Tributary effects on the ecological responses of a regulated river to experimental floods

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ABSTRACT

Rivers regulated by dams display several ecosystem alterations due to modified flow and sediment regimes. Downstream from a dam, ecosystem degradation occurs because of reduced disturbance, mostly derived from limitations on flow variability and sediment supply. In the last decade, most flow restoration/dam impact mitigation was oriented towards the development of environmental flows. Flow variability (and consequent disturbance) can be reintroduced by releasing artificial high flows (experimental floods). Flow-sediment interactions during experimental floods represent strong ecosystem drivers, influencing nutrient dynamics, and metabolic and functional properties. In river networks, sediment and water inputs from tributaries generate points of discontinuity that can drive major changes in environmental conditions, affecting habitat structure and determining functional differences between upstream and downstream. However, despite the relevance for management, flow/sediment relations during environmental flows – and more importantly during experimental floods – remain poorly understood, mostly due to the lack of empirical evidence. In this study, we examined how a major tributary (source of water and sediments) modified the physical habitat template of a regulated river, thereby influencing ecological and geomorphological responses to experimental floods. Methods combined high-resolution drone mapping techniques with a wide range of biological samples collected in field surveys before, during, and after experimental floods in an alpine river. Data were used to quantify changes in relevant functional and structural ecosystem properties, relating ecological responses to geomorphological dynamics. Results highlight the importance of tributaries in restoring ecosystem properties lost after damming, enhancing the resilience of the system. In addition, we observed that disturbance legacy played a fundamental role in determining ecological conditions of a river prior to experimental floods, thus confirming that considering flow variability and sediment availability is crucial in adaptive dam management and environmental flows design.

1. Introduction

A rivers’ natural flow regime is characterized by fluctuations in discharge between low flows/droughts and high flows/floods, tightly coupled with precipitation/snowmelt events, generally with some degree of seasonal predictability (Poff et al., 1997). Rivers also have a natural sediment regime, which depends on the geological configuration and topography of the catchment (Wohl et al., 2015). Floods are among the main sources of natural disturbance in streams (Lake, 2000) and function as an important connection between flow and sediment regimes, controlling stream geomorphological dynamics and ensuring lateral connectivity with adjacent floodplains. The erosional forces by flow and sediment during a flood play a key role in sustaining the natural heterogeneity of fluvial habitat mosaics (Stanford et al., 2005), determining the spatial and temporal variation in the distribution of nutrients, organic resources, and habitat suitability for stream organisms, thereby influencing all trophic levels of aquatic food webs (Robinson et al., 2002). During floods, organic matter and biota are...

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transported and redistributed along the river network (Golladay et al., 1987; Small et al., 2008). Specific life-history traits allow stream and terrestrial organisms to benefit from floods—and low flow windows between floods (Lytle and Poff, 2004; Poiani, 2006), as well as providing resistance and resilience to extreme events (e.g., Robinson, 2012). Many aquatic organisms rely on floods for reproductive or dispersal cues (e.g., Jimenez-Segura et al., 2010), and fish spawning and fry-rearing habitats are in many cases dependent on the seasonal pulse of floods (Chapman, 1988; Junk et al., 1989; Melis, 2011).

Nowadays, many rivers are heavily regulated, modifying natural flow and sediment regimes. River regulation contributes greatly to the global crisis of freshwater ecosystems (Dudgeon et al., 2006; Reid et al., 2019), where severe hydropower-related morphological and hydrological modifications add to stressors related to other human activities (Malmqvist and Rundle, 2002). The scale of this alteration is testified by the proliferation of studies on the effects of river regulation at a global level (e.g., Maavara et al., 2020; Turgeon et al., 2019; Vörösmarty et al., 2003). The rush towards sustainable energy production, coupled with future uncertainty in water security due to climate change, has strongly increased the strategic function of dams (Vörösmarty et al., 2010).

The number of large dams constructed is increasing, now over 55,000 worldwide, especially in developing countries (Winemiller et al., 2016; Zarfl et al., 2015), and only slightly compensated by removal of old and obsolete structures (O’Connor et al., 2015). The main alterations of the hydrological regime and sediment regime caused by dams are a reduction in flow variability and water availability, and sediment supply (Poff et al., 1997). Modified flow and sediment regimes alter the physical habitat template of a river downstream of a dam (Ward and Stanford, 1983). For instance, the combined effect of disruption in flow variability and sediment transport reduces flow-generated disturbance, increases streambed stabilization and promotes channel incision, resulting in physical habitat degradation. As a response to new habitat conditions, the biotic properties of a river undergo major structural and functional adjustments (Bunn and Arthington, 2002). For instance, flow/sediment regulation influences rates of downstream metabolic processes (Aristi et al., 2014) and resource availability (e.g., increased algal growth, Lessard et al., 2013). Further, native riparian and aquatic organisms are affected during important life stages, thus species preferring stable conditions become dominant, and invasion by alien species is facilitated (Bunn and Arthington, 2002).

There is growing awareness of the importance of interactions and relationships between hydrological, geomorphological, and ecological processes (Grabowski and Gurnell, 2016) to face the profound environmental modification and ecological consequences of river regulation (Poff, 2018). In adaptive dam management, environmental flows aim at targeting functional components of the natural flow regime (i.e., functional flows after Yarnell et al., 2015) link flow dynamics to ecosystem processes. Flow experiments (Konrad et al., 2011; Olden et al., 2014) or managed environmental flows (Gillespie et al., 2015) have been increasingly designed and implemented to modify dam water-release schemes to restore river geomorphology and ecological properties. For example, stable residual flows from dams can be mitigated by the implementation of experimental floods, which are outflow manipulations aimed at reintroducing elements of flow variability in regulated rivers (Robinson et al., 2018). Experimental floods can be used as management actions to enhance geomorphological/ecological conditions below dams and serve as large-scale flood experiments to improve the understanding of geomorphological/ecological response mechanisms to such disturbances (Konrad et al., 2011). Prime examples of long-term experimental flood applications are those of the Colorado (Cross et al., 2011) and Spöl River (Robinson et al., 2018). These studies provide a foundation on the topic, showing that ecosystem responses to extreme flow manipulations depend on the time-frame, status of abiotic components, and the adaptation of organisms to flood disturbance.

Tributaries can have important roles in determining different upstream/downstream responses to flow experiments (Vinson, 2001). In fact, unregulated tributaries are points of discontinuity that can generate major changes in the environmental conditions of a river (Vannote et al., 1980), and can revert or mitigate some of the impacts of flow regulation (Milner et al., 2019; Ward and Stanford, 1983). Tributaries can introduce substantial amounts of water, sediments, and organic matter that partially restore flow variability, and function as recruitment sources of stream organisms (Rice et al., 2001; Rice et al., 2006). Sediment and water inputs from tributaries can modify key aspects of the physical template of streams, determining changes in bottom hydraulics and substrate stability, thus generating an ideal environmental gradient to study flow-sediment-ecology relationships and responses to experimental floods.

The identification of specific patterns of eco-morphological response at different time resolutions (e.g., immediate, short term, long term) to flow manipulation is essential for water managers to generate explicit endpoints for an ecologically sound flow/sediment restoration (Harrison et al., 2017; Kieman et al., 2012; Robinson et al., 2004a, 2015; Thompson et al., 2018). The understanding of mechanistic relationships between flow restoration and environmental/ecological responses is still insufficient to link restoration methods and goals. Environmental flow planning needs to be objective-driven (Davies et al., 2012) as outcomes are likely to be context-dependent (Gillespie et al., 2015; Kevic et al., 2018; Olden et al., 2014). Further, the need to understand how sediment availability influences ecological and morphological responses to flow manipulations is motivated by a risk of unsuccessful or even deleterious flow restorations if evaluations on sediments are neglected (Wohl et al., 2015).

