

This is the preprint of the contribution published as:

Pasqualini, J., Majdi, N., Weitere, M., Brauns, M. (2025):

The contribution of the hyporheos to whole-stream invertebrate secondary production
Freshw. Sci. **44** (2)

The publisher's version is available at:

<https://doi.org/10.1086/735823>

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The contribution of the hyporheos to whole-stream invertebrate secondary production

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Abstract: The hyporheic compartment is a hot spot of numerous biogeochemical processes. Yet, its role in invertebrate mediated energy fluxes is unclear, due to limited studies assessing benthic and hyporheic production simultaneously. We sampled the meio- and macrofauna from the benthic (from surface to 5 cm depth) and hyporheic (from 5 to 15 cm depth) compartments of an agricultural and forested stream and quantified whole-stream and compartmental production. In the forested stream, 27% of whole-stream invertebrate secondary production occurred in the hydrologically well-connected hyporheic compartment and was sustained by early larval stages of macrofauna, mainly Diptera (28%) and Coleoptera (20%), as well as by the permanent meiofauna like Rotifera (12%), Acari (9%) and Copepoda (9%). In contrast, the agricultural stream had a lower hyporheic contribution (6%) likely because of fine sediment deposition that reduced the water exchange between the water column and the hyporheic compartment, creating anoxic conditions. Despite the large reduction in hyporheic contribution, whole-stream production was approximately threefold higher in the agricultural stream than in the forested one. Only few taxa sustained most of production, with the invasive species *Potamopyrgus antipodarum* (J. E. Gray, 1843) accounting for 42% of whole-stream production. Our findings provide support for the conclusions that (i) the whole-stream production is significantly greater in the agricultural than in the forested stream, this is most likely attributable to the enhanced availability of resources; whereas (ii) the relative importance of the hyporheic compartment decreases at the same time because of unfavorable habitat conditions caused by fine sediment clogging. Therefore, failure to include hyporheic compartment may result in a significant underestimation of whole-stream invertebrate secondary production in streams with permeable sediments. Given that the benthic and hyporheic compartments host different communities and exhibit different production rates both under natural and degraded conditions, each compartment

provides a discrete set of information that cannot be inferred from the other. Consequently, for a comprehensive understanding of the overall functionality of the stream, it is necessary to sample the entire invertebrate length distribution present in both compartments.

Key words: stream functioning – meiofauna – clogging – agriculture – benthic – hyporheic

The hyporheic compartment is a hot spot of biogeochemical processes and serves several ecological functions including temperature regulation, degradation of contaminants, nutrient cycling, and habitat provisioning for invertebrates (e.g., Lewandowski et al. 2019, Boulton et al. 2010, Duff and Triska, 2000, Storey et al. 2004). However, its role for the whole-ecosystem invertebrate mediated energy fluxes is not fully understood because studies quantifying simultaneously benthic and hyporheic secondary production are scarce. Earlier studies suggested a limited contribution of the hyporheic compartment to whole-stream invertebrate production because abundance, biomass, and consequently productivity, of large invertebrates decreases with depth (e.g., Smock et al. 1992, Huryn 1996). Conversely, other studies that have included early larval stages of macrofaunal taxa (e.g., Collier et al. 2004, Reynolds and Benke, 2012, Wright-Stow et al. 2006) and the permanent meiofauna (Majdi et al. 2017, Tod and Schmid-Araya, 2009) showed that hyporheic invertebrates can contribute significantly to production. These apparently contrasting findings might reflect either intrinsic differences in the repartition of production among compartments in streams with different environmental conditions (e.g., pH, grain size distribution, nutrient concentration etc.), or could simply derive from an incomplete sampling of the entire invertebrate size-range in earlier studies.

The distribution of invertebrates between the benthic and hyporheic compartment is influenced by a multitude of environmental factors (Coleman and Hynes 1970, Dole-Olivier and Marmonier 1992, Peralta-Maraver et al. 2018, Dunscombe et al. 2018). Nevertheless, the characteristics of the sediment are amongst the most significant drivers (Williams et al. 1974, Strommer and Smock 1989, Strayer et al. 1997, Weigelhofer and Waringer 2003, Peralta-Maraver et al. 2018). Among the sediment characteristics, permeability and porosity are crucial, as they determine the access of invertebrates to the hyporheic compartment, while also regulating the rates of water exchange between the water column and the sediment (Sawyer and Cardenas 2009). Exchange rates, in turn, determine dissolved O₂ concentrations and the availability of food resources within the hyporheic compartment (Boulton et al. 1998, Hakenkamp and Morin, 2000, Strayer et al. 1997). Consequently, in highly permeable sediment, small-bodied invertebrates can occur at depths of up to 70 cm (Coleman and Hynes 1970, Reynolds and Benke 2012). However, the majority of them are found in the upper 15 cm (e.g., Williams et al. 1974). Such small-bodied invertebrates are expected to significantly contribute to whole-stream invertebrate production (Schmid-Araya et al. 2020) as they have higher biomass turnover rates compared to large-bodied taxa (Brown et al. 2004).

The permeability of the streambed can be reduced as a consequence of physical or biological clogging (Dubuis and De Cesare 2023). Such conditions may arise naturally in sandy or mud-silty streambeds, or may result from human activities such as agricultural land use that increase the deposition of fine sediment on the streambed as a consequence of runoff events (Allan et al. 1997). Most studies on the implications of fine sediment clogging have focused on invertebrate responses in the benthic compartment (e.g., Bo et al. 2007, Jones et al. 2012, Mathers et al. 2017). However, fine sediment deposition can drastically alter the environmental

conditions in the hyporheic compartment, by diminishing water fluxes between the water surface and the hyporheic compartment (Buendia et al. 2013). This creates steep redox gradients and the rapid depletion of O₂, which, in turn, have negative effects on the abundance, biomass and affects assemblages of hyporheic communities (Smock et al. 1992, Boulton et al. 1998, Reynolds and Benke 2012, Mathers et al. 2014). Several studies acknowledge the detrimental effects of fine sediment on the structural and compositional attributes of benthic and hyporheic invertebrates (Jones et al. 2012, Mathers et al. 2014, 2017). However, there is still a lack of empirical understanding of how this affects invertebrate mediated functions, particularly those occurring in the hyporheic compartment, and how this, ultimately, affects whole-stream invertebrate production. In field conditions, this is further complicated by the fact that streams surrounded by agricultural land are not only exposed to fine sediment deposition, but often also have altered levels of light and nutrients (Tank et al. 2021). These conditions have been demonstrated to increase benthic macroinvertebrate secondary production compared to more pristine systems, by stimulating primary production and increasing food quality (Shieh et al. 2002, Wild et al. 2022). However, it is unclear whether the observed increase in production will be as pronounced if the hyporheic compartment is also included in whole-stream estimates. An increase in fine sediment deposition is likely to degrade hyporheic habitat conditions, potentially impairing hyporheic communities, and, consequently, invertebrate mediated functions. If the hyporheic compartment is a significant contributor to whole-stream production, this would imply that when comparing estimates of whole-stream production between agricultural and forested streams, the observed differences may be less pronounced.

