Contribution of microbial and invertebrate communities to leaf litter colonization in a Mediterranean stream

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Abstract. Leaf litter inputs and retention play an important role in ecosystem functioning in forested streams. We examined colonization of leaves by microbes (bacteria, fungi, and protozoa) and fauna in Fuirosos, an intermittent forested Mediterranean stream. Black poplar (Populus nigra) and plane (Platanus acerifolia) leaf packs were placed in the stream for 4 mo. We measured the biomasses and calculated the densities of bacteria, fungi, protozoa, meiofauna, and macroinvertebrates to determine their dynamics and potential interactions throughout the colonization process. Colonization was strongly correlated with hydrological variability (defined mainly by water temperature and discharge). The 1st week of colonization was characterized by hydrological stability and warm water temperatures, and allocation of C from microbial to invertebrate compartments on the leaf packs was rapid. Clumps of fine particulate organic matter (FPOM) were retained by the leaf packs, and enhanced rapid colonization by microfauna and meiofaunal collector-gatherers (ostracods and copepods). After 2 wk, an autumnal flood caused a 20-fold increase in water flow. Higher discharge and lower water temperature caused FPOM-related fauna to drift away from the packs and modified the subsequent colonization sequence. Fungi showed the highest biomass, with similar values to those recorded at the beginning of the experiment. After 70 d of postflood colonization, fungi decreased to nearly 40% of the total C in the leaf packs, whereas invertebrates became more abundant and accounted for 60% of the C. Natural flood occurrence in Mediterranean streams could be a key factor in the colonization and processing of organic matter.

Key words: Mediterranean stream, meiofauna, protozoa, bacteria, fungi, macroinvertebrates, C budget.

Leaf litter is an important energy source for food webs in forested stream ecosystems (Fisher and Likens 1973, Webster and Benfield 1986), especially in low-order systems (Minshall et al. 1985). Input and retention of organic matter in Mediterranean forested streams are strongly affected by climate (Sabater et al. 2001, Acuña et al. 2005) because the annual peak of leaf fall often coincides with heavy flooding in autumn (Acuña et al. 2007, Gasith and Resh 1999). Thus, colonization and decomposition of leaf litter might be determined by the availability of organic matter and hydrology in Mediterranean forested streams (Gasith and Resh 1999). Floods are a major organizing factor of organic matter dynamics because they determine cycles of accumulation and removal. Consequently, they provoke changes in consumer communities.
Leaf structure and chemistry and streamwater chemistry are the main determinants of microbial colonization and metabolism (Bärlocher et al. 1995, Gulis and Suberkropp 2003). Fungi are the main colonizers during initial phases, whereas bacteria dominate during late stages (Baldy et al. 1995) and probably benefit from fungus-induced changes in leaf surfaces or from the release of labile compounds (Allan 1995).

The dominant role of fungi during colonization of large particulate organic matter (Gessner 1997, Findlay et al. 2002) and their contribution to overall loss of C from leaves (Hieber and Gessner 2002, Gulis and Suberkropp 2003, Pascoal and Cassio 2004) mean that net C transformation mostly reflects the capacity of the fungal community to metabolize leaf matter.

The shredding activity of macroinvertebrates on decaying leaves contributes to the breakdown process. Shredders feed on coarse particulate organic matter and can accelerate decomposition by increasing leaf tissue fragmentation (Graça 2001, Graça and Canhoto 2006). Macroinvertebrates feed preferentially on conditioned leaves, thereby ingesting nutrients from leaf tissue and from the attached microbial community (Slansky and Scriber 1985). However, little information is available on the contribution of protozoa and smaller invertebrates to colonization and decomposition (Franco et al. 1998, Robertson and Milner 2001). Micro- and meiofauna are mostly bacterial specialists (Meyer 1994, Borchardt and Bott 1995), and they can exert significant grazing pressure on bacteria and fungi that colonize detritus (Perlmutter and Meyer 1991). Given the interaction between distinct groups of consumers, leaf litter breakdown is not a simple linear cause-and-effect relationship, but a simultaneous complex interaction between groups that could lead to several outcomes (Gessner et al. 1999).