Our study aimed at measuring ecological and morphological responses to experimental floods in a regulated river, as influenced by an unregulated tributary. We compared ecosystem baseline, response, and recovery properties after two experimental floods above and below a tributary input, and assessed the immediate ecological response patterns during each flood. We expected the two reaches to respond differently, and to relate the differences to disturbance frequency and sediment supply. We hypothesized that ecological components below the tributary will show attributes of resistance and resilience to floods disturbance. Our ultimate goal was to improve the understanding of ecosystem responses and eco-morphological relationships during experimental floods and to highlight the importance of periodic disturbance in maintaining the ecological properties of regulated rivers.

2. Methods

2.1. Study site

The Spöl catchment (295 km²) is located in the central Alps as part of the Danube basin (Fig. 1). Some ~80% of the catchment lies in Stelvio National Park, Italy, and the Swiss National Park (SNP), Switzerland. The Spöl is 28-km long, originating from Forcola di Livigno at 2315 m above sea level, flows into the Inn River at Zernez, Switzerland. From 1960 to 1970, Punt dal Gall (164M m³ reservoir capacity) and Ova Spin (6.2M m³ reservoir capacity) dams were built along its course for energy production (Scheurer and Molinari, 2003). Before regulation, the Spöl had a glacio-nival regime with winter low flows and higher flows during the other seasons, including floods generated by rainfall events (see hydrological data presented in Robinson et al., 2018). Due to the eco-morphological impacts caused by the dams, a board composed of members of the Swiss National Park, the hydropower company Engadiner Kraftwerke (EKW), Federal and Cantonal authorities agreed to an experimental flood program starting in 2000. The flood program aimed to reduce colmatation, increase substrate porosity by reducing periphyton levels and flushing fine sediments from the riverbed, and restoring channel morphology. Since 2000, floods have been implemented regularly once or twice each year, with controlled water releases from Ova Spin and Punt dal Gall (Kevic et al., 2018; Robinson et al., 2018). This study was carried out on the lower segment of the river, from...
The tributary, Ova da Cluozza, is a left bank unregulated tributary of the lower Spöl with a catchment of 27 km² (Fig. 1). Its flow regime is characterized by marked seasonality of winter low flows and frequent summer spates (Fig. 2). During baseflow, it adds 0.2–0.8 m³/s to the Spöl discharge (~0.4 m³/s winter residual flow; ~0.9 m³/s summer residual flow), while during periods of snowmelt and rainfall, it generates floods of 4–8 m³/s (see Fig. 2) and adds substantial inputs of sediment. This offers the opportunity to study how sediment dynamics vary and how ecological resistance and resilience to experimental floods differ on the river Spöl above and below the confluence. The two reaches (upstream and downstream of the confluence) have comparable water physico-chemical conditions but are different in morphology, streambed structure, and to some extent substrate composition (Mathers et al., 2021). The upper 3.5-km long section flows between Ova Spin dam and the junction with Ova da Cluozza through a mainly confined valley. Trees (mostly Pinus sp.) cover the steep banks, and many side scree slopes are present along the reach. A small tributary (Ova da Lasciadura) enters at around halfway in the upper reach. The study reach is characterized mostly by a single thread channel with riffles, pools, and small glides. The lower section is ~2.5-km long and flows from the junction with Ova da Cluozza to the Inn. Riparian vegetation is composed of trees (mostly Pinus sp. and Alnus sp.) and pasture near the town of Zernez. This section flows through an open valley, with vegetated islands and long bars, where the channel often braids forming riffles and glides.

Two experimental floods were released (September 4, 2018, July 19,
2019) from Ova Spin dam from the outlet located at the bottom of the dam (Scheurer and Molinari, 2003). Duration of both floods was ~8 h with a maximum discharge of 25 m$^3$/s maintained for ~2 h. Volumes of water released during the floods were $4.6 \times 10^5$ m$^3$ (2018) and $5.1 \times 10^5$ m$^3$ (2019). Peak discharges were in the typical range of seasonal high flows in the system before impoundment for autumn rains (Sept. 2018) and snowmelt (July 2019) (Robinson and Uehlinger, 2008). The floods had similar magnitude but different release patterns, with the 2018 flood characterized by a steeper rising limb and shorter duration (see hydrographs in Fig. 2). Samples were collected before (BF), during, after (AF), and four weeks after each flood (LAF) to observe patterns of response of a series of ecological parameters. BF, AF, LAF sampling was carried out at four locations, two above (C, D) and two below (A, B) the junction with the tributary (Fig. 1). Sampling during the floods was carried out at two locations, one above (D) and one below (A, B) the confluence. In 2019, three days after the study flood, EKW, and SNP carried out at two locations, one above (D) and most downstream (A) sites were analyzed. In total, 21 samples were collected at each sampling site in 2018 and, respectively, 25 (site A) and 21 (site D) in 2019. In addition, two drift samples were collected at both sites the day after (morning/afternoon) and one four weeks after the second flood.

Seston/drift samples were stored in plastic bags and frozen at −20 °C until analysis. In the laboratory, macroinvertebrates were separated from organic matter using a dissecting microscope (10X) and preserved in 70% ethanol. Some large samples were subsampled by half or a quarter before sorting. Macroinvertebrates were identified to the lowest practical taxonomic level (mostly genus) using Tachet et al. (2010). Large pieces of wood were separated from the remaining organic matter before analysis of dry mass. Organic matter content was obtained by measuring ash-free dry mass (AFDM) by drying at 60 °C for 48h, incinerating at 500 °C for 5 h, and then recording the difference in weights. To quantify transport during the flood, the number of invertebrates transported, and organic matter biomass were standardized for the water volume filtered for each sample and expressed, respectively, as gAFDM/m$^3$ and N/m$^3$.
of Baetidae (active swimmers), Heptageniidae and Rhyacophilidae (clingers), Chironomidae (passive), Gammaridae (swimmer/passive), and Nemouridae (crawler/passive). Their relative abundances were plotted against discharge to identify trait patterns in functional response to floods.

2.2.4. Data deposition

Physico-chemical, seston, and drift data available on request from Zenodo (Consoli et al., 2021).

2.3. Measures collected before/after each flood

2.3.1. Benthic samples and sediment respiration

Quantitative benthic samples of organic matter and macroinvertebrates were collected at three random points at each site on each sampling date using a Hess sampler (sampling surface of 0.045 m², 250µ mesh), and stored in 70% ethanol. Macroinvertebrates were sorted from organic matter and identified as above. Organic matter was separated into coarse and fine particulate organic matter (FPOM < 1 mm < CPOM) and analyzed as ash-free dry mass (g/m²).

Five stones were collected at each site on each sampling date for measuring periphyton biomass and frozen at ~20 °C until analysis. Measurements of the two main axes (a-, b-axis) were taken for each stone to calculate the surface sampled using the equation a·b·π/4. Periphyton was scraped from the stone surface using a metal brush and rinsed with water. The sample volume was recorded, and a subsample was filtered through a glass fiber filter (Whatman GF/F). Filters were dried at 60 °C, weighed, ashed at 500 °C for 3 h, and reweighed to estimate periphyton biomass as AFDM (gAFDM/m²).