The aim of this study was to investigate how the relative contribution of the benthic and hyporheic compartments varies in a forested and agricultural stream, and how this, in turn,

affects whole-stream invertebrate production. Therefore, we sampled the macro- and meiofauna inhabiting the benthic and hyporheic compartments in a forested stream, with a highly-permeable sediment, and in an agricultural stream, with a reduced permeability due to fine sediment deposition. Specifically, our research aims were to 1) to compare environmental controls on benthic and hyporheic invertebrate communities in an agricultural vs. a forested stream, 2) to quantify the relative contribution of the benthic and hyporheic compartments to whole-stream production in both cases. We expect that the less permeable conditions in the hyporheic compartment of the agricultural stream should reduce the proportional contribution of the hyporheic compartment to whole-stream invertebrate secondary production.

METHODS

We conducted a field study from August 2019 to June 2020 in 2 headwater streams located in the Harz region in Germany. At each stream site, we concurrently sampled the benthic and hyporheic invertebrates and measured environmental characteristics of water column, benthic, and hyporheic compartments bimonthly over 1 y. We then calculated invertebrate secondary production and compared the relative contribution of the benthic and hyporheic compartments to whole-stream production in these streams.

Site selection

The forested stream (Drängetalbach, 51°48'21.02"N, 10°43'51.82"E) is surrounded by coniferous forest, has a natural hydromorphology with pool-riffle sequences, gravel bars, and woody debris (Jähkel et al. 2022) (Fig. S1-A, Appendix S1). Water column nutrient concentrations are relatively low (Table 1) and its streambed consisted of cobbles (range: 63-200

mm) and coarse gravel (range: 20-63 mm). In contrast, the agricultural stream (Asse, 51°55' 23.357" N, 11°1'53.443" E) is channelized, surrounded by herbaceous vegetation and alder on the river bank (Fig. S1-B, Appendix S1). The stream exhibits higher nutrient concentrations and conductivity in the water column (Table 1, Fig. 1), and the streambed is predominantly fine sand (range: 0.2-0.063 mm) and coarse silt (range: 0.063-0.02 mm). Water temperature is slightly higher in the agricultural stream than in the forested one (Table 1). In addition, the agricultural stream has a higher light exposure (Table 1), as the stream flows through open fields and has a less dense canopy cover compared to the forested stream which flows through a steep valley and exhibit a denser canopy cover. Conductivity, temperature, and pH were measured in the water column with a multimeter (Multi 3630 IDS SET F, Xylem Analytics GmbH, Weilheim, Germany). Light intensity was measured over 1 y by 5 light intensity data loggers (MX2202 Onset, Bourne, Massachusetts, USA) placed along the stream reaches.

Table 1 Environmental characteristics of the 2 studied streams. Values are annual means \pm 1 SD of bimonthly measurements. Except discharge values which are annual means, minimum and maximum.

Variable	Forested	Agricultural
Land use (Forest % – Arable % – Other %)	100 – 0 – 0 ^a	12 – 85 – 3 ^a
DN (mg L ⁻¹)	2.2 \pm 0.6	10.4 \pm 2.0
N-NO ₃ ⁻ (mg L ⁻¹)	1.9 \pm 0.6	9.3 \pm 2.3
N-NH ₄ ⁺ (mg L ⁻¹)	0.01 \pm 0.01	0.06 \pm 0.05
SRP (μ g L ⁻¹)	7 \pm 3.2	31 \pm 15
DOC (mg L ⁻¹)	2.7 \pm 0.7	2.6 \pm 0.5
Dissolved O ₂ (mg/L)	11.5 \pm 1.5	10.0 \pm 2.6
Chl <i>a</i> (μ g/L)	< 1.1	4.0 \pm 3.2
Discharge (L/s)	42 (5-196) ^a	23 (4-59) ^a
Water temperature (°C)	7.6 \pm 3.2	9.3 \pm 6.2
pH	7.9 \pm 0.4	8.2 \pm 0.1
Conductivity (μ S/cm)	314 \pm 37	1402 \pm 155
Light (PAR)	40 \pm 61 ^b	97 \pm 148 ^b

^a Jähkel et al. 2022, ^b Jähkel A., Helmholtz Centre for Environmental Research, Magdeburg, Germany, personal communication.

1 **Environmental characteristics**

2 We sampled the streams every 2 months over 1 y. On each sampling occasion ($n = 6$), we
3 measured environmental variables related to water quality (i.e., nutrients and O₂) and sediment
4 characteristics, as important drivers of invertebrate assemblages and productivity. We aimed to
5 assess whether differences in nutrients, O₂ and sediment characteristics occurred between
6 compartments (i.e., water column, benthic and hyporheic) within each stream.

7

8 **Nutrients** We followed the same sampling design in both streams. On each sampling occasion
9 we collected 5 water samples from the benthic and hyporheic compartments and 1 from the
10 water column. The need to collect a higher number of replicate samples from the sediment than
11 from the water column arose from the higher variability in nutrient concentration within the
12 former than in the latter (Hartwig 2016). A parallel study indicated that nutrient concentrations in
13 the water column of both streams remained constant during each sampling date (Jähkel A.,
14 Helmholtz Centre for Environmental Research, Magdeburg, Germany, personal communication,
15 Coefficient of Variation of nutrient concentration are in Table S1, Appendix S1).

