will directly affect pH, alkalinity, and ion concentrations and can potentially change the proportion of essential nutrients available to organisms (44, 67). Studies addressing the relationship between sedimentary microbial communities and water chemistry have focused on pH, nutrients, or anthropogenic influences (3, 4, 20, 46, 53). It is evident that differences in water chemistry influence microbial community structure, but to date natural variation in cation/anion concentrations remains to be investigated. This study’s primary objective was to determine if patterns in sedimentary microbial community structure in forested low-order streams is influenced by geological differences in stream channel bedrock.

Mosher et al. (49) showed that dissolved organic matter...
(DOM) quality was correlated to geological variation in the same streams used for this study. Since previous studies have shown bacterial uptake and growth efficiency are dependent of the source of DOM (37, 39) and DOM quality can influence bacterial community composition (36), this study investigated a possible indirect linkage of geology and microbial community structure via DOM quality. To accomplish this second objective, factor scores from a principal component analysis describing the variation in DOM quality in the six streams (49) were included in the environmental descriptions used in this study.

**MATERIALS AND METHODS**

**Site description.** The study area was in the William B. Bankhead National Forest, in northern Alabama directly south of the Tennessee Valley Divide. All streams are second- or third-order tributaries to the Sipsey Fork of the Black Warrior River system in the Mobile River drainage basin. All streams have similar elevations, discharge, and sediment grain size and are located within 21 km of each other in the north central portion of the forest. The creek beds consist of exposed bedrock, cobble, and unconsolidated sediment.

The forest is located in the Appalachian Plateau physiographic province, and the primary bedrock within the region is Pennsylvanian-age Pottsville formation comprised of sandstone/shale with thin, discontinuous layers of coal (11). Some valleys and streambeds in this region have eroded through the Pottsville bedrock to older Mississippian-age bedrock, specifically Parkwood sandstone and Bangor limestone formations. Two of the streams (Beech and Brushy Creeks: 34°17′49.85″N, 87°18′44.03″W and 34°17′58.8″N, 87°16′29.41″W, see reference 49 for geologic map of study site) flow over exposed Pottsville formation bedrock, characterized by shale/shale and clayey coal. Three streams (Thompson, Flannagin, and Borden Creeks: 34°20′26.89″N, 87°28′15.82″W; 34°20′19.79″N, 87°23′18.31″W; 34°19′42.53″N, 87°22′30.15″W, respectively) have eroded through both the Pottsville and Parkwood bedrock and flow over exposed Bangor limestone. Only one stream in this study (Hubbard Creek: 34°18′30.22″N, 87°30′0.38″W) flows solely over the overlying Parkwood sandstone formation. The streams that have eroded through the upper layer(s) still contain Pottsville formation sandstone in the upland portion of their watershed (45). The catchments are heavily forested, primarily by mixed hardwoods interspersed with pine stands.

**Sampling.** Sampling was conducted on a quarterly basis for 1.25 years under base flow conditions, at least 2 weeks following the last recorded precipitation in the area. Sediment from each stream was sampled in triplicate with a push core. The top 1 cm of sediment of each core was homogenized, and subsamples were taken for each measurement. Stream water was collected in duplicate, and physical parameters were measured in situ with portable meters. All samples were placed on ice and processed immediately upon return to the laboratory; some samples were preserved (frozen [sediments for lipid analysis] and/or acidified [water for DOC analysis]) and stored prior to analysis (16).

**Physical parameters.** Total suspended solids (TSS) were determined by filtering well-mixed stream water samples through dried, preweighed glass fiber filters (GF/F) and drying them at 103°C to a constant mass (14). Specific surface area of combusted sediments was analyzed by the absorption isotherm of nitrogen under vacuum (7). Average weekly discharge in the streams was estimated by a rating curve constructed from instantaneous measurements of discharge and the placement of barologgers in each stream (Levelogger 3001, Solinst Canada Ltd., Georgetown, ON, Canada). The rating curve was constructed from monthly (n = 15) measurements of stream velocity (Marsh McBirney, Frederick, MD), depth measurements, and width. The average weekly gauge height obtained from the barologgers was applied to the rating curve, and discharge for the week preceding the date of sampling was estimated. Canopy cover was estimated using a hand-held convex spherical crown densimeter (Forestry Suppliers, Jackson, MS).