We examined the complex interaction between decomposers and detritivores with regard to colonization of organic matter. We analyzed the successional colonization dynamics of microbes (including bacteria, fungi, and protozoa) and meio- and macrofauna on decaying leaves during a leaf-pack experiment in a forested Mediterranean stream. We addressed the following: 1) the most important factors that affect colonization dynamics, 2) the settlement sequence of leaf litter colonizers, and 3) the contribution of microbial organisms and meio- and macrofauna to the leaf litter biofilm and to the total C budget of the leaf packs. We hypothesized that the sequence of colonization would occur in the order: microbes, meiofauna, and macrofauna. If this hypothesis is correct, C allocation should change between compartments over time. We also speculated that co-occurrence of organic matter processing and high climatic instability could introduce disruptions or accelerations into this sequence in Mediterranean streams.

**Methods**

**Study site**

We worked in an intermittent 3rd-order stream (Fuirosos) in northeastern Spain (lat 41°42′N, long 2°34′W). Fuirosos drains a 10.9-km² forested catchment that lies in the Natural Park of the Montnegre–Corredor range. Precipitation occurs mostly in autumn and spring, which are periods of frequent flooding, with occasional storms in summer. High variation in rainfall is characteristic of this typical Mediterranean system, which has large deviations in mean monthly precipitation and considerable interannual differences. The stream usually has a mean discharge of 30 L/s, width of 3 to 4 m, and depths ranging from 0.1 to 0.5 m (see details in Acuña et al. 2005).

We did the colonization study in a 50-m-long reach. The nearby riparian area (10–15-m width) formed a closed canopy from May to October. Dominant vegetation was alder (Alnus glutinosa L.), black poplar (Populus nigra L.), hazelnut (Corylus avellana L.), and plane (Platanus acerifolia Aiton–Willd.). Direct and lateral inputs of organic material (OM) ranged from 0.1 to 4.45 g C m⁻² d⁻¹ (Acuña et al. 2007). The canopy reduced incident light in summer, and total light increased in winter (Román et al. 2004).

**Sampling strategy**

We used the litter bag technique to monitor the microbial and faunal colonization of leaf litter. We collected recently abscised leaves of P. acerifolia and Populus nigra from the riparian floor in September 2003. We dried the leaves (room temperature, 48 h) and sterilized them in an autoclave (121°C, 30 min) before placing them in plastic mesh bags (1-mm mesh). The sterilization procedure can affect processing rates (Godshalk and Wetzel 1978), but we used it to eliminate previous colonizers. We used the relatively small mesh to facilitate analysis of the C pathway throughout the biofilm where the microbial loop occurs and to prevent ingestion of the leaf tissue by larger external macroinvertebrates. We placed 11 thin ropes supporting 3 bags each (a total of 33 bags, each containing 4 Platanus and 6 Populus leaves) every 3 to 5 m (depending on the presence of natural leaf accumulation zones) along the 50-m reach. We retrieved 3 randomly selected bags on days 1, 2, 4, 7, 17, 28, 44, 58, 73, 93 and 112. Day 0 was 6 October 2003 when litter bags were immersed.