16 In both streams, we collected 1-L of surface water and filtered it through 0.22- μ m filters
17 (Sartorius, Minisart Syringe Polycarbonate Filters). Samples were then transported at 4°C to the
18 laboratory and DN, N-NO₃⁻, N-NH₄⁺, SRP, DOC concentration measured following standard
19 protocols (DIN EN 1484, DIN EN ISO 11732, DIN EN ISO 13395, DIN EN ISO 15681-2). An
20 additional 500 mL water sample was collected for chl *a* determination. The water was stored at 4°C,
21 transported in the dark, and filtrated onto a glass microfiber filter (GFF: 47 mm diameter, 0.7
22 μ m pore size, Whatman, England) in the laboratory on the same day. Chl *a* concentration was
23 determined photometrically (DIN 38412-16).

24 To collect the pore water from the sediment of the agricultural stream, we extracted 5
25 sediment cores using a polyvinyl chloride corer (PVC) (UWITEC, 019011, Mondsee, Austria,
26 inner diameter = 9 cm) pushed to a depth of 30 cm. After sealing the top with a rubber stopper,
27 the core was extracted, and a second stopper placed underneath to prevent sediment loss. Cores
28 were then cut into 0-5 cm and 5-15 cm sections, the sediment transferred to centrifuge tubes, and
29 transported to the lab at 4°C. On the same day, the tubes were centrifugated (Allegra X-15R
30 Centrifuge, Beckman Coulter, Krefeld, Germany) at 1500 rpm for 10 min to separate the pore
31 water from the sediment. Extracted pore water was filtered and analyzed as previously described.
32 Some samples lacked sufficient pore water volume to measure all parameters (n DN = 56/60, n
33 N-NO₃⁻ = 56/60, n SRP = 60/60, n DOC = 58/60, n N-NH₄⁺ = 60/60).

34 For the forested stream, we used a different technique to extract pore water due to the
35 cobble-gravel lithology. We planned to install 2 PVC tubes (1 cm inner diameter, screened at the
36 bottom over 5 cm) at 5 locations a week before sampling. One tube was to reach 5 cm depth, the
37 other 15 cm. However, due to the coarse lithology, we could not install a tube firmly at 5 cm
38 depth, thus we only installed the tubes at a 15 cm depth. A week after the installation, pore water
39 was retrieved using a peristaltic pump (Peristaltic pump 12 VDC, Eijkelkamp, Netherlands) at a
40 constant rate of 1.8 mL/s, after discarding the first 20 mL. Collected pore water was filtered and
41 stored as previously described.

42 To compare the nutrient concentration across compartments (i.e., water column, benthic
43 and hyporheic), we averaged the measurements obtained from the 5 samples collected from each
44 compartment during each sampling campaign, resulting in a final sample size of $n = 6$. Then, we
45 used a linear mixed-effects model (function *lme*, package *lme4*, (Bates et al. 2015)) to examine
46 the relationship between the measured nutrient concentrations and the compartments (i.e., water

47 column, benthic, hyporheic). We set “*campaign*” as a random factor to account for lack of
48 independency of water samples collected during the same sampling campaign. The model was
49 formulated as: *nutrient concentration* ~ *compartment*, random = ~1 | *campaign*. When statistical
50 significance was determined, we additionally ran a Tukey’s Honest Significant Difference
51 (Tukey’s HSD) post hoc test to detect specific differences among the compartments. The normal
52 distribution of residuals and homogeneity of variances of the data was visually inspected by
53 plotting residuals vs fitted values.

54

55 ***Oxygen*** In both streams, we measured O₂ concentrations in the water column and sediment
56 pore water. O₂ in the water column was measured with the Winkler method (DIN EN 25813) by
57 collecting an additional 250 mL water sample. To assess O₂ in the pore water we employed 2
58 distinct approaches due to varying streambed lithologies. In the forested stream, we inserted a
59 PVC tube (inner diameter = 1 cm) at 15 and 30 cm depth. A week after the installation, we
60 collected pore water as previously described and measured O₂ using the Winkler method. In the
61 agricultural stream, we retrieved an additional sediment core and determined the depth at which
62 the sediment got anoxic (i.e., O₂ < 0.5 mg/L) by using a pre-calibrated Firesting needle (Oxygen
63 Micro/Minisensor, sensor code: ZA7-521-197, Pyroscience, Aachen, Germany) attached to a
64 house-made micromanipulator. The micromanipulator served to gradually insert the needle into
65 the sediment. The O₂ concentration and the depth reached by the needle in the sediment were
66 manually recorded. This procedure was repeated at 3 distinct spots within the same core during
67 each sampling event.

68

69 ***Sediment*** Our aim was to characterize sediment grain size distribution in both streams.

70 The sediment collection is described in detail in the next paragraph as it is associated with the
71 invertebrate sampling. In the forested stream, sediment was sieved using a nested column of
72 sieves (details in Pasqualini et al. 2023), while the sediment of the agricultural stream was
73 analyzed with a particle analyzer by laser diffraction (CILAS 1190d, CPS USA) due to the very
74 fine grain size of the sediment. Then, the 90, 50, and 10 percentiles (d90, d50, d10) of the grain
75 size accumulation curves of the sediment were calculated. A paired *t*-test was run to test for
76 differences in d90, d50 and d10 and fine sediment content (i.e., percentage < 2 mm) among
77 compartments in each sediment sample. The *t*-test was paired to account for the non-
78 independency of benthic and hyporheic samples collected from the same core.

79 To ensure uniform hydrological conditions in the hyporheic zone of the forested stream, we
80 additionally installed to a depth of 15 and 30 cm 5 high-density polyethylene (HDPE, 4 cm outer
81 diameter) piezometers screened at the bottom over a 5 cm range, in an area located 1-1.5 m from
82 the area where the invertebrates were planned to be collected. We measured vertical head
83 gradients (VHG) between surface water and piezometer water. We recorded only downwelling
84 conditions. This procedure could not be conducted in the agricultural stream due to the
85 displacement of non-cohesive fine sediment when hammering the piezometer into the sediment.

86

87 **Invertebrates**

88 At each stream site, we sampled the benthic (0-5 cm depth) and hyporheic (5-15 cm
89 depth) invertebrate communities every 2 months along a 300-meter reach. On each sampling
90 occasion ($n = 6$) at each stream, we collected 5 samples from 5 sites using a Surber sampler and
91 5 samples using a sediment corer. The sediment corer served two purposes: to collect smaller
92 benthic invertebrates that were not retained by the Surber net; and to sample invertebrates in the

93 hyporheic compartment. After extracting invertebrates from the sediment in the laboratory,
94 secondary production was calculated for both the stream and compartments using Plante and
95 Downing regression (1989).