Hyporheic sediment respiration was measured before and after each flood to evaluate microbial responses to flooding. In situ respiration chambers were used to measure oxygen depletion after Uehlinger et al. (2002). Four replicates per location were used to carry out the experiment. The upper layer of sediment (~10 cm) was removed before sample collection to exclude the effect of primary producers on respiration. Sediments were sieved with an 8-mm sieve to standardize measures following Uehlinger et al. (2003). Plexiglas® tubes (678 cm³) were half-filled with sediments, then filled with stream water and sealed. Tubes were incubated in darkness for at least 4 h in the stream channel at each site. Dissolved oxygen content and temperature were measured before and after incubation with a dissolved oxygen meter (Hach HQ40d equipped with LD0101 oxygen probe). After measurement, the content of each tube was stored in plastic bags and kept at ~20 °C until analyzed for sediment volume, grain size distribution, and loose (LPOM, expressed as gAFDM/l) and attached particulate organic matter (APOM, expressed as g/kgsediment). Oxygen depletion was standardized for temperature, sediment weight, and organic matter content.

2.3.2. Data analysis

Results from sediment respiration, periphyton, CPOM, FPOM, and macroinvertebrate density from benthic samples were grouped relative to their location (A, B downstream; C, D upstream), and the mean and standard error then calculated. A two-way ANOVA was used to test individual responses to experimental floods between upstream and downstream sites (factors: reach, sampling date, and sampling date × reach) on (log -1) transformed data to meet assumptions. Of interest for this study were reach and the interaction between sampling date and reach. In the case of significant results (alpha = 0.05), Tukey's HSD test was used to identify significant differences.

2.3.3. Drone surveys

Drone surveys were flown over the study segment (Fig. 1) before and after each flood, under baseflow conditions, to collect data (images) to assess morphological changes following the floods. A Wingtra One drone equipped with a high-resolution RGB camera (Sony RX1; 42 MP) and an Intel® Falcon™ 8 with a high-resolution RGB camera (Sony NEX-7; 24.3 MP) were used, respectively, in 2018 and 2019. Ground control points measured with a dGPS (n = 10 in 2018; n = 50 in 2019) were used to reach high accuracies in the rectification of the ortho-images. Generation of orthophotos was done with the Software Pix4D. The software relied on automatic feature detection and matching algorithms to retrieve the internal and external orientation of oblique images. Photogrammetric processing subsequently provided dense point clouds, digital elevation models (DEM), and ortho-images. The resolution of ortho-images was < 3 cm.

Large-scale morphological responses to each flood were tracked on the aerial images in QGIS 3.10.2 by separating the following macro-unit surfaces within the bankfull area: emerged sediment, wet channel, and instream vegetation (Geomorphic Units Survey (GUS) — broad level, Belletti et al., 2017). Polygons were hand-drawn in a vector layer to extract the area of each macro-unit type to estimate changes in mean size, variance, and relative abundance before and after a flood. Macro-units were identified at a scale of 1:250, and a scale of 1:125 was used in cases of difficult or ambiguous interpretation. Four maps (Fig. S1 in supplementary material) were generated as follows: before 2018 flood, after 2018 flood, after 2019 first flood, after 2019 s flood. Reach length was of ~800 m upstream and ~1500 m downstream. Based on these results, ratios between wet/dry surfaces were calculated for each map, and comparisons between reaches were carried out by adjusting macro-units surfaces to reach length (upstream/downstream).

Reach-scale changes in geomorphic units assemblage over the study period were assessed by applying the GUS method to the basic level. In this case, three sub-reaches of interest of ~500 m were identified, one upstream and two downstream of the confluence (mid and lower); see Fig. S1 in supplementary material for the location of the sub-reaches. Polygons were hand-drawn on a vector layer to extract the area of each geomorphic unit and sub-unit types relevant to the habitat level, to estimate changes in relative abundance of each geomorphic unit after the three floods (i.e., before 2018 flood and after the second 2019 flood). The following geomorphic units and sub-units were identified at a scale of 1:125, zooming in when necessary: rapid, riffle, pool/backwater, glide, deposition of boulders/cobbles (S – partially submerged, E – emerged), mid channel bar, bank attached bar, vegetated bar, vegetated island (shrubs and trees), trees in active channel. Two sets of maps were generated as follows: before 2018 flood, after 2019 s flood (Fig. 7). To estimate changes in channel width, measurements were taken every 5 m, perpendicular to a centerline with the Geometric Attributes tool (Nyberg et al., 2015) in QGIS 3.10.2, and mean and standard deviation were calculated. For each sub-reach wet/dry surface changes of the active channel were calculated based on geomorphic units areas. Change in surface occupied from each unit was calculated as the difference between maps and shown in Fig. 7.

3. Results

3.1. Immediate responses to the floods

3.1.1. Physico-chemical responses during the floods

Patterns of temperature, conductivity, total nitrogen, and total phosphorus showed minimal differences between upstream and downstream sites, but notable differences between floods (Fig. 3). During the 2018 flood, changes in temperature were in the range of ~1.5 °C. Water baseline temperature was ~8 °C at both reaches. Temperature increased by ~0.5 °C (upstream) and 1.0 °C (downstream) during the early phases of the flood, likely due to the presence of cooler water layers inside the reservoir. Upstream, temperature stabilized around 8.3 °C and slowly increased to 9 °C during the falling limb. Downstream, water temperature remained stable at around 9 °C during the flood, with slight oscillations of ±0.3 °C. During the 2019 floods, the patterns of response were comparable between reaches, (downstream temperatures shifted ~1 °C due to the input of cooler water from the tributary), and showed a gradual increase in temperature consistent with the daily solar heating,
as confirmed by the measurements taken the day after the flood. In this case, water temperature inside the reservoir did not seem to differ from water temperature in the stream.

Conductivity showed comparable patterns between reaches, but differences between years (Fig. 3). Water input from Ova da Cluozza added $+10 \mu S/cm^2$ to downstream waters. In 2018, water release from the dam caused a decrease from baseline values between 270 and 280 to $200-210 \mu S/cm^2$. Downstream, we measured a drop in conductivity to a minimum value of $185 \mu S/cm^2$ at maximum discharge, which corresponded to the second turbidity peak (see below) and an increase of $\sim0.5 ^\circ C$ in temperature. In 2019, baseline values of $200-210 \mu S/cm^2$ decreased to $150-160 \mu S/cm^2$ during the flood at both reaches, then returned to pre-flood values as soon as discharge started decreasing.

Total phosphorus (TP) sharply increased at both reaches during each flood at an early stage of the rising limb (Fig. 3). Maximum TP differed between reaches, decreasing by $\sim$ half below the tributary (11.6–6.8 mg/l in 2018, 5.5 to 3.2 mg/l in 2019). This increase on the rising limb also was observed for total nitrogen (TN), which showed a sharp increase in 2018 at both reaches (with peaks around 2.2–2.3 mg/l), while values were always below 1.0 mg/l in 2019.

Turbidity quickly exceeded the instrument detection threshold (6000 NTUs) in 2018 at both reaches during the rising limb of the flood (Fig. 3). This increase was related to fine sediments being released from the dam along with fine sediments and organic matter being mobilized from the streambed in response to the increase in discharge. Response patterns between reaches were compatible with the flood wave moving downstream. Secondary turbidity peaks were observed at both reaches as well. In 2019, turbidity responded similarly to 2018 at the upper reach. In contrast, we observed two marked peaks below the tributary in 2019, the first with the increase in flow from the flood, the second as discharge increased to the maximum release value of 25 m$^3$/s.

