The response of Chironomidae (Diptera) to a long-term exclusion of terrestrial organic matter

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Abstract We examined the effects of a seven-year detrital exclusion on chironomid assemblages in an Appalachian headwater stream. We hypothesized that litter exclusion would lead to a reduction in all chironomids at both the subfamily and generic levels because organic matter serves as both food and habitat in these headwater streams. Tanytarsini total abundance and biomass significantly declined after litter exclusion. Before litter exclusion, Tanytarsini average abundance was 4271 ± 1135 S.E. m⁻² and 625 ± 98 after litter exclusion. Biomass was 3.57 ± 0.96 mg AFDM m⁻² before litter exclusion and 1.03 ± 0.9 after exclusion. In contrast, Orthocladiinae abundance and biomass did not change because a psammanophilic chironomid, Lopescladius sp., and other Orthocladiinae genera did not decline significantly. Overall chironomid taxa richness and diversity did not change as a result of litter exclusion. However, Canonical Correspondence Analysis (CCA) of genus-level biomass did show a clear separation between the litter exclusion stream and a reference stream. Separation of taxa between the two streams was due to differences in fine ($r^2 = 0.39$) and coarse ($r^2 = 0.36$) organic matter standing stocks and the proportion of small inorganic substrates ($r^2 = 0.39$) present within a sample. As organic matter declined in the litter exclusion stream, overall chironomid biomass declined and the chironomid community assemblage changed. Tanytarsini were replaced by Orthocladiinae in the litter exclusion stream because they were better able to live and feed on biofilm associated with inorganic substrates.

Keywords Benthos • Chironomidae • Detritus • Resource limitation • Bottom-up regulation

Introduction

Chironomidae are often the most productive primary consumers in an ecosystem; they represent an important link between basal food resources (e.g., algae, fungi, and leaf litter) and predators in freshwater habitats (Stites & Benke, 1989; Leeper & Taylor, 1998; Ramírez & Pringle, 1998). The success of chironomids is a result of their diverse physiological adaptations that allow
them to live in hot, cold, and low-oxygen environments. Chironomidae often have high densities, fast growth, and small size resulting in high colonization rates and production (Berg & Helenthal, 1992; Tokeshi, 1995; Huryn & Wallace, 2000).

Studies have shown that Chironomidae in forested streams depend on terrestrially derived detritus in the form of coarse (CPOM) and fine particulate organic matter (FPOM) as habitat and food (Huryn, 1990; Richardson, 1991; Grubbs & Cummins, 1994; Berg, 1995), in addition to dissolved organic matter taken up by microbes on organic and inorganic substrates (Hall & Meyer, 1998; Benke, 1998). But other studies have also shown Chironomidae production to be limited by food and habitat. For example, Johnson et al. (2003) found that growth rates of collector-gathering chironomids significantly declined in response to litter exclusion in a temperate headwater stream. Chironomids were also limited by nutrients in other aquatic ecosystems. For example, Chaloner et al. (2004) found that chironomid biomass increased, indirectly, in response to nutrients delivered from the ocean to nutrient-limited freshwater streams by salmon spawning upstream. However, these ecosystem studies placed chironomids into two groups based on their general feeding modes, the predaceous Tanypodinae and non-Tanypodinae collector-gatherers.

Other studies have evaluated mechanisms controlling chironomid assemblages at the generic and species levels (Wiley, 1981; Ladle et al., 1984; Ruse & Davidson, 2000; Silver et al., 2004) and they showed that important ecological information may be lost by using broad groupings. For example, Wiley (1981) experimentally determined that both habitat stability and competition among psammomophilic chironomids determined the benthic community structure. Furthermore, Hershey (1985) described the Chironomidae community in an Artic lake and identified stratification of specific genera based on distribution of macrophytes and soft sediments. Our goal was to evaluate possible changes in the chironomid community assemblage in response to an ecosystem-level exclusion of leaf litter and wood that decoupled the riparian and stream ecosystems without altering the riparian habitat.