96

97 *Sample collection*

98 In both streams, the Surber sampler (0.0625 m², 250- μ m mesh) was placed on the
99 sediment and the sediment was vigorously stirred to 5 cm depth. The collected material was
100 rinsed on a white plastic tray to separate organic and inorganic fractions and the organic fraction
101 preserved in 70% ethanol. A corresponding sample was collected with the sediment corer 3-7 d
102 later at a distance between 0.5-1.5 m from the Surber sampling area, ensuring similar
103 microhabitats (i.e., water depth, flow velocity, sediment characteristics, shading) to those
104 sampled with the Surber sampler.

105 In the agricultural stream, we used a PVC corer (UWITEC, 019011, Mondsee, Austria,
106 inner diameter = 90 mm) manually inserted in the sediment to a depth of 30 cm. The cores were
107 extracted and cut into 2 parts (0-5 cm depth and 5-15 cm depth), stored in plastic bags, and
108 transported frozen at -20°C to the laboratory. In the forested stream, we used a freeze-corer
109 (UWITEC, freeze-corer type 1, Mondsee, Austria), which is the only quantitative method for
110 collecting hyporheic invertebrates in cobble-gravel streambeds (Bretschko 1985). Freeze-corer
111 tubes were installed to a depth of 45 cm, after the Surber samples were collected. After 3-7 d, the
112 cores were extracted with a tripod, and the cores were cut into 2 segments (0-5 cm depth and 5-
113 15 cm depth). The samples were transported at -20°C to the laboratory for subsequent
114 processing. The details of the extraction with the freeze-corer are presented in Pasqualini et al.
115 2023.

116

117 **Sample processing** Invertebrates collected with the sediment corers were extracted using the
118 flotation method described in the protocol by Traunspurger and Majdi 2017. After thawing, the
119 sediment was mixed, weighed and a weighted subsample was sieved on a nested column of
120 stainless-steel sieves (2-mm, 1.12-mm, and 20- μ m). The fraction retained on the 20- μ m sieve
121 was collected and centrifuged at 1500 g for 5 min (Allegra X-15R Centrifuge, Beck-man
122 Coulter, Krefeld, Germany) to remove excess pore water which was poured onto a 20- μ m sieve.
123 Ludox HS-40 (Sigma Aldrich, 420816), diluted to 1.14 g/mL, was added to the sediment. Mixing
124 of the sediment and diluted Ludox solution was done by placing the samples in a mechanical
125 vertical rotor (Reax 2, Heidolph Instruments GmbH and CO. KG, Schwabach, Germany) for 10
126 min at 20 rpm, and then centrifuging them at 800 g for 5 min to separate the organic from the
127 inorganic fraction. Extracted invertebrates and the Ludox were poured onto the 20- μ m sieve, and
128 the extracted were preserved in a 4% formaldehyde solution with a few drops of Rose Bengal
129 dye. This method enabled the extraction of both temporary and permanent meiofauna.
130 Invertebrates collected with the Surber sampler and the corer were counted and identified under a
131 stereomicroscope (Leica S8AP0, Wetzlar, Germany) with a 10-80x magnification. Identification
132 of macrofauna was done to the lowest possible level, while permanent meiofauna was identified
133 to the major group level (i.e., Rotifera, Nematoda, Copepoda, Ostracoda, Cladocera, Tardigrada,
134 Acari). Subsampling was performed for Rotifera and Nematoda when more than 200 individuals
135 were counted in the same sample. We measured the body length (BL) or head width (HW) of 30
136 randomly selected individuals from each taxon and sample. During the processing phase, we lost
137 1 benthic sample collected in August in the agricultural stream, and 1 hyporheic sample collected
138 in October in the forested stream.

139

140 **Data preparation** We created a benthic sample by combining the benthic sample obtained
141 from the Surber sampler with the upper 5 cm portion of the corresponding sediment sample
142 collected with the corer. This step was crucial to include small-size invertebrates not retained by
143 the Surber net. However, to avoid overestimation of production due to counting individuals
144 twice, we corrected the abundance values of those benthic taxa which were collected
145 simultaneously by both techniques. For this, we analyzed the length size distributions of taxa
146 collected with both techniques and implemented a selection criterion (detailed in Pasqualini et al.
147 2023). Briefly, if the length distributions did not overlap, we considered the techniques
148 complementary and summed abundances. However, if there was overlap, we recalculated the
149 numerical abundances of individuals within the overlapping range. Our approach assumed that
150 coring techniques provided a better estimate of small-size invertebrate abundance, while the
151 Surber sampler was more accurate for larger ones.

152

153 **Community composition analysis** We aimed to assess differences in the composition of
154 invertebrate communities among the compartments of the two streams. To visualize differences,
155 we used non-metric multidimensional scaling NMDS (function *metaMDS*, package *vegan*
156 version 2.6-4 (Oksanen et al. 2020)). This was supplemented by permutational multivariate
157 ANOVAs PERMANOVA (Anderson 2006) (function *adonis2*, package *vegan* version 2.6-4
158 (Oksanen et al. 2020)) on Bray-Curtis similarity matrixes generated from Hellinger-transformed
159 abundance data for macrofauna, and square-root transformed data for permanent meiofauna. To
160 address the lack of independence between benthic and hyporheic samples retrieved from the

161 same sediment core, we incorporated the blocking factor "strata = id" into the formula. The
162 average contribution of each taxon to differences among compartments was evaluated with the
163 *SIMPER* function (package *vegan* version 2.6-4 (Oksanen et al. 2020)). Data was checked to
164 meet the assumption of homogeneity of multivariate dispersions with the function *betadisper*
165 (package *vegan* version 2.6-4 (Oksanen et al. 2020)). The analyses were performed separately for
166 the macrofauna and the permanent meiofauna fraction due to the different taxonomical
167 resolution. All tests and analysis were performed in R (R version 4.3.2, R Project for Statistical
168 Computing, Vienna, Austria).