3.1.2. Drift and seston during the floods
Transported organic matter (seston) and macroinvertebrates (drift) showed corresponding patterns during both floods, where benthic organic matter and macroinvertebrates were entrained in the flow as the
water level increased (Fig. 3). In 2018, early peaks in seston and drift occurred simultaneously with peaks in turbidity, TP, and TN; i.e., during the early stages of the flood at ~5–10 m³/s discharge. Above the tributary, the first seston (~60 g/m³) and drift (~800 ind/m³) peak was followed shortly after by a second, less marked peak (~30 g/m³ and ~400 ind/m³, respectively) that corresponded to the maximum food discharge. Below the tributary, seston and drift showed comparable response patterns (first peak of ~15 g/m³ and ~200 ind/m³; second peak of ~3 g/m³ and ~50 ind/m³, respectively). The slower rising limb in 2019 resulted in a more gradual response in seston and drift than in 2018. Above the tributary, a double peak in drift (~500 and ~600 ind/m³, respectively) followed the first two discharge increments (~10–20 m³/s) and dropped shortly after to <100 ind/m³. Seston concentration reached ~30 g/m³ during the first increment and remained at 15–25 g/m³, showing two additional peaks with the second and last increase in discharge.

The total organic matter transported during the floods was 3197 and 4938 kg above the tributary, and 699 and 3044 kg below the tributary, in 2018 and 2019, respectively (Fig. 3). Total macroinvertebrate drift above the tributary was around 45–48 × 10⁶ individuals during both floods (Table 1). Below the tributary in 2019, we recorded almost double the number of drifting macroinvertebrates than in 2018 (~13 and ~23 × 10⁶ in 2018 and 2019, respectively) (Table 1). Gammaridae represented the largest fraction in the drift, being around 50% or higher at all sites in both floods. The remaining drift was composed of Chironomidae, Leuctridae, Baetidae, and Ephemeroptera other than Baetidae (mostly Heptageniidae). Lastly, Gammaridae and Chironomidae drift patterns corresponded to those of seston, reflecting their inclination to passive drift (Fig. 4a and c). Baetidae, Nemouridae, and Heptageniidae/Rhyacophilidae, in contrast, showed both a passive response to increasing discharge and an active response as discharge receded (Fig. 4b and d). However, this response was not uniform and mediated by location, abundance, and shape of the flood hydrograph.

Background drift for the entire study section was calculated based on the first three driftsamples from 2018 from both reaches, first three drift samples from 2019 (only downstream, as the first sample from upstream was collected as the flood already started – Fig. 3), and from three drift samples collected four weeks after the flood at both reaches. Results show very low background drift densities (1.7 ± 0.5 N/m³), mostly constituted by Chironomidae (1.33 ± 0.45 N/m³), Simuliidae (0.12 ± 0.04 N/m³), and Baetidae (0.1 ± 0.03 N/m³), with the sporadic presence of other taxa.

3.2. Before/after effects of the floods

3.2.1. Benthic samples and sediment respiration

ANOVA test on periphyton data revealed significant differences for date (p < 0.001) and the interaction date × reach (p < 0.001). Before the flood in 2018, periphyton biomass was on average higher above than below the tributary (16.8 vs 10.7 gAFDM/m², respectively) (Fig. 5a), albeit not significantly (Tukey’s HSD p = 0.09). The 2018 flood significantly reduced periphyton biomass in upstream sites to ~ half (7.6 gAFDM/m²; p < 0.001), and levels returned to pre-flood conditions within four weeks (16.7 gAFDM/m²; AF-LAF p < 0.01; BF-LAF p = 1). Below the tributary, periphyton biomass remained at similar levels for the duration of the study, although slightly increasing after the flood (15.8 gAFDM/m²) and returned to an intermediate level (13.7 gAFDM/m²) within four weeks. In 2019, patterns were similar at both reaches with no significant differences among sampling dates (Fig. 5a). Before the 2019 flood, periphyton biomass was significantly different to that found before the 2018 flood (Tukey’s HSD p = 0.01 upstream and p < 0.01 downstream), approximately 2/3 lower (5.1 gAFDM/m² upstream and 4.2 downstream). There was no effect of the flood on periphyton biomass in 2019 at both reaches, and biomass almost doubled four weeks after the flood to 9.8 and 7.3 gAFDM/m², respectively, upstream and downstream (Tukey’s HSD test AF-LAF, p < 0.01 and p < 0.01, respectively), following higher growth rates during summer months. Periphyton biomass upstream LAF was also significantly different from BF (Tukey’s HSD p < 0.01).

The ANOVA test for benthic CPOM showed no significant difference related to date, reach, or their interaction. CPOM response pattern varied greatly between sites and floods (Fig. 5b). Benthic FPOM patterns were similar to those of periphyton (Fig. 5c). ANOVA test revealed significant differences for date (p < 0.001) and the interaction date × reach (p < 0.05). Upstream, the 2018 flood reduced FPOM (Tukey’s HSD BF-AF, p < 0.001), then it increased four weeks after the flood (Tukey’s HSD AF-LAF, p < 0.05). Downstream in 2018, FPOM levels remained relatively stable between 1.2 and 1.6 gAFDM/m². In 2019, both reaches show an increase in FPOM over time, and the flood had no significant effect.

We found that macroinvertebrate density differences were significant for date (p < 0.001) and reach (p < 0.01) (Fig. 5d). Before the 2018 flood, Tukey’s HSD test revealed differences between sites (p < 0.05), with higher densities upstream. The impact of the flood was most evident upstream, where density went from ~5000 to ~100 ind/m², but density (at ~7000 ind/m²) recovered to or exceeded pre-flood levels within four weeks (Tukey’s HSD BF-AF p < 0.001; AF-LAF, p < 0.001; BF-LAF, p = 1.0). Downstream showed a similar pattern, however, the impact of the flood was smaller (~1000–~250 ind/m²) and not significant, and density reached ~4000 ind/m² within four weeks after the flood (Tukey’s HSD AF-LAF, p < 0.001). Before the 2019 flood, baseline densities were extremely low at both sites and significantly different to those found in 2018 (~200 and ~50 ind/m², respectively, upstream and downstream; tukey’s HSD in both cases p < 0.01), and remained comparably low after the flood (~400 and ~80 ind/m²). Four weeks after the flood, densities were significantly higher (~8000 and ~2300 ind/m²) (Tukey’s HSD upstream AF-LAF and BF-LAF, both p < 0.01; downstream AF-LAF, p < 0.01 and BF-LAF, p < 0.001).