We hypothesized that the exclusion of allochthonous organic matter could lead to a change in the Chironomidae assemblage. In a previous litter exclusion study, Wallace et al. (1997, 1999) found that long-term litter exclusion (i.e., 4 years) from a headwater stream resulted in a significant decline in biomass and abundance of most non-Tanypodinae chironomids (76% decline). We expanded on this study by looking at chironomids at a finer taxonomic level and through three additional years of litter exclusion. We examined the chironomid change in assemblage to a long-term litter exclusion and subsequent removal and exclusion of small and large wood in a headwater stream. Our objectives were to describe changes in taxa assemblage at the subfamily and generic levels before and after litter exclusion and to identify temporal changes in chironomid assemblage and biomass as a result of litter exclusion.

**Methods**

**Study sites**

A seven-year litter exclusion experiment (LE) was conducted in a first-order forested, headwater stream (C55) at Coweeta Hydrologic Laboratory (Macon Co., N.C.) in the southern Appalachians. For more information about the study sites see Wallace et al. (1999). A similar stream nearby was used as a reference (C53). The riparian canopy was similar in both streams and was composed primarily of tulip poplar (*Liriodendron tulipifera* L.), white oak (*Quercus rubra* L.), and dogwood (*Cornus florida* L.). *Rhododendron (Rhododendron maximum* L.) was the primary understory species and provided dense shade all year, which reduced in-stream primary production to very low levels. Both streams were similar with respect to their climate and physical parameters (Table 1); however, the litter exclusion stream had a higher percentage of pebble and sand. In August 1993, a 2.5-cm gill mesh net was built over the first 170 m of stream to exclude direct leaf fall and a 20-cm high fence of plastic mesh was erected on the
stream banks to exclude lateral litter blow-in. After three years of litter exclusion, small and large wood was removed and excluded (LE + WR, see Wallace et al., 2001). As a result of the litter exclusions, organic matter standing stock was reduced in the stream by ~95% compared to pre-litter exclusion values (Wallace et al., 1997, 1999; Eggert & Wallace, 2003b).

Chironomidae sampling

Samples were collected monthly from 1992 through October 2001. For each sample date, a 400-cm² stovepipe corer was used to collect four benthic samples, composed primarily of mixed substrate (sand, silt, cobble, pebble, gravel). All material was collected, placed on ice, and taken back to the laboratory for processing. At the time of collection, a visual assessment of the benthic substrate was also recorded as percent sand, gravel, pebble, silt and cobble. In the laboratory, organic matter was separated into two size fractions: >1 mm and <1 mm >250 µm. Individual size fractions were placed in plastic bags, dyed with Phloxin B, and preserved with 6–8% formalin. The formalin fixed samples were picked for invertebrates under a 15x dissecting microscope. All invertebrates were picked from the >1-mm samples. If more than 100 individual invertebrates were present in the 250-µm samples, they were subsampled using a sample splitter (Waters, 1969; Lugthart & Wallace, 1992).

Chironomid larvae were separated from the other invertebrate taxa, identified to genus, body lengths measured to the nearest 1 mm, and counted. Length–mass regressions were then applied to estimate biomass (Benke et al., 1999). Chironomids were identified from January, April, July, and October samples, each month representing the winter, spring, summer, or autumn season, respectively. We examined chironomids from 1992, 1993, 1995, 1998, and 2000. Chironomids were sorted under a dissecting microscope into like-groups and all individuals were mounted, unless the totals were >20, then 50% were mounted and identified. If different taxa were found mounted within like-groups, taxa were further separated and mounted. All chironomids were mounted using CMC-10 and identified to genus using Wiederholm (1983), Coffman & Ferrington (1996), and Epler (2001).

Data analysis

All abundance and biomass data were log_{10} transformed to meet the assumptions of ANOVA. We compared total differences in Chironomidae abundance and biomass over time by subtracting the mean of the reference (C53) from the mean of the Litter Exclusion (C55) stream for each sampling date. Zero indicated no difference in abundance or biomass of chironomids between the two streams. Positive numbers indicated higher chironomid abundance or biomass in the LE stream. Negative numbers indicated higher chironomid abundance or biomass in the reference stream. Least-squares regression was used to examine the change in treatment versus control.