**Chemical parameters.** Stream water samples were taken using high-density polyethylene (HDPE) bottles rinsed with a sulfuric acid-noroxin mixture and washed with phosphate-free soap. The first 20 ml of sample was filtered over a combusted GF/F and acidified with two drops of 2 N HCl and dissolved organic carbon (DOC) concentration was analyzed by flash combustion in a Shimadzu total organic carbon analyzer TOC-500. The remainder of the stream water was filtered through the same GF/F and analyzed for NO3-N, NO2-N, NH4-N, and PO4-P concentrations in a Lachat QuickChem 8000.

Water samples for anions (Cl− and SO42−) and cations (Al3+, Ba2+, Ca2+, Fe3+, and Mg2+) [hereafter Fe], Na+, Ca2+, Mg2+, and Mn2+[hereafter Mn], SiO2− and SiO42− [hereafter Si], and K+] were collected in HDPE bottles previously washed with ultrapure water (>18.0 MQ) and filtered over a combusted GF/F filter. Anion samples were stored at −20°C, and cation samples were acidified (2% [vol/vol]) with optimum-grade nitric acid until analysis. Samples for anion concentrations were analyzed using ion chromatography, and the cation concentration samples were subjected to inductively coupled plasma spectrometry.

Acid-neutralizing capacity (ANC) of the stream water was estimated by the Gran titration method; pH was determined after incremental (0.025-mL) additions of 0.1 N HCl to 60 mL stream water while stirring until full protonation had occurred. The Gran function was calculated using the following equation (68):

\[ F_1 = V_{equil} + V[H^+] \]

where \( V \) is the volume and \([H^+]\) is the hydrogen ion concentration.

The equivalence point was found by plotting \( F_1 \) versus acid volume. Dissolved inorganic carbon (DIC) was calculated from the ANC by using the following equation (68):

\[ \text{DIC} = ([\text{Alk}]) - [OH^-] + [H^+]\left(\epsilon_1 + 2\epsilon_2\right) \]

where Alk is alkalinity and \( \epsilon_1 \) and \( \epsilon_2 \) are the decimal percentages of carbonate and bicarbonate ions, respectively, in the total DIC.

Conductivity (Fishier Scientific, Waltham, MA), pH (Oakton Instruments, Vernon Hills, IL), dissolved oxygen (DO), and temperature (model 95, YSI, Yellow Springs, OH) were measured in situ using hand-held meters.

**Microbial parameters.** Microbial biomass and microbial community structure were characterized by phospholipid phosphate (PLP) and phospholipid fatty acid (PLFA) analyses. Cellular lipids were extracted from the sediments by a modified (dichloromethane-methanol-water) Bligh-Dyer lipid extraction (5, 26). Total microbial biomass was determined by the oxidation of lipids and quantification of the resulting orthophosphate colorimetrically (26). Fractionation of phospholipid fatty acid composition into fatty acid methyl esters (FAMEs) using allylation allowed identification of functional groups comprising the microbial community structure. Purified FAMEs were separated and quantified using a gas chromatograph (23). FAMEs were identified by coelution with known standards and mass spectral analysis. A fraction of total lipids, protected from light, was used to determine chlorophyll a colorimetrically. Dried samples were suspended in 90% aqueous acetone, and absorbance was measured at 663, 645, and 630 nm. Chlorophyll a abundance was calculated using the following equation (62):

\[ \text{chlorophyll a} = 11.85(\text{OD}_{663}) - 1.54(\text{OD}_{645}) + 0.08(\text{OD}_{630}) \]

where OD663, for example, represents the optical density at 663 nm.

Percent contributions of eukaryotes and prokaryotes to the microbial community biomass were determined using the approach described by Findlay and Dobbs (24), and it was assumed that 50% of the PLFAs from eukaryotes were polyenoic (25, 69) and eukaryotic biomass was the sum of the polyenoic PLFAs × 2. Prokaryotic biomass was calculated as the difference between the total microbial biomass and eukaryotic biomass. Eukaryotic biomass in terms of C was calculated from PLP concentration × the percent eukaryotic biomass (expressed as a decimal fraction) × 0.02 g C µmol P−1 (24). Prokaryotic biomass in terms of C was calculated as the PLP concentration × the percent prokaryotic biomass (expressed as a decimal fraction) × 0.01 g C µmol P−1 (24).