| Order       | Taxa          | 2018 Upstream | 2019 Upstream | 2018 Downstream | 2019 Downstream |
|-------------|---------------|---------------|---------------|----------------|-----------------|
|             | N (×10⁶)      | %             | N (×10⁶)      | %              | N (×10⁶)        | %              |
| Amphipoda   | Gammaridae    | 28.8          | 62.8          | 6.6            | 31.1            | 24.9           | 51.5           | 11.4           | 49.6           |
|             | Chironomidae  | 5.9           | 12.8          | 1.3            | 10.1            | 11.0           | 22.9           | 2.8            | 12.2           |
|             | Other Dipt.   | 0.6           | 1.4           | 0.5            | 4.2             | 0.2            | 0.3            | 0.1            | 0.6            |
| Ephemeroptera | Baetidae     | 1.5           | 3.2           | 1.6            | 12.3            | 4.2            | 8.7            | 1.1            | 4.8            |
|             | Other Ephem.  | 6.4           | 14.0          | 1.5            | 12.0            | 1.0            | 2.0            | 1.3            | 5.6            |
| Plecoptera  | Lamellibranchiidae | 0.2       | 0.4           | 0.1            | 0.5             | 4.4            | 9.1            | 2.3            | 9.8            |
|             | Leuctridae    | 0.9           | 1.9           | 0.8            | 6.0             | 1.7            | 3.5            | 2.5            | 10.7           |
|             | Other Plecopt. | 0.6       | 1.2           | 0.1            | 0.8             | 0.0            | 0.0            | 0.2            | 0.8            |
|             | Other taxa    | 1.1           | 2.3           | 0.4            | 3.0             | 0.9            | 1.9            | 1.4            | 5.9            |
| Total       |               | 46.0          | 100           | 12.9           | 100             | 48.3           | 100            | 23.1           | 100            |
Macroinvertebrate community composition (Table 2) varied greatly between reaches before the flood in 2018, with marked dominance of Gammaridae in the upstream reach, as opposed with the higher occurrence of Heptageniidae, Baetidae and Plecoptera downstream. The flood had an impact on taxa richness in upstream riffle habitats, while it remained substantially unchanged downstream. Four weeks after the flood, communities recovered to a similar pre-flood composition, with a notable increase of Leuctridae density in both reaches, and Simuliidae downstream. In 2019, pre-flood community composition and density were strongly influenced by the snowmelt-induced disturbance in both reaches. Community composition was substantially left unchanged by the flood, with some density variations. After four weeks, community composition was dominated at both sites by Chironomidae and Simuliidae. This determined notable differences in relative densities from those observed before the 2018 flood, at a similar time (end of August – beginning of September).

Sediment respiration responses to the floods differed significantly among dates (p < 0.001), reaches (p < 0.001), and their interaction (p < 0.001) (Fig. 6a). Before the flood in 2018, sediment respiration was, on average, four times higher upstream than downstream (Tukey’s HSD p < 0.05). Sediment respiration upstream of the tributary was reduced by the flood, although not significantly (Tukey’s HSD p = 0.45), while it remained the same downstream (Tukey’s HSD p = 0.99). In 2019, sediment respiration in the upstream site was significantly lower than that found before 2018 flood (Tukey’s HSD p < 0.001). The 2019 flood enhanced sediment respiration at both reaches, albeit significantly only upstream (Tukey’s HSD p < 0.05). Concentration of LPOM differed significantly among dates (p < 0.001), while APOM differed significantly among reaches (p < 0.01) and dates (p < 0.001). In 2018, before the flood, the loose organic matter content (LPOM) of sediments upstream was double that of downstream sediment, although not significantly (Tukey’s HSD p = 0.16); after the flood, levels were similar in both reaches (Fig. 6c). The flood significantly reduced LPOM content at both sites (respectively, Tukey’s HSD p < 0.001 upstream and p < 0.05 downstream), while attached organic matter (APOM) decreased at both sites after the flood, although not significantly (Fig. 6b). In 2019, both reaches had similarly low LPOM concentrations (Fig. 6c), which differed significantly from those in 2018 before the flood only in the upstream reach (Tukey’s HSD p < 0.001). LPOM concentrations remained similar after the 2019 flood (all Tukey’s HSD p ~ 0.95). This flood significantly reduced only APOM in the upstream reach (p < 0.01). Concentrations of APOM before 2018 and 2019 floods were comparable at both sites. The content of APOM in sediments was generally comparable between reaches and before each flood (min = 4.18; max = 5.66), and was reduced ~ 1 g/kg after the floods (Fig. 6b). We found that respiration and LPOM concentrations were highly correlated before the floods (r = 0.57, p < 0.001) but not after (r = 0.02, p = 0.89), and that respiration was negatively correlated with APOM after the flood (r = −0.52, p < 0.01).
Table 2
Benthic macroinvertebrate composition expressed as mean density (N/m²) ± standard error and relative abundance (%) before (BF), after (AF) and four weeks after (LAF) the floods in 2018 and 2019, upstream and downstream of the tributary.

| Taxa          | 2018 Upstream |     |     |     | 2019 Upstream |     |     |     | 2019 Downstream |     |     |     |
|---------------|---------------|-----|-----|-----|---------------|-----|-----|-----|---------------|-----|-----|-----|
|               | BF (AF)       | BF  | AF  | LAF | BF (AF)       | BF  | AF  | LAF | BF (AF)       | BF  | AF  | LAF |
| Gammaridae    | 2119.3 ± 822.4 (43.4%) | 58.7 ± 19.2 (51.6%) | 3127.7 ± 997.6 (46.2%) | 110.0 ± 50.8 (9.3%) | 44.0 ± 32.4 (17.1%) | 168.7 ± 98.7 (4.1%) | 0.5 ± 0.5 (5%) | 2.0 ± 1.0 (10.7%) | 330.0 ± 106.8 (4.2%) | 0.3 ± 0.2 (16.7%) | 0.5 ± 0.2 (13.0%) | 7.3 ± 6.7 (0.3%) |
| Baetidae      | 843.3 ± 337.1 (17.3%) | 7.3 ± 4.2 (6.4%) | 330 ± 125.8 (4.9%) | 326.3 ± 124.3 (27.5%) | 47.7 ± 21.6 (18.6%) | 425.3 ± 99.6 (10.2%) | 0 | 0 | 550.0 ± 162.8 (7.0%) | 0 | 0 | 124.7 ± 24.1 (5.5%) |
| Heptagenidae  | 117.3 ± 27.3 (24.1%) | 14.7 ± 6.7 (12.9%) | 117.3 ± 31.8 (1.7%) | 223.7 ± 79.7 (18.8%) | 40.3 ± 17.5 (15.7%) | 150.3 ± 19.0 (3.6%) | 2.3 ± 1.6 (23.3%) | 6.2 ± 2.8 (32.1%) | 77.0 ± 18.5 (1.0%) | 0.3 ± 0.2 (16.7%) | 1.2 ± 0.6 (30.4%) | 99.0 ± 18.5 (4.3%) |
| Rhyacophilid. | 36.7 ± 11.2 (6.4%) | 7.3 ± 4.2 (1.7%) | 260.3 ± 51.7 (3.8%) | 33.0 ± 18.5 (2.8%) | 7.3 ± 4.2 (2.9%) | 143 ± 37.7 (3.4%) | 0.2 ± 0.1 (1.7%) | 0.7 ± 0.6 (3.6%) | 40.3 ± 14.1 (0.5%) | 0 | 0 | 14.7 ± 8.5 (0.6%) |
| Other Nemouridae | 36.7 ± 11.2 (4.4%) | 0 | 29.3 ± 12.3 (0.4%) | 0 | 3.7 ± 3.3 (1.4%) | 0 | 0 | 0 | 0 | 14.7 ± 4.2 (0.2%) | 0 | 0 | 3.7 ± 3.3 (0.2%) |
| Leuctridae    | 451.0 ± 287.8 (9.2%) | 3.7 ± 3.3 (3.2%) | 1316.3 ± 416.6 (19.4%) | 51.3 ± 35.4 (4.3%) | 14.7 ± 6.7 (5.7%) | 1085.3 ± 312.3 (26.1%) | 0 | 0 | 234.7 ± 76.7 (3.0%) | 0 | 0 | 113.7 ± 29.5 (5.0%) |
| Other Plecopt. | 462.0 ± 132.4 (9.5%) | 0 | 315.3 ± 86.1 (4.7%) | 238.3 ± 140.7 (20.1%) | 7.3 ± 6.7 (2.8%) | 498.7 ± 122.5 (12.0%) | 0.8 ± 0.3 (8.3%) | 6.8 ± 3.0 (36.6%) | 66.0 ± 24.0 (8.8%) | 0.3 ± 0.2 (16.7%) | 1.0 ± 0.6 (16.6%) | 3.7 ± 3.3 (0.2%) |
| Simulidae     | 47.7 ± 21.0 (1%) | 0 | 234.7 ± 115.0 (3.5%) | 91.7 ± 30.9 (7.7%) | 11.0 ± 6.8 (4.3%) | 1063.5 ± 715.0 (25.6%) | 0 | 0 | 4825.3 ± 2452.1 (61.1%) | 0 | 0 | 458.3 ± 118.2 (20.1%) |
| Chironomidae  | 135.7 ± 43.82 (2.8%) | 0 | 652.7 ± 433.5 (9.7%) | 25.7 ± 13.1 (2.2%) | 51.3 ± 16.9 (20%) | 458.3 ± 83.2 (11.0%) | 1.0 ± 0.5 (10%) | 1.2 ± 0.4 (6.2%) | 1514.3 ± 366.1 (19.2%) | 0.5 ± 0.3 (25%) | 0.2 ± 0.1 (4.3%) | 1045.0 ± 101.3 (45.7%) |
| Other Diptera | 253.0 ± 58.15 (5.2%) | 14.7 ± 4.2 (12.9%) | 62.3 ± 12.1 (0.9%) | 40.3 ± 18.3 (3.4%) | 7.3 ± 4.2 (2.8%) | 139.3 ± 53.3 (3.3%) | 2.3 ± 0.8 (23.3%) | 1.3 ± 0.7 (7.1%) | 47.7 ± 12.1 (0.6%) | 0.2 ± 0.1 (8.3%) | 0.7 ± 0.2 (17.4%) | 66.0 ± 24.3 (2.9%) |
| Other taxa    | 165.0 ± 39.09 (3.4%) | 7.3 ± 4.2 (6.4%) | 319.0 ± 215.3 (4.7%) | 11.0 ± 6.9 (0.9%) | 18.3 ± 10.9 (7.1%) | 18.3 ± 13.1 (0.4%) | 2.8 ± 1.0 (28.3%) | 0.7 ± 0.4 (3.6%) | 176 ± 99.3 (2.2%) | 0.3 ± 0.2 (16.7%) | 0.3 ± 0.4 (8.7%) | 282.3 ± 117.5 (12.4%) |
but not before \((r = 0.13, p = 0.48)\). LPOM and APOM content were also significantly correlated before \((r = 0.64, p < 0.001)\) but not after the floods \((r = 0.01, p = 0.95)\).