We used a Before-After-Control-Intervention-Paired (BACIP) design (Osenberg et al., 1994; Smith, 2002) to detect litter exclusion effects, and to compare changes in chironomid abundance and biomass between the reference and litter exclusion streams. A two-way ANOVA with litter exclusion effects (i.e., pre-litter exclusion (P), litter excluded (LE), and litter and wood excluded (LE + WR)) plus control (C53) versus Intervention (C55) represented our class level variables. Because there were an uneven number of observations for each litter exclusion period (i.e., P: n = 4, LE: n = 7, and LE + WR: n = 10),
a General Linear Model was used (SAS v. 8.02, SAS Institute Inc., Cary, North Carolina). Only if a significant interaction between the Before-After and Control-Intervention occurred was a litter exclusion effect inferred.

Shannon–Weiner Index (SWI) values, represented as $H' = -\sum p_i \log p_i$ where $p_i$ represents proportion of total number of individuals in the $i$th species, values were compared using BACIP design and differences were detected using Tukey's multiple comparison test. Generic differences in abundance and biomass were examined using the BACIP factorial ANOVA. Least square means percent differences were used to detect paired differences in litter exclusion periods and stream, but a Tukey–Kramer correction was used to correct for multiple comparisons.

Genus-level biomass dynamics for each sampling period were explored using Canonical Correspondence Analysis (CCA, PC-ORD Version 4, MJM Software Design, Glenden Beach, OR, USA: http://home.centurytel.net/~mjm/pcordwin.htm). CCA was used to relate measured environmental variables to the seasonal distribution of chironomid biomass at the genus level (PC-ORD, McCune, 1997). We used ordination to graphically explore the changes in chironomid community biomass in response to LE. $\log_{10}$ transformed generic mean biomass was used to create a matrix for all taxa represented in at least 5% of the total samples. A secondary matrix of environmental variables (e.g., CPOM, FBOM, and % sand, gravel, and pebble, maximum stream discharge, and mean water temperature) were correlated with each axis and graphed as vectors. The closer a vector was to an axis, the greater the correlation to that axis. A longer vector represented a stronger correlation between that environmental variable and chironomid biomass within a stream and season.

**Results**

**Chironomidae abundance, biomass, and diversity**

Total chironomid abundance (Fig. 1a) and biomass (Fig. 1b) became progressively lower in the LE compared to the reference streams over the study period. Taxon richness and Shannon–Weiner Index values did not decline in response to LE or LE + WR (Fig. 2a, b). A total of 35 genera were identified in the litter exclusion stream before and 43 genera were identified after LE + WR. In the reference stream, 43 genera were identified. Chironominae, Orthocladiinae, Tanypodinae, Diamesinae, and Podinominae were identified, but because Diamesinae and Podinominae were so rare, their responses are not discussed.

**Chironomidae response**

Chironomini were neither abundant nor dominant in either stream (Fig. 3, Table 2). In the litter exclusion stream, Chironomini (predominantly Polypedilum sp., Dicrotendipes sp., and Steno-
Fig. 2 Species richness (a) over the study period and (b) Shannon Weiner Index of Diversity of the chironomid assemblages in the treatment and reference streams. Lines (solid = reference and dashed = treatment) represent the average for each treatment period (PRE pretreatment, LE litter exclusion, LE + WR wood removal).

*chironomus* sp.) abundance significantly declined following LE + WR ($F = 5.17, P < 0.001$, Fig. 3), but not LE alone ($P > 0.05$). This decline could not be attributed to litter exclusion as Chironomini abundance also declined in the reference stream (i.e., no significant interaction). There was also no significant decline of Chironomini biomass after LE or LE + WR and variability among seasons was high within the tribe and among genera.