169

170 **Secondary production** Macrofauna and permanent meiofauna secondary production were
171 calculated separately for both compartments using Plante and Downing (1989) regression (P&D)
172 (Eq. 1). While P&D regression is commonly used for permanent meiofauna (e.g., Majdi et al.
173 2017), it is less commonly applied for macrofauna. Usually, macrofauna estimates are obtained
174 using the size frequency (SF) method following Hynes and Coleman (1968) and Hamilton
175 (1969), with correction for cohort production intervals (Benke 1979). However, SF tend to
176 overestimate production compared to P&D (Butkas et al. 2011). Given the prevalence of
177 macrofauna in the benthic compartment and permanent meiofauna in the hyporheic
178 compartment, the use of different methods could increase the differences between compartments.
179 To prevent such artifacts, we applied the same method to both invertebrate fractions. Taxa with a
180 mean annual abundance $> 50 \text{ ind./m}^2$ were included in the calculation. For those, dry mass (DM)
181 was calculated using published length–mass (Table S1, Appendix S2). Taxon-specific production
182 P_i ($\text{g DM m}^{-2} \text{ y}^{-1}$), was calculated based on mean annual taxon-specific biomass values B_i (g

183 DM/m²), maximum mass per individual M_{max} (mg DM/ind.) and annual average temperature T
184 (°C) measured in the water column:

$$185 \quad \text{Log}_{10}(P_i) = 0.06 + 0.79\text{Log}_{10}(Bi) - 0.16\text{Log}_{10}(M_{max}) + 0.05T. \quad (\text{Eq. 1})$$

186 To account for spatial variability in abundance and biomass in our production estimates, we
187 considered the 5 samples collected during each sampling campaign at each depth as replicates.
188 We permuted these replicate samples using a randomized block design, generating all possible
189 combinations across the 6 sampling campaigns (5^6 unique combinations, details in Tables S2-3,
190 Appendix S1). For each combination, we estimated mean, median, SD, 95% CI of taxon-specific
191 production. Compartmental and whole-stream production for each combination was obtained by
192 summing mean taxon-specific production and propagating the error with the *propagate* function
193 (package *propagate* version 1.0-6). The relative contribution of each taxon to compartmental
194 production was calculated by summing the medians and determining the percentage contribution
195 of each taxon. To assess differences in whole-stream and compartmental secondary production
196 we compared 95% CI. Medians with nonoverlapping CI were considered significantly different
197 (Cross et al. 2013, Brabender et al. 2016, Wild et al. 2022).

198

199 **RESULTS**

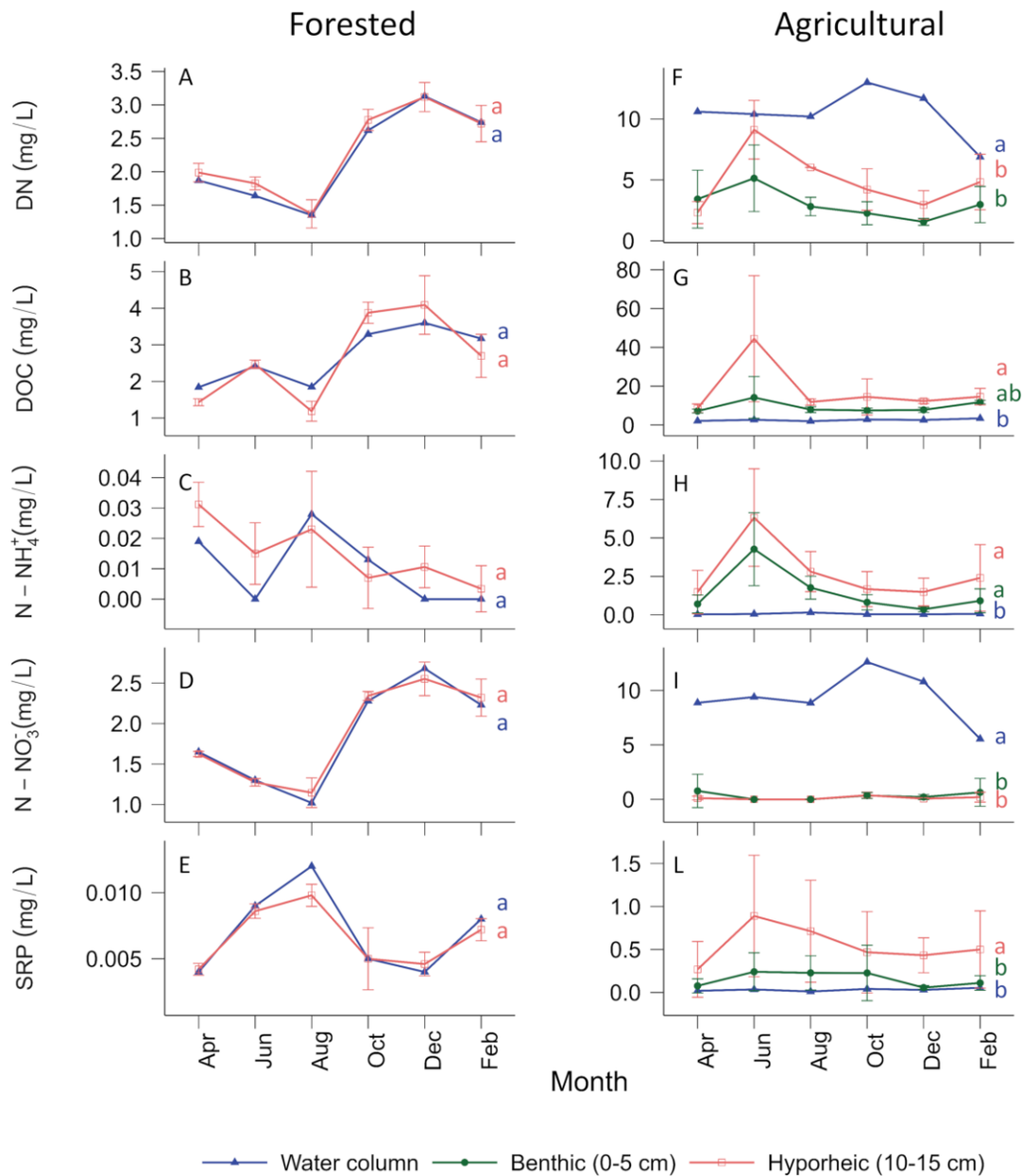
200 **Environmental conditions**

201 Nutrient concentrations did not differ among the water column, benthic and hyporheic
202 compartment in the forested stream (Fig. 1A-E, Table S4 Appendix S1). Conversely strong
203 differences occurred in the agricultural stream (Fig. 1F-L). DOC, N-NH₄⁺, and SRP concentrations
204 were substantially higher in the hyporheic compartment than in the water column (Tukey's HSD
205 post hoc tests, Table S5 Appendix S1; Fig. 1G-H-L). Conversely, DN, N-NO₃⁻ concentrations

206 were substantially lower in the hyporheic compartment than in the water column (Tukey's HSD
207 post hoc tests, Table S5 Appendix S1; Fig. 1F-I). In the agricultural stream, anoxic conditions (<
208 0.05 mg O₂/L) were recorded in the upper 5 cm of sediment on every sampling occasion (Table
209 S6, Appendix S1). In contrast, the sediment of the forested stream was oxygenated at each
210 sampling occasion at least up to a depth of 30 cm (Fig. S2, Appendix S1). Sediment d₉₀, d₅₀, d₁₀,
211 and fine sediment content did not differ among compartments in both streams (paired *t*-test, *p* >
212 0.05; data not shown).