### 3.2.2. Flood induced changes in channel morphology

The two reaches showed substantial baseline morphological and morphodynamics differences \((\text{Fig. S1 in supplementary material})\) in response to the experimental floods. Above the tributary, the reach is characterized by a narrow valley and showed signs of flow regulation and sediment deficit, with instream emergent sediment patches limited in extent and vegetated side bars disconnected from the active channel. Instream vegetation patches were limited in number and size \((\text{Table 3})\).
and mostly comprised of small shrubs and grass. With the floods, upstream emerged sediment and vegetation patches were relatively stable in terms of mean size, with limited changes in number and area. The analysis of geomorphic units in the upstream sub-reach confirmed no major morphological changes (Fig. 7, Table 4), and limited lateral dynamics. Active channel width and surface remained substantially unchanged after the floods, showing only small increases related to local redistribution of sediments and exposure of bank attached sediment patches (Fig. 7). Before the floods, the sub-reach was characterized by the presence of relatively large areas of slow-flow (pools/backwater), some of which were formed by large wood deposits, that were greatly reduced after the floods. Riffle surface was also reduced, in favour of glides, as a consequence of gravel-sized sediment deposition in the channel. This sub-reach was characterized by the presence of partly submerged boulder-cobble depositions that remained substantially unchanged after the floods.

In the downstream reach, the valley gradually opens up and lateral controls give room to the alluvial valley. Here, sediment is supplied by Ova da Cluozza, which is also responsible for enhanced channel dynamics downstream. A braided channel formed across large patches of emergent sediment in the form of bars and islands. Instream vegetation here is mostly represented by trees and shrubs on vegetated islands and patches (Fig. 7, Table 4), mostly related to lateral deposition. After the floods, riffle area declined in favour of rapids, while other submerged units area remained substantially unchanged. This riffle-rapid turnover was driven by erosive dynamics in the area surrounding one of the vegetated islands (Fig. 7), as indicated by the removal of the vegetated sediment surrounding the island.

The floods activated interesting lateral dynamics in the lower sub-reach as well, with the formation and stabilization of a lateral secondary channel on the left bank (Fig. 7), and the inundation of the right bank, with an associated increase of vegetation (trees) presence in the active channel. The formation of this secondary channel after the first 2019 flood, and consolidation after the second 2019 flood, increased the active channel width and area, with a sharp growth of submerged areas (glides and riffles) (Table 4 and Fig. 7). This was associated with the formation of two large mid-channel bars (Fig. 7).

### Table 3

| Sub-reach          | Before 2018 | After 2nd flood 2019 | Before 2019 | After 2nd flood 2019 | Before 2019 | After 2nd flood 2019 |
|--------------------|-------------|----------------------|-------------|----------------------|-------------|----------------------|
| Mean area ± sd (m²) | 19 ± 52     | 12 ± 4               | 245 ± 2957  | 17 ± 59              | 165 ± 2751  | 23 ± 65              |
| N (tot m²)         | 140 (2635)  | 22 (255)             | 8 (118)     | 10 ± 25              | 28 (288)    | 9 ± 12               |
| Upstream sediment  |             |                      |             |                      |             |                      |
| Upstream vegetation| 98 ± 152    | 143                  | 186         | 131 ± 416            | 117         | 170 ± 492            |
| Downstream sediment|             |                      |             |                      |             |                      |
| Downstream vegetation| 184 ± 349 | 15 (2764)            | 73 ± 198    | 61 ± 160             | 89 (5434)   | 74 ± 165             |
| Downstream channel |             |                      |             |                      |             |                      |
| Downstream channel | 8937        | 8878                 | 9365        | 27618                |             | 25482                |

### Table 4

| Sub-reach          | Before flood 2018 | After 2nd flood 2019 | Area (m²) | Submerged/emerged |
|--------------------|-------------------|----------------------|-----------|-------------------|
| Sub-reach length (m) | 486               | 546                  |           |                   |
| Width (m) mean ± sd | 14 ± 4            | 15 ± 3               | 23 ± 5    | 26 ± 3            | 24 ± 2    |
| Before flood 2018 | 7128              | 7873                 | 12656     | 13809             | 11788     |
| Area (m²)         | 0.31              | 0.46                 | 0.57      | 0.43              | 0.52      |
| After 2nd flood 2019 | 7128           | 7873                 | 12656     | 13809             | 11788     |