**Statistical analyses.** Fully nested analysis of variance (ANOVA) with Tukey’s honest significant difference test (HSD) was performed for all chemical, and biological parameter to determine significant differences of the parameters by substrate type (limestone versus sandstone) and sampling date (Minitab 14.13). Pearson correlation coefficient was used to determine colinearity among the environmental parameters (SPSS 14.0).

Constrained ordination techniques were utilized to identify patterns of variation in microbial community structure among streams and correlations between microbial community structure and environmental descriptors. Detrended correspondence analysis (DCA), an indirect gradient analysis based on segment length, was performed to determine the modality of the PLFA data and environmental predictor variables. The analysis resulted in short (<1.0) segment lengths, indicating the data set was linear and suitable for indirect gradient analysis; therefore, redundancy analysis (RDA) was applied (Canoco 4.5). In the RDA, the response variables were individual PLFAs (transformed using In[weight percentPLFA + 1]), and the predictor variables were measured environmental parameters. Forward selection of the predictor variables followed by Monte Carlo permutation tests were used to prevent artificial inflation of variation due to autocorrelation in the constrained ordination model (42). A second RDA, limited to data collected during the March 2006 sampling, describing the influence of DOM quality on sedimentary microbial community structure was performed and included factor scores (factors 1 and 2) from a principal component analysis that examined variation in DOM quality of the six streams, DOM was sampled concurrently with the March 2006 sampling, and the analysis of the influence of bedrock geology on dissolved organic matter quality has been published elsewhere (49).
TABLE 1. Summary of chemical and environmental characteristics of the six study streams located in the Bankhead National Forest, AL*