Before litter exclusion, Tanytarsini (predominantly *Cladotanytarsus*, sp., *Constempellina* sp., *Micropsectra* sp., *Rheotanytarsus* sp., *Stempellinella* sp., and *Tanytarsus* sp.) composed over 60% of the chironomid assemblage in the litter exclusion stream (Table 2). Following LE and LE + WR, Tanytarsini abundance ($F = 5.17$, $P = 0.0002$) and biomass declined significantly ($F = 33.9$, $P < 0.0001$, Fig. 3). Overall, Tanytarsini abundance and biomass declined by 85% after litter exclusion and wood removal. However, at the genus level, only *Constempellina* sp. biomass declined significantly following both litter and wood exclusion ($F = 12.76, P < 0.001$). *Cladotanytarsus*, sp., *Constempellina* sp., *Micropsectra* sp., *Rheotanytarsus* sp., *Stempellinella* sp., and *Tanytarsus* sp. were also most abundant in the
Fig. 3 Log mean abundance and biomass of major Chironomidae tribes Chironomini and Tanytarsini, and subfamilies Orthocladiinae and Tanypodinae in the treatment and reference streams. Lines represent the average for each treatment period (PRE pretreatment, LE litter exclusion, and LE + WR wood removal). Different letters indicate significant differences between treatment periods, taking into account the difference in the values in the reference stream using BACIP.

In contrast to Tanytarsini, Orthocladiinae abundance and biomass did not significantly change in the litter exclusion stream (Fig. 3). Before litter exclusion, Orthocladiinae (predominantly Lopescladius sp., Parachaetocladius sp., and Synorthocladius sp.) were 24% of the chironomid assemblage (Table 2). Following LE and WR, Orthocladiinae were 73% of the total assemblage but total abundance and biomass did not show a significant increase. Abundance was 1817 ± 1135 before litter exclusion and 3271 ± 221 following LE + WR. Biomass was 6.67 ± 1.54 before litter exclusion and 11.54 ± 3.15 after LE and WR. Genus level abundance and biomass did not change and the Orthocladiinae assemblage showed less seasonal variation in the reference stream.

Predaceous Tanypodinae represented 4% of the taxa in the litter exclusion stream, with no change in relative abundance, absolute abundance, or biomass in either the litter exclusion or reference streams over time (Fig. 3, Table 2).

Ordination results: organic matter dynamics and seasonal changes in chironomid community assemblage and biomass

Ordination of mean generic biomass of the most abundant taxa in each year, over seven years, showed a clear separation between the reference and litter exclusion streams (Fig. 4). Axis 1 described 42% and axis 2 described 16% of the variation among samples. Fine ($r^2 = 0.36$) and coarse ($r^2 = 0.39$) organic matter was positively related to axis 1. The proportion of sand, gravel, and pebbles ($r^2 = 0.39$) was negatively related to axis 1. Axis 2 accounted for little variation among samples (16%). Mean monthly temperature ($r^2 = 0.17$) and mean monthly discharge ($r^2 = 0.15$) were weakly related to axis 2 (Table 3). Separation between the two streams corresponded to changes in the reference stream. Abundance, biomass, and relative abundance either increased or did not change significantly through time in the reference stream (Fig. 3).
Table 2  Average biomass (mg AFDM m⁻², ± standard error) of most abundant (>5% of total abundance) chironomid taxa within each stream and treatment period