We reduced the loss of nonattached micro- and
meiofauna from the leaf packs during retrieval by placing the still-immersed bags in a plastic container, which was then removed from the stream. We removed macroscopic colonizers from the outside of the bags if they were present. We took a leaf subsample and its surrounding water from the container and counted attached and nonattached protozoa. We used a metal borer to cut 1.1-cm-diameter disks from the leaves and biofilm to sample fungi and bacteria. We placed samples for measurement of ergosterol (to estimate fungal biomass) in plastic vials and froze them (−20°C) until analysis. We preserved samples for bacterial density in formalin (2%). We dried additional leaf disks (70% preserved samples for bacterial density in formalin used empirical bacterial biovolumes of 0.147 μm³ to estimate bacterial density in formalin). We passed the rest of the contents of the mesh bag through 500-μm and 32-μm nested sieves. We scraped the leaf fraction retained by the 500-μm sieve to detach macroinvertebrates and stored the leaves for later measurement of surface area with a scanner and ImageJ software (v. 1.40; http://rsbweb.nih.gov/ij/index.html), biomass, and C content. We fixed macroinvertebrates retained in the 500-μm sieve immediately in 4% formalin, and stored them for further counting and identification. We retained the fraction between the 500-μm and 32-μm sieves and stored it at 4°C for counting live meiofauna.

**Physicochemical measurements**

We measured O₂, pH, conductivity, and temperature with handheld meters (MultiLine F/SET-3; WTW, Weilheim, Germany) in the field on each sampling date. We measured current velocity (MiniAir2; Schiltknecht, Zurich, Switzerland) beside the litter bags before they were collected. We calculated stream discharge by using the slug-injection method with NaCl as the tracer (Gordon et al. 1992).

**Bacterial density and biomass**

We estimated bacterial density in triplicate in each litter bag after sonicating the samples (2 + 2 min, 40-W power, 40-kHz frequency; Ultrason, Selecta, Abrera, Spain). After appropriate dilution, we stained fixed samples for 5 min with 4′,6-diamidino-2-phenylindole (final concentration 2 μg/μL) and passed them through 0.2-μm irgalan black-stained polycarbonate filters (Nuclepore; Whatman, Maidstone, UK). We counted bacteria in 15 fields/filter (400–800 organisms) with a fluorescence microscope (Eclipse E-600; Nikon, Tokyo, Japan) at 1250× magnification. We measured the volume of cells with a Soft Imaging System (analySIS®; Olympus, Münster, Germany). We used empirical bacterial biovolumes of 0.147 μm³ (Populus) and 0.163 μm³ (Platanus) and a conversion factor of 2.2 × 10⁻¹³ g C/μm³ to estimate bacterial C (Bratbak and Dundas 1984).

**Ergosterol content**

We lyophilized frozen leaf samples and used 3 subsamples from each litter bag for ergosterol extraction by saponification with methanol (80°C, 60 min) in a shaking bath. We purified the resulting extracts by solid-phase extraction (Gessner and Schmitt 1996) and measured ergosterol by high-performance liquid chromatography (Waters, Milford, Massachusetts). We detected ergosterol at 282 nm and quantified it by comparison with ergosterol standards (0–200 μg/mL; Fluka Chemical Co., Milwaulkee, Wisconsin) (Gessner and Schmitt 1996). We estimated fungal C biomass on the basis of an ergosterol content of 5.5 mg/g fungal biomass (Gessner and Chauvet 1993) and 43% C content in fungal dry mass (Baldy and Gessner 1997).

**Determination of microfauna, meiofauna, and macroinvertebrates**

We counted live microfauna (protozoan) samples on the day of collection with a microscope at 400× magnification (Polivar; Reichert-Jung, Wien, Austria). We identified and measured soft-bodied meiofauna in formalin for further length and width measurements. We estimated biomass from exponential equations when they were available, or otherwise from biovolumes. We found published power equations or values for Ephemeroptera, Ple-
coptera, Coleoptera, Diptera (Meyer 1989, Benke et al. 1999), Rotifera (Bott and Borchardt 1999), Copepoda, and Cladocera (Bottrell et al. 1976). We used biovolum estimates for Nematoda (after Andrassy 1956), Oligochaeta (Smit et al. 1993), Ostracoda, Hydracarina, Tardigrada, and Microturbellaria (Ramsay et al. 1997). We sorted fixed macroinvertebrates with a dissecting microscope, dried them (70°C) to a constant mass, and burned them (450°C, 4 h) to obtain biomass as ash-free dry mass. We expressed all densities (meiofauna and macroinvertebrates) per microgram of leaf DM. However, we obtained biomass (micrograms of organism C) through standard conversions (Waters 1977) and expressed it per microgram of leaf C in each litter bag.