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Borden</th>
<th>Flannagain</th>
<th>Thompson</th>
<th>Beech</th>
<th>Brushy</th>
<th>Hubbard</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al³⁺ (mg/liter)</td>
<td>0.04 ± 0.03</td>
<td>0.04 ± 0.03</td>
<td>0.05 ± 0.04</td>
<td>0.08 ± 0.04</td>
<td>0.05 ± 0.02</td>
<td>0.06 ± 0.02</td>
</tr>
<tr>
<td>ANC (mg/liter)</td>
<td>2.17 ± 0.49</td>
<td>2.31 ± 0.39</td>
<td>0.90 ± 0.34</td>
<td>0.25 ± 0.12</td>
<td>0.14 ± 0.07</td>
<td>0.12 ± 0.05</td>
</tr>
<tr>
<td>Ba²⁺ (mg/liter)</td>
<td>0.03 ± 0.00</td>
<td>0.03 ± 0.01</td>
<td>0.02 ± 0.00</td>
<td>0.03 ± 0.02</td>
<td>0.02 ± 0.00</td>
<td>0.02 ± 0.00</td>
</tr>
<tr>
<td>Ca²⁺ (mg/liter)</td>
<td>41.24 ± 8.16</td>
<td>42.27 ± 8.76</td>
<td>14.05 ± 5.32</td>
<td>2.60 ± 0.79</td>
<td>1.50 ± 0.48</td>
<td>1.35 ± 0.11</td>
</tr>
<tr>
<td>Canopy (%)</td>
<td>61.83 ± 33.40</td>
<td>44.38 ± 41.68</td>
<td>40.27 ± 34.85</td>
<td>66.99 ± 31.25</td>
<td>96.53 ± 3.69</td>
<td>69.11 ± 26.66</td>
</tr>
<tr>
<td>Cl⁻ (mg/liter)</td>
<td>1.42 ± 0.28</td>
<td>1.28 ± 0.28</td>
<td>1.11 ± 0.13</td>
<td>1.33 ± 0.23</td>
<td>1.09 ± 0.15</td>
<td>1.34 ± 0.28</td>
</tr>
<tr>
<td>Conductance (µS/m)</td>
<td>207.09 ± 46.55</td>
<td>223.39 ± 46.35</td>
<td>77.80 ± 29.77</td>
<td>31.72 ± 5.83</td>
<td>23.47 ± 5.07</td>
<td>20.75 ± 1.56</td>
</tr>
<tr>
<td>DIC (mg/liter)</td>
<td>0.53 ± 0.12</td>
<td>0.56 ± 0.09</td>
<td>0.37 ± 0.31</td>
<td>0.17 ± 0.13</td>
<td>0.14 ± 0.14</td>
<td>0.23 ± 0.13</td>
</tr>
<tr>
<td>Discharge (m³/s)</td>
<td>10.88 ± 3.99</td>
<td>11.65 ± 4.34</td>
<td>9.94 ± 3.62</td>
<td>10.52 ± 3.90</td>
<td>10.54 ± 4.42</td>
<td>10.40 ± 3.36</td>
</tr>
<tr>
<td>DO (mg/liter)</td>
<td>0.12 ± 0.06</td>
<td>1.17 ± 0.37</td>
<td>1.64 ± 1.02</td>
<td>1.33 ± 0.14</td>
<td>1.39 ± 0.50</td>
<td>1.08 ± 0.18</td>
</tr>
<tr>
<td>DOC (mg/liter)</td>
<td>0.08 ± 0.02</td>
<td>0.05 ± 0.02</td>
<td>0.05 ± 0.03</td>
<td>0.50 ± 0.36</td>
<td>0.38 ± 0.31</td>
<td>0.18 ± 0.09</td>
</tr>
<tr>
<td>Fe (mg/liter)</td>
<td>0.86 ± 0.25</td>
<td>0.71 ± 0.25</td>
<td>0.85 ± 0.16</td>
<td>0.80 ± 0.21</td>
<td>0.71 ± 0.19</td>
<td>0.83 ± 0.20</td>
</tr>
<tr>
<td>K⁺ (mg/liter)</td>
<td>14.50</td>
<td>13.50</td>
<td>6.00</td>
<td>20.00</td>
<td>23.00</td>
<td>2.50</td>
</tr>
<tr>
<td>Mg²⁺ (mg/liter)</td>
<td>0.61</td>
<td>1.07</td>
<td>0.36</td>
<td>0.60</td>
<td>0.31</td>
<td>0.32</td>
</tr>
<tr>
<td>Mn (mg/liter)</td>
<td>2.16 ± 0.42</td>
<td>2.45 ± 0.56</td>
<td>1.44 ± 0.31</td>
<td>1.26 ± 0.20</td>
<td>1.07 ± 0.26</td>
<td>0.80 ± 0.07</td>
</tr>
<tr>
<td>Na⁺ (mg/liter)</td>
<td>0.23 ± 0.01</td>
<td>0.01 ± 0.01</td>
<td>0.01 ± 0.00</td>
<td>0.06 ± 0.03</td>
<td>0.01 ± 0.01</td>
<td>0.01 ± 0.01</td>
</tr>
<tr>
<td>NH₄-N (µg/liter)</td>
<td>1.36 ± 0.14</td>
<td>1.20 ± 0.10</td>
<td>1.20 ± 0.16</td>
<td>1.16 ± 0.15</td>
<td>1.03 ± 0.15</td>
<td>0.89 ± 0.12</td>
</tr>
<tr>
<td>NO₃-N (µg/liter)</td>
<td>2.95 ± 2.45</td>
<td>5.80 ± 6.15</td>
<td>2.52 ± 2.20</td>
<td>13.97 ± 19.41</td>
<td>3.39 ± 2.11</td>
<td>3.03 ± 2.75</td>
</tr>
<tr>
<td>NO₂⁻ (µg/liter)</td>
<td>17.07 ± 16.53</td>
<td>20.22 ± 17.27</td>
<td>42.94 ± 21.62</td>
<td>99.27 ± 41.86</td>
<td>33.33 ± 41.33</td>
<td>190.62 ± 71.51</td>
</tr>
<tr>
<td>pH</td>
<td>0.11 ± 0.13</td>
<td>0.19 ± 0.29</td>
<td>0.16 ± 0.27</td>
<td>0.54 ± 0.78</td>
<td>0.36 ± 0.60</td>
<td>0.31 ± 0.32</td>
</tr>
<tr>
<td>PO₄-P (µg/liter)</td>
<td>8.09 ± 0.16</td>
<td>8.17 ± 0.66</td>
<td>7.73 ± 0.13</td>
<td>7.20 ± 0.19</td>
<td>7.17 ± 0.13</td>
<td>7.24 ± 0.24</td>
</tr>
<tr>
<td>Position* (km)</td>
<td>0.42 ± 0.65</td>
<td>0.36 ± 0.56</td>
<td>0.92 ± 1.04</td>
<td>0.45 ± 0.49</td>
<td>0.26 ± 0.36</td>
<td>0.43 ± 0.42</td>
</tr>
<tr>
<td>Si (mg/liter)</td>
<td>2.66 ± 0.27</td>
<td>2.62 ± 0.24</td>
<td>2.87 ± 0.25</td>
<td>3.00 ± 4.42</td>
<td>3.09 ± 0.46</td>
<td>2.65 ± 0.34</td>
</tr>
<tr>
<td>SO₄²⁻ (mg/liter)</td>
<td>3.94 ± 0.81</td>
<td>5.50 ± 1.44</td>
<td>4.30 ± 0.14</td>
<td>2.96 ± 0.45</td>
<td>3.44 ± 0.31</td>
<td>2.52 ± 0.71</td>
</tr>
<tr>
<td>Surface area (m²/g of sediment⁻¹)</td>
<td>1.79 ± 0.28</td>
<td>1.86 ± 0.81</td>
<td>2.88 ± 0.61</td>
<td>1.07 ± 0.77</td>
<td>0.31 ± 0.35</td>
<td>1.64 ± 2.21</td>
</tr>
<tr>
<td>Temp (°C)</td>
<td>15.05 ± 6.27</td>
<td>14.54 ± 6.48</td>
<td>16.75 ± 5.79</td>
<td>15.59 ± 6.72</td>
<td>15.08 ± 7.16</td>
<td>14.79 ± 6.09</td>
</tr>
<tr>
<td>TSS (mg/liter)</td>
<td>1.59 ± 1.10</td>
<td>3.62 ± 3.07</td>
<td>1.37 ± 0.82</td>
<td>10.80 ± 9.97</td>
<td>3.23 ± 3.14</td>
<td>1.58 ± 0.42</td>
</tr>
</tbody>
</table>