<table>
<thead>
<tr>
<th>Taxa</th>
<th>Reference</th>
<th>LE</th>
<th>LE + WR</th>
<th>Litter exclusion</th>
<th>Pre</th>
<th>LE</th>
<th>LE + WR</th>
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<td><strong>Chironomini</strong></td>
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<tr>
<td><em>Demicryptochironomus</em></td>
<td>0.05 ± 0.02</td>
<td>0.70 ± 0.65</td>
<td>0.41 ± 0.20</td>
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<td><em>Microtendipes</em></td>
<td>0.55 ± 0.47</td>
<td>0.34 ± 0.34</td>
<td>15.53 ± 14.73</td>
<td>0.13 ± 0.12</td>
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<td><em>Polyplecton</em></td>
<td>1.34 ± 0.98</td>
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<td>0.74 ± 0.22</td>
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<td>0.06 ± 0.03</td>
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<td><em>Stenochironomus</em></td>
<td>1.31 ± 1.31</td>
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<td>0.00 ± 0.00</td>
<td>0.14 ± 0.14</td>
<td>0.92 ± 0.66</td>
<td>0.04 ± 0.04</td>
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<td><strong>Tanytarsini</strong></td>
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<tr>
<td><em>Cladotanytarsus</em></td>
<td>0.69 ± 0.29</td>
<td>0.16 ± 0.11</td>
<td>6.39 ± 2.92</td>
<td>0.69 ± 0.26</td>
<td>0.44 ± 0.17</td>
<td>0.46 ± 0.28</td>
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<td><em>Constempellina</em></td>
<td>1.21 ± 0.92</td>
<td>11.04 ± 6.27</td>
<td>6.42 ± 1.74</td>
<td>0.72 ± 0.54</td>
<td>0.72 ± 0.23</td>
<td>0.11 ± 0.04</td>
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<tr>
<td><em>Microspectra</em></td>
<td>6.69 ± 1.56</td>
<td>29.91 ± 13.26</td>
<td>26.27 ± 11.18</td>
<td>1.3 ± 10.79</td>
<td>0.59 ± 0.13</td>
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<td><em>Rheotanytarsus</em></td>
<td>0.89 ± 0.41</td>
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<td>0.02 ± 0.02</td>
<td>0.23 ± 0.22</td>
<td>0.32 ± 0.30</td>
<td>0.01 ± 0.01</td>
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<td><em>Stempellina</em></td>
<td>0.16 ± 0.15</td>
<td>2.75 ± 2.49</td>
<td>1.07 ± 0.75</td>
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<tr>
<td><em>Tanytarsus</em></td>
<td>1.38 ± 1.07</td>
<td>0.11 ± 0.09</td>
<td>0.05 ± 0.04</td>
<td>0.15 ± 0.13</td>
<td>0.53 ± 0.20</td>
<td>0.02 ± 0.01</td>
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<td><strong>Orthocladiinae</strong></td>
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<td><em>Corynoneura</em></td>
<td>0.07 ± 0.05</td>
<td>0.43 ± 0.33</td>
<td>0.63 ± 0.25</td>
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<td><em>Brilia</em></td>
<td>1.67 ± 0.44</td>
<td>1.07 ± 0.63</td>
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<td><em>Hydrobaenus</em></td>
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<td>0.24 ± 0.17</td>
<td>0.28 ± 0.20</td>
<td>0.02 ± 0.01</td>
<td>0.20 ± 0.09</td>
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<td><em>Parametricrion</em></td>
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<td>2.07 ± 1.79</td>
<td>0.70 ± 0.32</td>
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<td><em>Rheosmitia</em></td>
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<td><strong>Tanytarsine</strong></td>
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<td><em>Krenopelopia</em></td>
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<td>8.79 ± 4.88</td>
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<td><em>Monopelopia</em></td>
<td>0.25 ± 0.17</td>
<td>0.42 ± 0.37</td>
<td>0.28 ± 0.28</td>
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<td><em>Nataria</em></td>
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<td>0.13 ± 0.08</td>
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Pre = pretreatment, LE = litter exclusion, LE + WR = litter and wood exclusion. Different letters indicate significant differences from BACI analysis (i.e., Before-After differences between the reference and treatment streams).

amount of organic matter and the proportion of inorganic substrates over time.

The taxonomic assemblage differed between the streams. At the genus level, Tanytarsini were positively associated with higher organic matter standing stocks in the reference stream. Tanytarsini genera associated with higher amounts of organic matter included *Microspectra* sp. and *Constempellina* sp. Some Orthocladiinae were also positively associated with axis 1, such as *Parachaetocladius* sp., *Brilia* sp., and *Parametricrionemus* sp. (Table 4). The Orthocladiinae, *Lopescladius* sp., was associated with low organic matter standing stocks and higher proportions of inorganic substrates in the litter exclusion stream. The Chironomini, *Stenochironomus* sp. and Tanytarsini, *Rheotanytarsus* sp., were associated with Axis 2. A decline in chironomid biomass corresponded with the decline in organic matter. For example, chironomid biomass in the LE stream was much closer to the chironomid biomass in the reference stream in autumn and summer prior to litter exclusion. Organic matter standing stocks and chironomid biomass and assemblage structure did not change appreciably in the reference stream by season or year, evidenced by the tight clustering of sampling points among years. Much more seasonal variability occurred in the litter exclusion stream (Fig. 4).
Table 3 Pearson and Kendall correlations of independent variables (CPOM, FPOM, and % Sand, Gravel, and Pebble) with CCA axes