213

214 *Figure 0-1 Nutrient concentrations were measured bimonthly at various depths in the forested (A-E) and*
 215 *agricultural (F-L) streams. Different lowercase letters indicate significantly different values at the $p <$*
 216 *0.05 level (Tukey's HSD post hoc test) within the same*



217

218 **Invertebrates**

219

220 **Community composition** NMDS ordination of macrofauna communities revealed marked

221 differences in assemblage composition between the benthic and hyporheic compartments in both

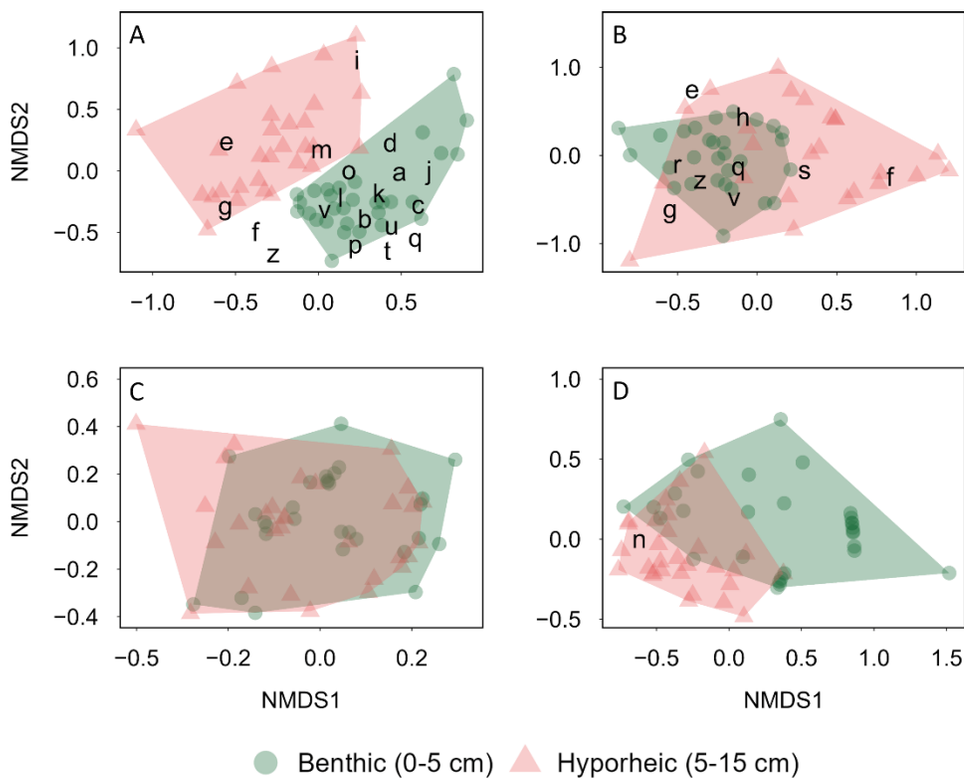
222 streams (Fig. 2A-B). In the forested stream, differences between benthic and hyporheic
223 communities were significant (Fig. 2A, PERMANOVA, $F = 13.36$, $R^2 = 0.19$, $p < 0.001$).
224 Ephemeroptera, Plecoptera, and Trichoptera were numerically more abundant in the benthic
225 compartment than in the hyporheic compartment (Fig. 2A, Table 2). In contrast, Coleoptera and
226 Diptera were more abundant in the hyporheic compartment and contributed to 35% to
227 differences among compartments (SIMPER) (Fig. 2A, Table 2). In the benthic compartment of
228 the agricultural stream, Diptera (8 taxa) and *Potamopyrgus antipodarum* [J. E. Gray, 1843]
229 (Gastropoda) dominated the assemblage. Differences between benthic and hyporheic
230 communities were significant (Fig. 2B, PERMANOVA, $F = 8.06$, $R^2 = 0.13$, $p < 0.001$) and
231 reflected a lower abundance of Diptera larvae in the hyporheic compartment (SIMPER, 56% of
232 variation attributable to Diptera larvae) (Fig. 2B, Table 3).

233 The composition of the benthic and hyporheic permanent meiofauna did not significantly
234 differ in the forested stream (PERMANOVA, $F = 8.06$, $R^2 = 0.008$, $p = 0.7$; Fig. 2C), but it did
235 in the agricultural stream (PERMANOVA, $F = 4.27$, $R^2 = 0.007$, $p = 0.01$; Fig. 2D). In the
236 agricultural stream, permanent meiofauna densities were considerably lower in the hyporheic
237 compartment than in the benthic one, differences were especially large for Nematoda (Fig. 2D,
238 Table 3).

239

240 Figure 2 NMDS ordination of macrofauna communities in the forested stream (A) and in the agricultural
 241 stream (B). NMDS of permanent meiofauna communities in the forested stream (C) and in the agricultural
 242 stream (D). Letters indicate the taxa identified with SIMPER that contributed the most to differences
 243 among zones. a *Agapetus fuscipes*, b *Amphinemura* spp., c *Ancylus fluviatilis*, d *Baetis* spp., e
 244 *Ceratopogonidae*, f *Chironomidae*, g *Chironomini* spp., h *Elmis* spp., i *Esolus* spp., j *Glossosoma* spp., k
 245 *Ibisia marginata*, l *Leuctra* spp., m *Limnius* spp., n *Nematoda*, o *Nemoura* spp., p *Oligochaeta*, q
 246 *Orthoclaadiinae*, r *Potamopyrgus antipodarum*, s *Prodiamesa olivacea*, t *Sericostoma* sp., u *Simuliidae*, v
 247 *Tanytarsini* spp., z *Tanypodinae*. NMDS = nonmetric multidimensional scaling.