* Adapted from reference 49.
** Significance of differences between substrate type (limestone versus sandstone) as determined by fully nested ANOVA. Abbreviations: NA, statistics not applied; NS, no significant difference.

RESULTS

Physical, chemical, and biological factors. The water chemistry of the streams reflected the nature of the bedrock over which the streams flowed. Limestone streams had significantly higher values for ANC, pH, conductivity, and concentrations of Ca²⁺, DIC, Mg²⁺, Na⁺, and SO₄²⁻ than those of the sandstone streams (Table 1). One of the limestone streams, Thompson Creek, had significantly lower values for these parameters than the other two limestone streams, but the values were significantly higher than those observed in the sandstone streams. Water samples taken from sandstone streams had significantly higher concentrations of NO₃-N and Fe than the limestone streams. The differences found in these chemical parameters, with the exception of NO₂⁻, were in accordance with previous published studies examining differences in ion concentrations in relation to geology. No significant differences were found between the remainder of the chemical parameters (DO and concentrations of Ba²⁺, Cl⁻, DOC, K⁺, Mn, NH₄-N, and Si) or any of the physical parameters (percent canopy cover, discharge, temperature, and TSS).

Total microbial and autotrophic biomass. There were no significant differences between substrate types or among sampling dates for total microbial biomass (Fig. 1). Total microbial biomass values ranged from 1.5 to 100 ng PLP g of sediment⁻¹ and the limestone streams averaged 17.5 ng PLP g of sediment⁻¹, while the sandstone streams averaged 21.34 ng PLP g of sediment⁻¹. There were also no significant differences for chlorophyll a concentrations between the substrate types or among sampling dates (Fig. 2). The values for chlorophyll a ranged from 1 to 101 µg g of sediment⁻¹, with a noticeable increase in concentrations for the December sampling date in both substrates. The average concentration of chlorophyll a in limestone streams was 6.23 µg g of sediment⁻¹ and 12.1 µg g of sediment⁻¹. There were also no significant differences for chlorophyll a concentrations between the substrate types or among sampling dates (Fig. 2).
of sediment\(^{-1}\) in sandstone streams. There was a positive correlation between chlorophyll \(a\) concentration and total microbial biomass (\(r = 0.673\)), and none of the environmental variables correlated with total microbial biomass or chlorophyll \(a\) concentration.

**Microbial community structure.** Microbial community structure displayed two distinct overlying patterns as indicated by the triplot of samples, environmental variables and individual fatty acid markers (Fig. 3). Starting at the top left of the triplot and moving toward the bottom right, samples were aligned by increasing stream water ion concentrations, which were significantly associated with Ca\(^{2+}\) and DOC concentrations (Table 2). Starting at the top right of the triplot and moving toward the bottom left, the samples were aligned by increasing water temperature and were significantly associated with water temperature and sampling date (\(F = 8.71; P = 0.002\)). RDA canonical axes 1 and 2 described 28.6\% of the variation in microbial community structure.