<table>
<thead>
<tr>
<th>Independent variables</th>
<th>Pearson correlation</th>
<th>r²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biomass</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Axis 1</td>
<td>0.60</td>
<td>0.36</td>
</tr>
<tr>
<td></td>
<td>0.59</td>
<td>0.39</td>
</tr>
<tr>
<td></td>
<td>-0.62</td>
<td>0.39</td>
</tr>
<tr>
<td>Axis 2</td>
<td>-0.42</td>
<td>0.17</td>
</tr>
<tr>
<td></td>
<td>-0.39</td>
<td>0.15</td>
</tr>
</tbody>
</table>

Discussion

Spatial and temporal differences in abundance and biomass

Chironomids are often bivoltine, fast colonizers found in high densities in disturbed habitats. These attributes allow them to be among the few taxa able to exploit patchy, ephemeral food resources (Palmer et al., 2000). In this study, chironomid abundance and biomass declined at the family level, resulting from seven years of litter exclusion, but it did not consistently decline at the subfamily and generic levels. However, our results did show a clear decline of Tanytarsini at the tribe-level following both leaf litter and leaf litter and wood exclusion. The most abundant genera within the tribe Tanytarsini: Cladotanytarsus sp., Constempellina sp., and Micropsectra sp., were expected to decline significantly but only one genus did, Constempellina sp. Conversely, Orthocladiinae abundance and biomass did not change and seasonal variation was high. This spatial and temporal variation can be attributed to seasonal fluctuations in available food resources coupled with the phenology of different taxa.

Other studies have shown that high quality organic matter patches attract invertebrates when

Fig. 4 Canonical correspondence analysis ordination of mean biomass for all taxa. Taxa representing < 5% in total samples were omitted in each stream from 1992 through 2000. Taxon names are abbreviated with BRIL = Brilia, CONS = Constempellina sp., HELE = Heleniella sp., LOPE = Lopescladius sp., MICR = Microsectra sp., PARA = Parachaetocladius sp., PARM = Parametriocnemus sp., RHEO = Rheotanytarsus sp., and STEN = Stenochironomus sp. Independent variables shown are FBOM = fine benthic organic matter, CBOM = coarse benthic organic matter, and % sand, gravel, and pebble
carbon is limited (e.g., Palmer et al., 2000; Eggert & Wallace, 2003b). Because some particulate organic matter was present within the litter exclusion stream, patches of fine particles created 'resource islands' for chironomids. Fine particulate organic matter did decline in the LE stream; however, concentrations may have been high enough, especially in spring, to support a small and spatially variable population of chironomids (Hurny, 1990; Berg, 1995). Tanytarsini, specifically, have been shown to be associated with organic matter (Oliver, 1971; Pinder, 1986) and were likely able to track organic matter even at low densities. For example, Eggert and Wallace (2003b) previously found that a shredding caddisfly, Pycnopsyche gentilis, was able to closely track leaf litter during litter exclusion until standing stocks declined below 25–50 g AFDM m⁻². Consequently, Pycnopsyche gentilis production declined significantly. Chironomids may be able to track even lower amounts of organic matter and this could have maintained a link in the detrital food web when other taxa were unable to persist (Wallace et al., 1999).