Data analyses

We used nonparametric Spearman’s rank coefficient to identify possible correlations between environmental and biological variables. We applied a false discovery rate correction (FDR; Benjamini and Hochberg 1995) because of the large number of comparisons made. We did these statistical analyses with STATISTICA (version 8.0; StatSoft, Tulsa, Oklahoma). We used ANOSIM on 4th-root(x)-transformed data to compare average ranked Bray–Curtis similarities among sampling dates with average ranked similarities within a sampling date (between replicates). We quantified dissimilarity between community assemblages through the colonization period with similarity percentages (SIMPER). We did these analyses with PRIMER (version 6.1.6; PRIMER-E, Plymouth, UK).

Results

Physicochemical

Water temperature decreased throughout the study period. It ranged from ~15°C in October to ~4°C in January (Table 1). On day 11, water flow suddenly increased from 1.2 to 90.4 L/s as a result of rain, and dissolved O₂ content increased and conductivity decreased. Water velocity next to litter bags ranged from 0.01 to 0.64 m/s. The flood caused significant differences among bags retrieved on consecutive sampling dates (Kruskal–Wallis statistic = 25.86, p < 0.005), but not among bags retrieved on the same day.

Bacteria and fungi

Fungi and bacteria accumulated on the leaf material following a logistic pattern (Fig. 1A). However, bacteria began a 2nd increase after day 73 of the experiment, whereas fungi remained stable until the end of the experiment. Fungal biomass was ~200× higher than bacterial biomass except during week 2 (Fig. 1A).

Microfauna

The most abundant groups of microfauna were flagellates, testamoebae, and ciliates (Fig. 2A). Microfauna density and biomass increased during week 1 (Fig. 1A) to 1.12 × 10⁵ individuals (ind.)/g leaf DM (testamoebae) and 7.3 × 10⁴ ind./g leaf DM (flagellates) (Fig. 2A). Ciliate density was moderate (240–~5000 ind./g leaf DM), except on day 93 when it peaked at 1.95 × 10⁴ ind./g leaf DM. Microfauna community composition differed among days (ANOSIM, R = 0.52, p = 0.001), and these differences were greatest (with dissimilarities >70%) immediately after the flood (day 17; SIMPER), when microfauna density was almost 0 (Fig 2A). Microfauna densities remained low until day 93, when they increased again. On day 93, ciliate density was high, but taxon richness was lower than before day 17. Hypotrichia (UFF) and Hymenostomata (DFF) were the most abundant ciliates from day 93 to day 112, when Pleurostomatida (RF) also were present.

Meiofauna

Densities of Chironomidae, Rotifera, and Oligochaeta were higher than those of other groups of meiofauna. Community composition differed significantly among sampling days (R = 0.835, p = 0.001) and between samples collected before and after the flood (R = 0.73, p = 0.001). Microcrustaceans were the most common group of meiofauna only during the 1st week. This group consisted mainly of ostracods and copepods (with more nauplii than copepodites and adults) and, to a lesser extent, cladocerans. Two patterns were observed during colonization (Fig. 2B), corresponding to temporary and permanent meiofauna. Temporary meiofauna, including early larval stages of Chironomidae, Oligochaeta, Plecoptera, and Ephemeroptera (Robertson et al. 2000), followed the dynamics of their macroinvertebrate representatives (Table 2, Fig. 1B) and increased during the late phases of colonization. Densities of permanent meiofauna (Rotifera, Nematoda, Microcrustacea, Microturbellaria, and Tardigrada) were lower than those of temporary meiofauna (Fig. 2B). Permanent meiofauna contributed to total meiofauna density, but most meiofaunal biomass was made up by temporary meiofauna (Fig. 3).