The samples grouped into three clusters in the triplot. Samples taken in March were clustered together and contained two subgroups. Samples from the sandstone streams and the lowest-conductivity limestone stream (Thompson Creek) formed one subgroup within the cluster, and the other subgroup contained Borden and Flannagin Creeks. The March samples were significantly associated with fatty acids found in autotrophic eukaryotes (16:4\(_1\), 18:4\(_3\), 16:3, and 20:5\(_3\)). The second cluster consisted of non-March sandstone streams and was described by bacterial fatty acid markers (18:1\(_9\), br19:1a, br19:1b, and 17:0). The third cluster was comprised of non-March Borden and Flannagin Creeks (the two highest-conductivity limestone streams) and was described by autotrophic (16:1\(_{13t}\)) and heterotrophic (20:4\(_6\)) eukaryotic markers. Thompson Creek results plotted either with the sandstone streams or between the non-March clusters.

Notably, the significant fatty acid markers describing the variation among the clusters were differences in eukaryotic and bacterial fatty acids. Samples having positive RDA axis 1 scores were characterized by a significant presence of eukaryotic fatty acids, while the samples having negative RDA axis 1 scores were characterized by bacterial fatty acids. A significantly higher percentage of eukaryotes were found in the samples taken in March (\(P = 0.002\)) (Fig. 4). Removal of eukaryotic microorganisms from the streambed sediments was significantly greater for the March sampling date (fully nested ANOVA with Tukey's honestly significant difference, \(P < 0.05\)).

**TABLE 2.** Correlation matrices (\(r\)) of environmental variables from the constrained ordination analysis (RDA) performed on PLFA profiles and environmental variables from the six streams in the Bankhead National Forest

<table>
<thead>
<tr>
<th>Factor</th>
<th>RDA 1</th>
<th>RDA 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ca(^{2+}) concn</td>
<td>0.4177</td>
<td>-0.5782</td>
</tr>
<tr>
<td>Sampling date</td>
<td>-0.3838</td>
<td>-0.3666</td>
</tr>
<tr>
<td>DOC concn</td>
<td>-0.0761</td>
<td>0.2609</td>
</tr>
<tr>
<td>Temp</td>
<td>-0.2204</td>
<td>-0.2404</td>
</tr>
</tbody>
</table>

**FIG. 2.** Chlorophyll \(a\) concentrations of streambed sediments by sampling date from six streams (three limestone, three sandstone) located in the Bankhead National Forest. Error bars represent 1 standard deviation.

**FIG. 3.** Triplot of the redundancy analysis of PLFA profiles of stream bed sediments of six streams (three limestone, three sandstone) located in the Bankhead National Forest, with forward selection of predictor variables followed by Monte Carlo permutations. Solid arrows represent predictor variables significantly associated with variation in microbial community structure. Dashed arrows represent individual fatty acid markers (\(r > 0.65\)) significantly associated with the variation among samples. The lengths of the arrows are correlated with the degree of relation between the response variable (exact values are shown in Table 2). The arrows point in the direction of the maximum change for the associated variable. Legend: closed symbols, limestone streams; open symbols, sandstone streams; stippled symbols, intermediate limestone (Thompson Creek); circles, March; squares, June; diamonds, September; triangles, December.

**FIG. 4.** Percentage of total microbial biomass that was comprised of microeukaryotes, by date in the streambed sediments of six streams (three limestone, three sandstone) located in the Bankhead National Forest. Error bars represent 1 standard deviation. *, significantly greater eukaryotic contribution to total biomass for the March sampling date (fully nested ANOVA with Tukey's honestly significant difference, \(P < 0.05\)).
otic fatty acids from the response variable matrix allowed examination of bacterial community response and showed a clear clustering of samples along the RDA axis 1, with samples from limestone streams having positive scores and those from sandstone streams with negative scores ($F = 7.28; P = 0.002$). Again, samples taken from Thompson Creek were either grouped with the sandstone streams or in between the sandstone and other limestone streams (Fig. 5).

ANC was the environmental parameter most closely associated with the variation describing limestone streams along the RDA axis 1, and DOC concentrations and sampling date described the variation on RDA axis 2 (Table 3). Individual fatty acids describing the variation in this analysis were a mixture of branched monoenoic and saturated fatty acids.