Life history differences among taxa helped to promote the success of Orthocladiinae in the LE stream. Orthocladiinae are often most abundant in winter and spring when temperatures are cool and algae and DOC are most abundant (Boerger, 1981; Berg & Hellenthal, 1992). Consequently, Orthocladiinae were able to maintain higher densities than other groups following litter exclusion. Conversely, Tanytarsini are often abundant in summer and have been shown to feed on slow-breakdown leaf litter and fine particulates. The absence of summer leaf litter may exclude some Tanytarsini genera, while others would feed on FPOM (Grubbs & Cummins, 1994). Grubbs & Cummins (1994) found chironomids to be important especially important in processing organic matter in streams with low summer leaf standing stocks when few other shredders were present. These studies emphasize the importance of chironomids in processing organic matter in streams when standing stocks may be too low to support other larger organisms (i.e., shredding Plecoptera).

Taxonomic composition

Chironomid taxonomic richness and evenness did not change. Recruitment still occurred and our results are supported by those of Lugmuth et al. (1990) in a nearby Coweeta stream that was treated with insecticide. In their study, chironomid secondary production declined but continuous recruitment from aerial adults maintained benthic diversity. Baer et al. (2001) experimentally confirmed that the litter exclusion stream was continually colonized by aerial adults from downstream, but colonization was lower than before litter exclusion. Our results suggest that emergence and recruitment might have declined due to limited resources for hatched larvae or oviposition of less fecund adults from within the litter exclusion reach. Furthermore, individuals hatched in the litter exclusion stream experienced increased predation pressures (Hall et al., 2000; Wallace et al., 1999) and an elevated per capita drift of late-instar Siler et al., 2001) in response to limited food and habitat in the litter exclusion stream.

Chironomidae response to habitat alteration

Tanytarsini responded most dramatically to litter exclusion. Several unique characteristics made them particularly vulnerable to litter and wood exclusion. First, Tanytarsini are small, rheophilic chironomids that depend on stable habitat for either case (e.g., Constempellina sp.) or tube (e.g., Micropsectra sp.) attachment (Oliver, 1971; Pinder, 1986; Coffman & Ferrington, 1996). The loss
of leaves and wood likely eliminated Tanytarsini habitat (Chaloner & Wotton, 1996) and created an environment less buffered from disturbances (Bilby & Likens, 1980; Smock et al., 1989). Food was also reduced and became more limited. Fine particulate organic matter (FPOM), a known food source for many Tanytarsini, declined from about 1500 g to 200 g AFDM m\(^{-2}\) from the start of LE through the end of the seven-year study period (as cited in Wallace et al., 1999; Eggert & Wallace, 2003a). The temporal and spatial heterogeneity of FPOM would have contributed to the decline and variability found in Tanytarsini abundance and biomass.

The additional removal and exclusion of small and large wood had the largest impact on the chironomids due to the combined loss of habitat, retention structures, and the direct reduction of food resources. The physical attributes of wood in streams are well known, but less is known about the role it plays as a food resource (Bilby & Bisson, 2000). Wood provides critical habitat for biofilms, a complex matrix of algae, bacteria, and fungi (Golladay & Sinsabaugh, 1991; Couch & Meyer, 1992) and retains fine particles. Bacteria and algae on the wood provided a labile food source for chironomids in the LE stream (Hall & Meyer, 1998; Hart & Lovvorn, 2003). Some of the Orthocladiinae in the LE stream had scraping mouthparts able to feed on biofilms growing on wood and inorganic substrate (i.e., Lopescladius sp., Parachaeotcladius sp., and Rheosmittia sp.). Furthermore, Meyer et al. (1998) found that overall, dissolved organic carbon (DOC) concentrations declined as leaf standing stock declined; however, DOC concentrations did not significantly decline in the spring and summer because leaves were not the primary DOC source. This DOC would have been an important nutrient source for bacteria and algae on FPOM, sand, and rocks in the litter exclusion stream during summer.

Many Orthocladiinae persisted in the litter exclusion stream after litter and wood exclusion, the genus Lopescladius sp. being most successful. Lopescladius sp. abundance and biomass increased during spring, autumn, and winter in the litter exclusion stream; however, their absence during summer confounded statistical analyses.