4. Discussion

4.1. Immediate flood responses

#### 4.1.1. Physico-chemistry

Transient physico-chemical changes of surface waters during an experimental flood are a function of reservoir characteristics. Patterns of temperature and conductivity change measured during the flood reflect temperature and conductivity of reservoir waters that vary depending on the season. Small reservoirs like Ova Spin are generally characterized by short hydraulic residence time that strongly limit functional processes in the reservoir. For some nutrients, however, even short residence times will result in some sequestration. For example, phosphorus (burial) and to a lesser extent nitrogen (denitrification and burial) will accumulate in reservoir sediments (Maavara et al., 2020; Maranger et al., 2018). During the study floods, deep-water releases suspended and mobilized reservoir sediments. TN and TP concentrations measured during the floods appeared to be tightly coupled with sediment flushing, as observed also in previous studies on the system (e.g., Kevic et al., 2018). Suspended sediment load measured as turbidity was higher in the initial ramping phase, as fine sediments were flushed with reservoir
While turbidity patterns were comparable between the two floods, TP and TN patterns differed greatly between floods, indicating the influence of season and time since a previous flood on nutrients adsorbed to sediments deposited in the reservoir. Transport TP data might suggest the occurrence of deposition along the study segment; here, ~6 and ~3 mg/l, respectively, were deposited in ~2 km during each flood. This large input of phosphorus might influence recovery patterns to experimental floods by subsidizing biofilm assemblages (Odum et al., 1979). Nutrient inputs during experimental floods could have important implications in nuisance algae management (e.g., Lessard et al., 2013; Uehlinger et al., 2003), but require more detailed studies on mechanisms of accrual.

4.1.2. Seston and drift
Temporal patterns of seston during each flood were generally consistent with those observed in previous floods in the upper Spöl (Jakob et al., 2003; Robinson, 2012; Robinson et al., 2004a) and lower Spöl (Kevic et al., 2018), and followed a typical hysteresis curve (i.e., a decline of response intensity over time) (Scheffer et al., 2001). Although floods had identical peak flows and overall released comparable volumes of water (~5 × 10^3 m^3), we observed variability in flow-dependent entrainment and transportation of seston, where differences in the ramping rate and pre-flood conditions (this last factor also supported by benthic samples and field observations, see par. 3) appear to be important in shaping the flood response (Fig. 3). The 2018 flood had a shorter duration and a much steeper rising limb than the 2019 flood. During the 2018 flood, seston response was earlier and stronger and peaked at a lower discharge compared to the 2019 flood, which had stepwise increases in discharge. During the flood of 2019, seston response also could have been influenced by pre-flood conditions determined by the intense spring snowmelt (Fig. 5b and c; see also section 4.2.2 below).

We hypothesize that the lateral contribution of snowmelt water from side valleys and small tributaries, as well as increased discharge from the tributary Ova da Cluozza, likely caused a moderate disturbance in the system (both study reaches), which influenced the distribution of organic matter and macroinvertebrates in the channel before the 2019 flood (discharge of Ova da Cluozza during the study period is reported in Fig. 2). Benthic samples collected before the 2019 flood had extremely low macroinvertebrate abundances in riffle habitats at both study sites (Fig. 5d, Table 2). A closer look revealed that a higher number of macroinvertebrates moved from these areas into lateral habitats (GC and CTR personal observation), which can be important refugia during disturbance events (e.g., Lancaster and Hildrew, 1993a). As drift density and taxa richness were comparable between floods, we suggest that macroinvertebrates were present at similar densities as found before the earlier flood (2018), (Table 1), but likely distributed mostly in refugia habitat. Secondary peaks of drifting macroinvertebrates, seston and turbidity, as observed in other large-magnitude experimental floods in the system (>12 m^3/s peaks) (see Kevic et al., 2018; Robinson, 2012; Robinson et al., 2004b), occurred at both sites during the first flood at approximately peak discharge. These secondary peaks resulted from lateral erosion of scree slopes, bank inundation, and secondary mobilization of the streambed, which likely recruited organisms from hyporheic and side-channel refugia (Lancaster and Hildrew, 1995b; Matthaei et al., 2000; Stubbington, 2012), as well as mobilizing organic matter from banks.

Drift was dominated by Gammaridae (G. fossarum being the only species present in the system). Flood-driven drift composition quickly diverged from background drift in the system - mostly constituted by Chironomidae, Simuliidae and Baetidae at low densities, with the sporadic presence of Gammaridae and other taxa - and converged to resemble benthic composition (Gibbins et al., 2007) with the addition of other limnophilic taxa (e.g., Psychodidae). Taxa drift response varied across floods and locations (Fig. 4). Response to high flows has been shown to be strongly taxon- and habitat-specific, depending on a combination of habitat structure, substrate stability, and physical or behavioral attributes that confer resistance (Holomuzki and Biggs, 2003; Irvine and Henriques, 1984; McMullen and Lytle, 2012; Naman et al., 2017). In our study, Chironomidae and Gammaridae showed a purely passive response, with patterns closely resembling those of organic matter. Baetidae, Nemouridae, and Heptageniidae show mixed patterns of response, where the initial passive response in some cases was followed by a growing number of drifting individuals as flood waters receded. Poff et al. (2018) observed that some traits, such as high mobility (i.e., Baetidae) are important for the persistence of taxa after disturbance, and can be determinant in habitat recolonization and stranding avoidance during water recession. However, this response was variable across floods and sites, suggesting an influence of morphology and water recession velocity (Bond and Downes, 2003; Sedell et al., 1990). The latter response observed might indicate that active drift was initiated by receding water as a mechanism to recolonize instream habitats after disturbance. Changes in water temperature following deep-water releases (so-called thermoequating; Zolezzi et al., 2011), as during 2018 flood, could add up to flow effects and influence taxon-specific drift responses (Schülting et al., 2016).

4.2. Tributary effect

4.2.1. Morphological dynamics and sediment respiration
Although sedimentological differences were not quantitatively assessed in this study (but see Mathers et al., 2021), the environmental discontinuity (sensu Rice et al., 2001) generated by the presence of the tributary appears evident. Above the tributary, morphology is dominated by a single thread channel with riffle-pool sequences, and stable flow conditions associated with a sediment deficit suggest channel degradation and a reduction of lateral dynamics. In this reach, the lack of a major sediment source resulted in negligible morphological changes in response to experimental floods. However, a closer look at instream changes (Fig. 7) suggests that the intense snowmelt before the 2019 flood activated lateral sediment sources also above the confluence with Ova da Cluozza. These sediments were redistributed during the 2019 floods and caused observable changes in habitat structure. A substantial increase in glides, and pools reduction, occurred as a consequence of local gravel depositions, resulting in more homogeneous habitat conditions. However, these changes did not alter the overall geomorphological component of the reach. The floods did not result in enhanced lateral dynamics, suggesting that channel incision had inevitably disconnected the channel from riparian areas. These results highlight the influence of river regulation on morphology, as well as the importance of local sediment sources (i.e., scree slopes and minor tributaries) in sediment-deprived reaches (Vetts, 1979) for instream physical habitat rejuvenation through disturbance.