Influence of DOM quality on bacterial community structure. The number, nature, and relative abundance of molecules that comprise stream water DOM (here referred to as DOM quality) from the six streams, sampled March 2006, was significantly influenced by stream water chemistry and the presence of bituminous coal beds within the watershed (49). These data were added to the March 2006 description of the environment, and RDA was performed to determine if DOM quality contributed to the variation in bacterial community structure.

RDA axes 1 and 2 described 55.9% of the variation (Fig. 6). Stream water ion composition ($Na^+$ and $Fe^{2+}$ concentration) and DOM quality were significantly associated with the variation (Table 4) ($F = 3.31; P = 0.004$). The samples plotted according to stream water ion concentrations. The sandstone samples, containing lower ion concentrations, had negative RDA factor scores, and the samples from two of the limestone streams (Borden and Flannagin Creeks), containing higher stream water ion concentrations, had positive RDA factor scores. Thompson Creek, with intermediate stream water ion concentrations, had values near zero and the results were between the sandstone and the two limestone streams that had higher stream water ion concentrations. There was a mixture of individual fatty acids significantly associated with the variation, the majority being indicative of anaerobic and/or Gram-positive bacteria.

**DISCUSSION**

Total microbial and autotrophic biomass levels. No significant differences were observed in total microbial biomass and chlorophyll $a$ concentrations between the two substrate types.

<table>
<thead>
<tr>
<th>TABLE 3. Correlation matrices ($r$) of environmental variables from the constrained ordination analysis (RDA) performed on bacterial PLFA profiles and environmental variables from the six streams in the Bankhead National Forest</th>
</tr>
</thead>
<tbody>
<tr>
<td>Factor</td>
</tr>
<tr>
<td>ANC</td>
</tr>
<tr>
<td>DOC concn</td>
</tr>
<tr>
<td>Sampling date</td>
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</tbody>
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<table>
<thead>
<tr>
<th>TABLE 4. Correlation matrices ($r$) of environmental variables from the constrained ordination analysis (RDA) performed on bacterial PLFA profiles and environmental variables including DOM quality from the six streams in the Bankhead National Forest on March 2006</th>
</tr>
</thead>
<tbody>
<tr>
<td>Factor</td>
</tr>
<tr>
<td>DOM quality</td>
</tr>
<tr>
<td>$Fe$ concn</td>
</tr>
<tr>
<td>$Na^+$ concn</td>
</tr>
</tbody>
</table>
Microbial biomass was comparable to reports from previous studies of streambed sediments (6, 12, 63), and chlorophyll a concentrations were also within ranges previously reported (31, 58). These previous studies found environmental and physical factors that correlated with microbial and autotrophic biomass levels. For instance, variation in sediment particle size (59) and DOC concentrations (6) have been found to be influential on microbial biomass, while inorganic nutrients (41) and light attenuation (17) have been associated with variation in chlorophyll a concentrations. With this knowledge, 6 streams were chosen within a single watershed that had similar physical and environmental characteristics, with the exception of bedrock geology, to eliminate potential confounding factors and to maximize the ability to detect the influence of geological formations. We conclude that there was no direct influence of geology on microbial or autotrophic biomass in streambed sediments stemming from variations in stream water ion concentrations.

**Microbial community structure.** In contrast to microbial and autotrophic biomass, the variation described in streambed microbial community structure was associated with geological formations, specifically, stream water ion concentrations, DOM quality, and sampling date. Streams flowing over sandstone bedrock consistently showed lower pH and cation concentrations, while streams flowing over limestone bedrock showed higher pH and cation concentrations. Previously, Findlay et al. (28) showed differences in microbial community structure in stream sediments at the biome level; while they noted differences in stream water chemistry among biomes they did not determine proximate causes contributing to the variations in microbial community structure. The current study describes variation in stream water chemistry and streambed microbial community structure of six streams within one biome. Again, by confining this study to a single forest, many physical and environmental parameters that contribute to variations in microbial community structure were minimized, allowing the demonstration of the role of geologic bedrock in structuring microbial communities.

One parameter not controlled for in the study design was the presence of geologically influenced macroinvertebrate grazers. Reproduction and growth of pleurocerid snails (*Elimia carineferia*) are dependent on sufficient concentrations of stream water ions, specifically Ca$^{2+}$, and therefore these organisms were only found in the limestone streams of this study (35). In a related study, an enclosure/exposure experiment was conducted in two of the study streams, one limestone (Flannagin Creek) and one sandstone (Brushy Creek) stream, to determine if any of the variation observed in microbial community structure could be attributed to the effects of macroinvertebrate grazing. No significant differences in total microbial biomass, chlorophyll a biomass, or microbial community structure for any of the treatments were detected, thus eliminating grazing as a geology-induced effect (48).