Huryn (1990) found that the most abundant chironomid taxa in Coweeta streams were present throughout the year, but Lopescladius sp. was not abundant in his study. However, in this study, chironomid abundance and biomass declined, particularly Lopescladius sp., in summer when canopy was closed and fine organic matter export was low. The Orthocladiinae that persisted in the litter exclusion stream were small, bivoltine species, making them very fast colonizers. In fact, when leaves were added back in autumn to the stream, they were among the first to colonize (J.B. Wallace & S.L. Eggert, University of Georgia, unpublished data). They are also good swimmers, adapted to fast-flowing, cold water streams that often experience frequent hydrologic disturbance (Coffman & Ferrington, 1996; Lods-Crozet et al., 2001). Some researchers have found Lopescladius sp. and Rheosmittia sp. to be indicative of disturbance in southeastern U.S. streams and to dominate sandy, coastal plain blackwater rivers (see also Soluk, 1985; Benke, 1998, Rinella & Feminella, 2005).

Community response to resource reduction

A combination of parametric statistics and ordination proved to be an insightful approach for exploring generic-level community dynamics within the litter exclusion stream and between the reference and litter exclusion streams. Ordination analyses clearly illustrated differences in the chironomid community structure between the reference and litter exclusion streams. It also showed that the biomass and community assemblage in the litter exclusion stream was more closely related to the reference stream before litter exclusion, especially in autumn. Differences in chironomid biomass between the two streams increased with each passing year of litter exclusion. These differences were related to the amount of fine and coarse particulate organic matter in the streams and the amount of small inorganic substrates. Differences in inorganic substrate were more important in structuring the community before LE and seemed to become less important with the continued exclusion of leaf litter and wood.
Functional feeding

Macroinvertebrates are often classified into functional feeding groups (FFGs) to clarify their ecological significance (Cummins, 1974), but chironomids are typically grouped as predators (Tanypodinae) or collector-gatherers (non-Tanypodinae) in ecosystem-level manipulations (e.g., Wallace et al., 1999). The use of FFGs to categorize chironomid feeding habits is problematic because chironomids are often facultative feeders that shift their diet with the most abundant food available. The particles they ingest are so small it can only be identified as amorphous detritus (Berg, 1995; Tavares-Comar & Williams, 1996; Rosi-Marshall & Wallace, 2002; Henriques-Oliveira et al., 2003). Our results show that categorizing chironomid feeding into a functional mode can lead to an oversimplification of their response to environmental change.

Chironomidae facultative feeding habits make categorization into functional feeding groups difficult. Out of 22 dominant chironomid genera in this study, 12 were classified as collector-gatherers, six had not been classified, and two were predators. However, categorizing chironomids by subfamily was informative and the additional classification of feeding modes and other life history attributes of subfamilies provided more insight into the overall chironomid response to litter exclusion. A few key genera were sensitive to changes in organic matter standing stocks (i.e., Constempellina sp. and Lopescladius sp.) and loss of wood (e.g., Micropsectra sp.). These taxa may be useful as indicator species in altered stream ecosystems.

Conclusions

The chironomid assemblage changed as a result of excluding terrestrial detrital inputs. Lowered standing stocks of CPOM, FPOM, and DOC concentrations negatively affected the Tanytarsini, but not Tanypodinae or Orthocladiinae, in fact, Orthocladiinae abundance and biomass did not change over the course of litter exclusion. The coupled removal of litter and wood had the largest affect on the chironomid community, due to a loss of habitat and stable retention structures, as well as the indirect reduction of food resources (e.g., biofilms, Golladay & Sinsabaugh, 1991; Couch & Meyer, 1992). Orthocladiinae were able to persist because of their habitat preferences, feeding strategies, and recolonization by aerial adults. From a management perspective, our results suggest a forested headwater stream community dominated by Orthocladiinae such as Lopescladius sp. and Rheosmittia sp. and a lack of common Tanytarsini could be indicative of an ecologically significant loss of detritus. This may occur in streams with disturbed riparian habitats or streams with increased inorganic sediments and loss of retention structures.

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