Macroinvertebrates

Macroinvertebrate densities were low and ranged between 0 and 800 ind./g leaf DM (Fig. 2C). Large
Ostracoda (>250 μm) were the earliest colonizers, followed by Plecoptera (mainly Nemouridae) and Chironomidae (abundant on day 44; Fig. 2C). Ephemeroptera, Oligochaeta, and Coleoptera increased late in the study. Macroinvertebrate biomass increased during colonization, and made up most of the total invertebrate biomass at the end of the study (Fig. 1B).

**TABLE 1. Physical and chemical characteristics of Fuirosos stream water during the 4-mo colonization experiment. Values are individual measures of variables on the 11 sampling days.**

| Day | Temperature (°C) | O₂ (mg/L) | Conductivity (µS/cm) | pH | Discharge (L/s) |
|-----|------------------|-----------|----------------------|----|----------------|
| 1   | 11.6             | 6.41      | 307.0                | 6.67 | 4.62          |
| 2   | 12.5             | 6.29      | 305.0                | 6.75 | 4.62          |
| 4   | 13.7             | 5.83      | 314.0                | 6.6  | 1.46          |
| 7   | 15.3             | 5.2       | 302.0                | 6.54 | 1.19          |
| 17  | 12.8             | 10        | 181.8                | 7.00 | 90.41         |
| 28  | 11.8             | 18.1      | 188.0                | 6.64 | 63.97         |
| 44  | 9.4              | 9.66      | 196.0                | 7.44 | 14.66         |
| 58  | 10.3             | 10.61     | 173.0                | 7.24 | 60.05         |
| 73  | 8.5              | 11.20     | 174.3                | 7.79 | 60.10         |
| 93  | 4.3              | 12.64     | 188.6                | 7.73 | 30.23         |
| 112 | 8.2              | 9.55      | 198.2                | 6.82 | 14.00         |
| Mean ±1 SE | 10.8 ± 0.9 | 9.59 ± 1.13 | 229.81 ± 18.61 | 7.02 ± 0.46 | 31.39 ± 31.64 |

**FIG. 1.** Mean (±1 SE) bacterial, fungal, and microfaunal biomass (A) and meiofaunal and macroinvertebrate biomass (B) during a 4-mo study of colonization of leaf litter in Fuirosos. Error bars are as follows: bacteria (positive thick line and long cap), fungi (negative thick line and short cap), microfauna (thin line and short cap), meiofauna (thin line and long cap), and macroinvertebrates (thick line and short cap).

**FIG. 2.** Mean density of microfauna (A), meiofauna (±1 SE) (B), and macroinvertebrates (±1 SE) (C) during a 4-mo study of colonization of leaf litter in Fuirosos. In panel A, densities of ciliates and Testamoebae are shown on y-axis 1 and densities of flagellates are shown on y-axis 2. In panels B and C error bars are as follows: temporary and permanent meiofauna, Oligochaeta, and Ephemeroptera (normal line and cap); Plecoptera (positive cap); and Chironomidae (negative cap). DM = dry mass.
Relationships between organisms and environmental characteristics

Microcrustacean biomass and conductivity were significantly negatively correlated with discharge (Table 3). Some ciliates also tended to be negatively correlated with discharge, but these relationships were not statistically significant after FDR correction. Oligochaeta, Chironomidae, Plecoptera, and Ephemeroptera biomasses were significantly positively correlated with bacterial biomass and negatively correlated with water temperature. Fungal biomass was significantly positively correlated with macroinvertebrate biomass and negatively correlated with water temperature (Table 3).