Seasonal variation in the microbial community structure was observed in both stream types. Remarkably, while distinct differences in sediment microbial communities occurred between stream types, the shifts in the microbial community structure throughout seasonal changes were similar. Eukaryotic biomarkers showed increased relative abundances in all samples taken in March. Previous studies have shown seasonal variation in microbial community structure caused by differing eukaryotic and bacterial contribution to total biomass. Most of these studies found that microeukaryotes were more abundant and responsible for describing the variation in microbial communities in colder months (marine sediments [27], reservoir sediments [61], and creosote-impacted stream sediments [40]). One study found a higher relative abundance of eukaryotes in sediments of a regulated stream in warmer months (63). In a comparison of streams within and among biomes, sampled to avoid any seasonal effects, Findlay et al. (28) also found the major component of variation in microbial community structure was caused by differing eukaryotic and bacterial contributions to total biomass. Variation occurred among biomes, among streams with a biome, and for one biome, within streams. Interestingly, the second component of variation in microbial community structure in the Findlay et al. (28) study appeared to be associated with stream water ion concentrations/geology. In our current study, the importance of eukaryotes versus bacteria was evident both seasonally and by bedrock type, with the variation in microbial community structure in the limestone streams characterized by higher relative abundances of microeukaryotic fatty acid markers, while bacterial fatty acids were significant in describing the variation in the sandstone streams. When eukaryotic biomarkers were removed from the analysis, the effects of geologically driven water chemistry parameters and seasonal variations were still significant, and DOC concentration became a significant determinant of bacterial community structure.

DOM in stream ecosystems is an essential component for many geochemical processes and is a major energy source for microbe-based food webs (66). DOM quantity and quality are important in shaping aquatic bacterial community structure and metabolism (15, 29). Judd et al. (36) cross-fed bacterial communities from stream and lake sediments with DOM from stream, lake, or soil water and found that the bacterial community shifted to reflect the source from which the DOM originated. An experiment manipulating DOM quality and quantity sources in planktonic bacterial communities from a lake and two streams reported similar results (13). While these previous studies showed evidence that the source of DOM is important for bacterial assemblages, none characterized DOM quality at the molecular level, nor did any of the studies identify factors influencing DOM quality. Mosher et al. (49) successfully accomplished both of these by using ultra-high-resolution mass spectrometry to determine DOM quality in the stream waters of Hubbard, Thompson, Flannagin, Borden, Beech, and Brushy creeks. The water in the sandstone streams was characterized by higher abundances of condensed hydrocarbons, while the limestone stream water contained more oxygenated molecules. Further, Mosher et al. (49) determined that geological variation, including the presence of coal, was the most influential factor behind the variation in DOM quality. The inclusion of the DOM quality data from Mosher et al. (49) demonstrated that DOM quality was a significant factor (along with the direct effects of bedrock type) behind the variations in bacterial community structure. This was the first study to show evidence of the influence of geological variation and DOM quality on bacterial and microbial community structures in stream sediments. In addition, this correlative experimental design clearly demonstrated the influence of geological
variation on DOM quality (49); however, it was unable to differentiate the effects of variation in microbial community structure on stream water DOM and vice versa. Stream microorganisms, via DOM processing, alter DOM quality (38), and it is reasonable to expect that different microbial communities alter DOM quality to a lesser or greater extent. Nonetheless, this study provides further support for the second half of Baas-Becking’s “everything is everywhere, but the environment selects” dictum. This support should be tempered by the understanding that while PLFA analysis produces quantitative data, unlike DNA-based methods (50), it is a phenotypic method (see reference 23 for a full discussion of the advantages and disadvantages of the PLFA approach). It should be noted that when PLFA and molecular methods are applied in tandem, they produce very similar analyses of microbial community structure (28, 32). These results definitively demonstrated that DOM quality and bedrock composition, via the influence on stream water chemistry, influenced microbial community structure in streambed sediments. While a complex relationship between geology, microbial community structure, and DOM quality may exist, it is clear that geologically mediated stream water ion concentrations and DOM quality selected for sedimentary microbial community structure and that bedrock type within fluvial networks should be added to the growing list of environmental determinants of microbial distribution and abundance.

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