Further downstream, the sediment input from Ova da Cluozza promotes braiding of the channel across gravel bars and the existence of a wider floodplain with vegetated islands. In unregulated alpine rivers, sediment input by the tributary would be regularly mobilized and redistributed by the receiving channel (Guillén Ludena et al., 2017), often simultaneously, as high flow events would affect the entire catchment. At the Spöl – Ova da Cluozza confluence, sediments accumulate and are partly redistributed solely during experimental floods, as the residual flow on the Spöl does not have enough competence for sediment mobilization. The study floods stimulated morphological processes below the tributary, where patterns of erosion and deposition of sediments, and reorganization of morphological units, were evident. During the study, we witnessed lateral channel movements because of sediment redistribution, the formation of a secondary channel, and the inclusion of bank vegetation in the active channel.
Experimental floods, when coupled with sediment input, can maintain patch dynamics in stream ecosystems (Matthaei et al., 1999a; Townsend, 1989). Stahly et al. (2019) observed the importance of sediment supply in combination with experimental floods in restoring fluvial habitats, however empirical studies in this sense are few. A previous study on this segment of river (Kevic et al., 2018) report that no major morphological changes (measurements based on permanent transects) occurred after the evident channel forming changes that followed the first floods (2000–2003; Mirle et al., 2003). Channel adjustments below a dam are gradual and occur over long timescales (Pets and Gurnell, 2005). In this lower section, lateral mobility could have been enhanced by aggradation with subsequent slope change over the years (Pets, 1984). The formation of secondary lateral channels and wet areas, with trees involved in channel dynamics within the floodplain, has an extraordinary value in increasing microhabitat heterogeneity in the reach, both for fish and macroinvertebrates (Benke and Wallace, 2003; Dolloff and Warren, 2003).

Sediment respiration showed response patterns consistent with the other benthic components (see below). The 2018 flood significantly reduced sediment respiration above the tributary, while leaving it unchanged below, and the 2019 flood resulted in an enhancement of respiration at both sites. In general, a flood’s scouring effect removes surface POM and small particles, thus enhancing the permeability of sediments and promoting POM penetration into deeper sediments (Naegeli et al., 1995). Floods, by modifying local hydraulic characteristics of streambeds, can influence POM presence in the sediment, thus affecting respiration (Naegeli et al., 1995). Studies carried out during early experimental floods on the upper Spöl found that floods greatly enhanced respiration rates, but had no effect on POM sediment content (Uehlinger et al., 2003). With the 2018 flood, we observed significant removal of POM at both sites, but no significant change in 2019. The positive correlation observed between sediment respiration rates and loose POM indicates that, before floods, heterotrophic activity is sustained by this pool of resources. This result suggests that a rapid response to floods was mediated by a combination of scouring of the small fraction of POM and time since the last flood disturbance. However, we also found a negative correlation between attached POM and respiration rate after the flood. Further studies will need to estimate changes in vertical gradients of surface/subsurface exchange after experimental floods to better understand mechanisms that affect responses of sediment metabolism.

4.2.2. Benthic responses

The contrasting benthic responses to the floods, with marked differences above/below the tributary in 2018 and 2019, confirm that short-term disturbance history is a strong determinant of resource distribution and macroinvertebrate community structure (Matthaei et al., 1999a). Below the confluence, periodic disturbances generated by Ova da Cluozza maintain the Spöl river in a dynamic state (i.e., shifting mosaic theory) and influence resistance and resilience of its ecological components. On the other hand, the constant residual flows released from the dam maintain stable environmental conditions above the confluence, with consequent ecological re-arrangements (Bunn and Arthington, 2002). These differences in hydrological conditions determine ecological baseline dissimilarities above and below the tributary, and determine different ecological response patterns to experimental floods, as observed before the 2018 flood. However, these dissimilarities were reversed after the 2019 intense snowmelt event that affected both reaches. The intense snowmelt induced extraordinary, although recurrent at small frequency, hydrological conditions in the system, as testified by the hydrological data from Ova da Cluozza (Fig. 2). The discharge peak generate by this event on Ova da Cluozza was the 4th highest since 2000, with an estimated 10 years return period (FOEN – Federal Office for the Environment flood statistics, available at www.hydrodaten.admin.ch). Based on field observations and the results from this study, we suggest that a major disturbance event was associated with snowmelt also above the confluence.

In 2018, we observed a strong location effect (i.e., tributary effect) driven by differences in disturbance history between the upper and lower reaches. Between the dam and the confluence, the prolonged stable flow conditions with no natural floods resulted in dramatic responses to the experimental flood. Streambed disturbance during the flood resulted in periphyton scouring and mobilization of organic matter (FPOM and CPOM) (Robinson et al., 2018). Macroinvertebrate density decreased 25-fold, suggesting limited system resistance to flood. Periphyton and macroinvertebrate responses were compatible with that observed in previous experimental floods on another residual flow reach on the Spöl (Robinson et al., 2018). Below the confluence, the change in morphology associated with water/sediment input from the tributary and periodic flow disturbances resulted in less marked response patterns to the experimental flood. Sample variability appeared to be higher, reflecting patchiness in a more dynamic system (Matthaei et al., 1999b), and overall changes were limited and generally non-significant for most environmental variables. For instance, periphyton and benthic organic matter were not flushed from the reach, and macroinvertebrates, which had relatively low densities, were in proportion less affected by the flood than in the upper reach. Periphyton persistence in the downstream reach, as opposed to the drastic reduction in biomass in the upstream reach, could also depend on differences in the algal assemblage. As observed by Biggs and Thomsen (1995), algal species show different resistance to scouring events, thus suggesting potential adaptation differences to disturbance in algal communities.

In 2019, ecological response patterns were largely dissimilar to those of 2018. Baseline conditions at both reaches were influenced by the recent natural disturbance, thus the experimental flood had limited effect. In general, benthic components were comparable above/below the tributary before the flood, and responded with similar patterns to the flood, except for CPOM, which is usually patchy in its distribution (Small et al., 2008). Periphyton development, as well as formation and deposition of FPOM, followed a gradual increase over time at both sites, favored by stable flows after the flood events (Biggs, 1988; Webster et al., 1987).

The environmental discontinuity generated at the confluence generally marks shifts in macroinvertebrate assemblages (Katano et al., 2009; Rice et al., 2001), following changes in hydrological conditions and physical habitat structure and distribution. In our system, these baseline differences were evident in 2018. Stable flow conditions before the flood favored Gammaridae, which dominated benthic assemblages (~50%) above the tributary. Gammaridae colonized the Spöl after dam construction and are local indicators of habitat degradation associated with flow regulation (Robinson et al., 2003). Below the tributary, limited availability of suitable habitat and disturbance contributes to maintaining low Gammaridae density. In contrast, suitable habitats for EPT taxa (Ephemeroptera, Plecoptera and Trichoptera) are present and they comprise ~80% of the benthic assemblage, confirming the mitigation function of post-impoundment tributaries (Milner et al., 2019; Ward and Stanford, 1983).

In 2019, the pre-flood disturbance from snowmelt could have triggered migration of invertebrates towards refugia, e.g. hyporheos or side areas of the channel in both reaches, in effect acting as “natural” flood precursor, attenuating the difference between natural, “predictable” high flows and unanticipated experimental floods. The differential persistence of taxa after the flood suggests that river morphology and habitat structure likely influence community attributes of resistance to floods (Gjerlov et al., 2003; Holomuzki and Biggs, 2003; Townsend, 1989). Disturbance history, as observed by Effenberger et al. (2006), has a fundamental importance in determining the distribution of invertebrates in different stream habitats. However, McMullen and Lytle (2012) observed a systematic underestimation in density following disturbance events, thus further studies are needed to quantify the influence of morphology on resistance (and resilience) of stream macroinvertebrate communities across habitats.
After each flood, regardless of location, Simuliidae and Chironomidae recolonized the system at high densities (measured four weeks after), taking advantage of newly available habitats. A strong response of these two taxa was observed after the 2019 flood with cumulative density between 65 and 80%. Early instars of Leuctridae were also found in large quantities below the tributary four weeks after 2018 flood, marking the transition to a winter community. Above the tributary in new system for the survey and classification of river geomorphic units. Geomorphology 283, 143–157. https://doi.org/10.1016/j.geomorph.2017.01.032.
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