Allocation of C during the colonization

Fungal biomass accounted for ~32 to ~93% of the total organismal (nonleaf) C in the litter bags during the experiment, bacterial biomass accounted for ~0.2 to ~6%, and faunal biomass accounted for ~0.03 to ~8.4% (meiofauna) and ~1.7 to ~58% (macroinvertebrates) (Table 4). Between days 1 and 7, C exchange between compartments was rapid and shifted from fungal (~93%) to bacterial (~6%) and faunal (mainly macroinvertebrate) compartments (~30%). The flood reset the system and shifted the highest percentage of C back to the fungal compartment. Fungi continued to have the highest percentage of total C until late in the experiment when macroinvertebrates and, to a lesser extent, temporary meiofauna, made up most of the nonleaf C in the litter bags. Between days 28 and 58, C in the temporary meiofaunal compartment was transferred to the macroinvertebrate compartment. After the flood, when water temperature was lower and discharge was higher than before the flood, almost 70% of the total nonleaf C in the litter bags consisted of invertebrate biomass and 30 to 40% corresponded to fungal biomass.

Discussion

Factors affecting colonization dynamics

Flow and temperature influence leaf litter breakdown. In desert streams, the duration of flood-free periods is an important factor regulating community biomass and efficiency (Grimm and Fisher 1989). In Mediterranean streams, low rainfall and high temperatures in summer cause early leaf fall in riparian forests (Acuña et al. 2004). Vázquez et al. (2007) reported an increase of dissolved organic C when flow resumed because of the partly decomposed leaves that had accumulated in the stream bed. Therefore, this accumulated organic C is available as a source of energy and shelter to aquatic organisms. Mechanical breakdown caused by the turbulent waters when flows resume provides more surface area for colonization, accelerates decomposition of leaf litter, and enhances its conditioning (Heard et al. 1999, Graça 2001). A similar process occurs in semiarid Australian rivers (Francis and Sheldon 2002).

In Fuirosos, biological succession on decaying leaf litter was strongly correlated with discharge and water temperature. Our study began shortly before a flood associated with the end of the summer drought. Fine particulate OM (FPOM) accumulation and rapid primary colonization occurred during the initial period of low discharge and moderate water temperature. During the flood, the FPOM drifted downstream. After the flood, Populus leaves broke apart and were almost entirely washed out of the mesh bags by day 58; Platanus leaves, though broken, remained in the bags. The flood reset the colonization process. Subsequent recolonization occurred under higher flow, cooler temperatures, and different leaf characteristics, and
these changes were associated with shifts in the composition of the colonizing community and slower transfer of C between compartments.

Settlement sequence of colonizers

Our choice of mesh size for the litter bags allowed us to study colonization of leaf litter by micro- and meiofauna, but also might have favored accumulation of detritus and its associated fauna. A procedural control might have permitted us to correct for litter-bag effects (Boulton and Boon 1991). However, Acuña et al. (2005) surveyed naturally deposited leaves in Fuirosos and reported similar invertebrate composition and densities of the same order of magnitude as those in our litter bags on days 2 and 4.

Hydrological stability (discharge between 1–5 L/s, water velocities <12 cm/s) and high water temperatures allowed initial colonization of litter bags by microfauna, which achieved maximum diversity in this period. UFF and DFF feeding types appeared during this early stage of colonization. Both groups have membranelles that allow them to capture small particles (DFF: ≤2 μm, UFF: ≤2 μm), such as bacteria and flagellates, that were abundant in this period. However, microfaunal densities were much lower than those reported on decomposing leaves in temperate streams (Bott and Kaplan 1989, Schönborn 1982) or in sandy sediments in Fuirosos (Domènech et al. 2006).