248



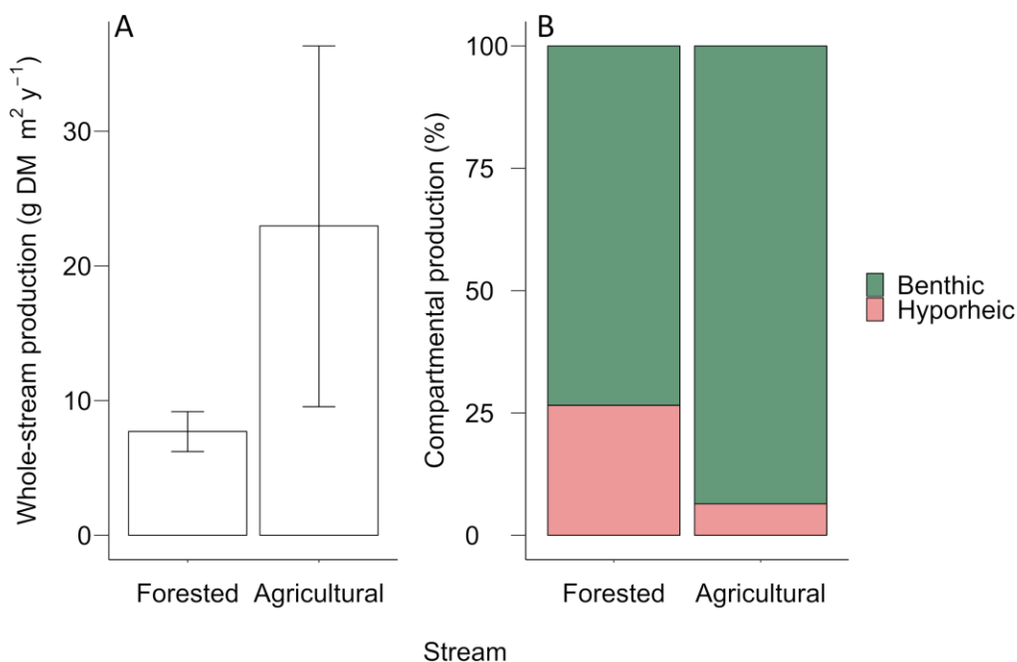
250

251 **Secondary production** In the forested stream median and 95% CI annual whole-stream
 252 invertebrate production was 7.71 (6.21-9.18) g DM m⁻² y⁻¹ (Fig. 3A). 73% of production
 253 occurred in the benthic compartment and 27% in the hyporheic one (Fig. 3B). No taxa
 254 contributed to more than 12% of compartmental production (Fig. 4). In the benthic compartment,
 255 the taxa with the highest contribution to production were *Ancylus fluviatilis* (Gasteropoda:

256 Planorbidae) (12%), *Sericostoma* sp. (Trichoptera: Sericostomatidae) (10%), Chironomidae
 257 (Diptera) (7%) (Table 2). Most of production in the hyporheic compartment was sustained by
 258 early larval stages of *Esolus* sp. (Elmidae) (14%) and Chironomidae (12%), but also by
 259 representatives of the permanent meiofauna such as Rotifera (12%), Acari (9%), and Copepoda
 260 (9%).

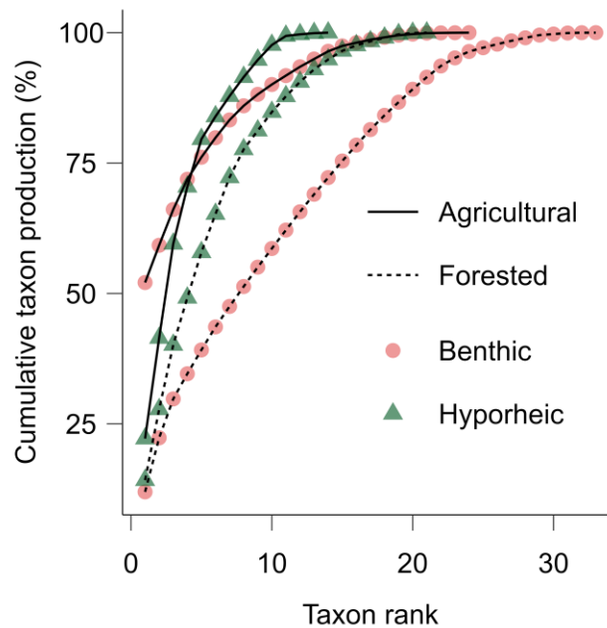
261 In the agricultural stream median and 95% CI annual invertebrate production was 22.97
 262 (9.54-36.33) g DM m⁻² y⁻¹ (Fig. 3A). 94% of production occurred in the benthic compartment
 263 and 6% in the hyporheic one (Fig. 3B). Benthic production was sustained by few taxa and *P.*
 264 *antipodarum* accounted for 52% of benthic production and 42% of whole-stream production
 265 (Fig. 4, Table 3), followed by *Pisidium* sp. (Bivalvia: Sphaeriidae) (7%) and Naididae (Clitellata)
 266 (7%) (Table 3). In the hyporheic compartment, Diptera (8 taxa) and Nematoda contributed
 267 respectively to 74% and 14% of hyporheic production.

268 *Figure 3 Median and 95 % CI of annual whole-stream invertebrate production (A). Mean relative*
 269 *766 contribution of the compartments to annual whole-stream invertebrate production (B).*
 270 *767 DM = dry mass.*



271

272 *Figure 4. Cumulative percentage of taxa's contribution to whole-stream production in the benthic*



273

274 **DISCUSSION**

275 The hyporheic compartment has been recognized as a hot spot for many ecosystem
276 functions, but its role in invertebrate mediated energy fluxes has not been fully understood. In
277 this study, we sampled the macro- and meiofauna inhabiting the benthic and hyporheic
278 compartments of a forested and an agricultural stream, and quantified compartmental and whole-
279 stream invertebrate secondary production.

280 The findings of this study indicate that the contribution of the hyporheic compartment to
281 whole-stream production is significantly reduced in the agricultural stream, with clogged
282 sediment, in comparison to the forested one, with more permeable sediment. Nevertheless,
283 whole-stream production in the agricultural stream was approximately threefold higher than in
284 the forested stream, with most of this difference reflecting increased production in the benthic
285 compartment. Many factors regulate levels of invertebrate productivity streams, and it is
286 challenging to disentangle the influence of individual components under field conditions.
287 Nevertheless, by comparing our findings with those of previous studies conducted in low-order

288 streams, we identify and discuss two main processes that may help to explain the observed
289 patterns.