During week 1, FPOM retained by the litter bags was colonized quickly by meiofauna. Microcrustaceans thrived in these conditions because of their feeding preference for FPOM-associated microflora (Perlmutter and Meyer 1991) and the shelter provided by the clumps of FPOM (Robertson and Milner 2001, Gaudes et al. 2006). Clumps of FPOM also facilitate

| Variable                  | Fungi | Bacteria | Temperature | Discharge |
|---------------------------|-------|----------|-------------|-----------|
| Chironomidae (TM)         | 0.56  | 0.83**   | −0.73**     | 0.47      |
| Chironomidae              | 0.62* | 0.81**   | −0.75**     | −0.03     |
| Ephemeroptera (TM)        | 0.66* | 0.86**   | −0.79**     | 0.34      |
| Ephemeroptera             | 0.76**| 0.88**   | −0.90**     | 0.24      |
| Gastropoda                | 0.74**| 0.83**   | −0.87**     | 0.13      |
| Microcrustacea (PM)       | −0.62*| 0.58     | 0.49        | −0.87**   |
| Oligochaeta (TM)          | 0.78**| 0.79**   | −0.75**     | 0.26      |
| Oligochaeta               | 0.69* | 0.88**   | −0.81**     | 0.22      |
| Plecoptera (TM)           | 0.90**| 0.77**   | −0.72**     | 0.53      |
| Plecoptera                | 0.75**| 0.92**   | −0.88**     | 0.49      |
| Rotifera (PM)             | 0.66* | 0.83**   | −0.73**     | 0.02      |
| Fungi                     | −0.67*| 0.67**   | −0.83**     | 0.48      |
| UFF ciliates              | −0.38 | 0.04     | 0.09        | −0.62*    |
| DO                        | 0.62* | 0.62*    | −0.60       | 0.88**    |
| Temperature               | −0.83**| −0.76** | −            | −0.36     |
| Conductivity              | −0.60 | −0.62*   | 0.39        | −0.85**   |

These changes were associated with shifts in the composition of the colonizing community and slower transfer of C between compartments.

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**Table 4.** Percentages of total organismal C in each group of organisms that colonized leaf litter bags during a 4-mo experiment. Total absolute C is expressed as mg organismal C/litter bag.

| Sampling day | Fungi | Bacteria | Protozoans | Permanent meiofauna | Temporary meiofauna | Macroinvertebrates | Total organismal C |
|--------------|-------|----------|------------|--------------------|--------------------|--------------------|--------------------|
| 1            | 92.68 | 0.33     | 0.09       | 0.03               | 0.03               | 6.84               | 142.95             |
| 7            | 59.80 | 6.18     | 2.11       | 1.87               | 0.58               | 29.46              | 13.29              |
| 17           | 93.65 | 2.29     | 0.01       | 0.02               | 2.35               | 12.11              | 65.61              |
| 28           | 88.64 | 0.81     | 0.08       | 0.10               | 4.33               | 6.03               | 184.63             |
| 58           | 86.95 | 0.35     | 0.01       | 0.06               | 0.51               | 12.11              | 404.56             |
| 93           | 32.51 | 0.43     | 0.04       | 0.08               | 8.45               | 58.49              | 71.42              |
| 112          | 41.28 | 0.26     | 0.01       | 0.11               | 3.28               | 55.05              | 127.71             |
Concluding remarks

Our results indicate that colonization of decaying leaf material is highly dependent on hydrology and temperature. In Mediterranean forested streams, the occurrence of floods during organic matter accumulation in the streambed could shape the colonization (and subsequent decomposition) process and be essential to ecosystem functioning. After the flood, almost 70 d were necessary to restore the C budget values in the different compartments to those achieved during the first 10 d of colonization. The Mediterranean region is expected to endure higher temperatures and lower precipitation, especially in summer, under predicted climatic changes (IPCC 2007). Under these conditions, the quantity of leaves in the streambed could increase, as happens now in the driest years (Sabater et al. 2001, Acuña et al. 2005), and colonization and decomposition dynamics might be accelerated with the first autumn rains. Changes in patterns of precipitation could alter the duration, frequency, and magnitude of flow and flood pulses. Modification of the inundation regime is expected to decelerate breakdown rates and reduce breakdown heterogeneity—and both factors influence the decomposition process (Langhans and Tockner 2006). The influence of these changes (temperature and flow) on the C budget during leaf-litter processing in the Mediterranean headwater streams remains to be seen.

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