290

291 **Resource availability is a major control of whole-stream invertebrate production**

292 Several studies showed that nutrient enrichment and higher food quality considerably
293 increase benthic macroinvertebrate production in agricultural (Shieh et al. 2002 and Wild et al.
294 2022), urban (Gücker et al. 2011), and detritus-based ecosystems (Cross et al. 2006) compared to
295 reference conditions. However, it is unclear whether this increase is representative of the whole-
296 stream, as these assessments are based on benthic macrofauna and do not include neither the
297 hyporheic compartment, nor the full range of invertebrate sizes.

298 Our study is among the few that have included both compartments and the entire length
299 distribution in estimating whole-stream production. Consequently, to contextualize our findings,
300 we compare our whole-stream and compartmental production estimates with those obtained in
301 studies with similar sampling strategies. Specifically, studies conducted in temperate low-order
302 streams with different environmental characteristics but using comparable sampling strategies
303 (i.e., included the whole invertebrate length distribution and the hyporheic compartment to
304 various depths (Table 4)). Our whole-stream production estimates for the forested stream are
305 comparable to those found in the Ems, Furlbach and Lone Oak, streams which have different
306 substrates but relatively low nutrient concentrations (Table 4). While the whole-stream
307 invertebrate production estimate for the sandy agricultural stream is comparable to values
308 estimated for the gravel habitat of the Lambourn, a hypernutrified chalk stream sampled to a
309 depth between 5 and 10 cm (Table 4).

310 Temperature can influence secondary production by stimulating invertebrate growth rates
311 (Brown et al. 2004). However, differences in water temperature between these streams do not
312 appear to be pronounced. Furthermore, similar production values are observed in streams with
313 different streambeds, thus, large differences in whole-stream estimates may be attributable to
314 differences in nutrient availability. This is consistent with previous conclusions based on benthic
315 macroinvertebrates production alone (Shieh et al. 2002, Cross et al. 2006, Wild et al. 2022).
316 Negishi et al. 2019 showed that benthic and hyporheic invertebrates assemblages responded
317 differently to changes in water quality in a stream with a gravel bed. In this study, the co-
318 occurrence of other stressors related to agricultural land use (e.g. fine sediment) prevents us from
319 concluding that enhanced nutrients alone will stimulate differently benthic and hyporheic
320 production. However, given the presence of different communities with different growth rates in
321 the compartments (i.e. macro- vs. meiofauna), we expect that similarly enhanced nutrient
322 availability will differentially affect invertebrate production, with further consequences for
323 whole-stream production.

324 In the agricultural stream, 52% of benthic production was supported by the invasive
325 species *P. antipodarum*. The success of *P. antipodarum* is attributable to a number of factors,
326 including its ability to colonise fine substrates, feed on filamentous algae, tolerate a wide range
327 of abiotic conditions, high fecundity, and its capacity to reproduce parthenogenetically (see
328 review in Alonso and Castro-Díez 2008). Similar patterns of extremely high production of *P.*
329 *antipodarum* have been observed in other streams with degraded environmental conditions (Hall
330 et al. 2003, Alonso and Castro-Díez 2008), thus our result aligns with previous conclusion that
331 agricultural practices may enhance stream production through the dominance of few generalist
332 species (Wild et al. 2022). The overdominance of a single species can destabilize fundamental

333 ecosystem processes such as secondary production, rendering these processes unstable and
334 susceptible to collapse (Dolbeth et al. 2003, Hall et al. 2006, Wild et al. 2022). Our study, by
335 including the hyporheic compartment, not only aligns with these conclusions, but also shows that
336 the dominance effect is even more pronounced due to the simultaneous reduction in hyporheic
337 functionality. This suggests that the susceptibility of these processes to collapse in the case of
338 strong environmental perturbations may be even higher than expected.

339

340 **Sediment characteristics are a major control of redistribution of invertebrate production**
341 **between stream compartments**

342 While resource availability appears to be the main driver of differences in whole-stream
343 productivity, sediment characteristics have a significant influence on the establishment of the
344 community in the hyporheic compartment, thereby affecting the relative contribution of the
345 hyporheic compartment to whole-stream production. This pattern was clearly visible in our
346 study. In the hyporheic compartment of the agricultural stream, the deposition of fine sediment
347 on the streambed resulted in the sediment being disconnected from the water column, resulting in
348 anoxic conditions (< 0.05 mg O₂/L) and elevated levels of N-NH₄⁺ in the hyporheic
349 compartment. These conditions impinge invertebrates over extended periods (Zhang et al. 2023),
350 and it is likely that they have prevented the establishment of a resident hyporheic community.
351 Indeed, in the agricultural stream, we consistently found lower abundance, biomass, and
352 production of almost all taxa in the hyporheic compartment in comparison to the benthic
353 compartment. We observed that the hyporheic community was mostly reduced to Diptera larvae
354 and nematodes, altogether making up only 6% of the overall stream production. Dipterans have
355 developed respiratory adaptations (e.g., synthesis of high-affinity respiratory pigments as

356 haemoglobin) to cope with hypoxic or anoxic conditions, while nematodes are known to thrive
357 under suboxic conditions (Taheri et al., 2014).

358 Given that no prior studies employed our sampling approach in streams with
359 environmental conditions similar to those present in the agricultural stream, we are unable to
360 compare hyporheic productivity estimates from the agricultural stream with previous research.
361 Although we did not replicate our measurements in streams with similar conditions, we
362 anticipate that the functionality of the hyporheic compartment will be similarly diminished in
363 streams affected by biological or physical clogging because of the detrimental effects of fine
364 sediment on the hyporheic compartment. This indicates that alterations in sediment permeability
365 resulting from agricultural land use not only affect macroinvertebrate structural metrics (Bo et al.
366 2007, Jones et al. 2012, Mathers et al. 2017), but also substantially diminish the functionality of
367 the ones inhabiting the hyporheic compartment. This necessitates the implementation of
368 measures to preserve of the hyporheic compartment's role as a habitat, as it sustains vital
369 ecosystem functions.

370 Unlike in the agricultural stream, the hyporheic compartment of the forested stream had
371 similar water quality and O₂ concentration to the water column, indicating that the hyporheic
372 compartment was well-connected with the surface water. In these conditions, we found a diverse
373 and productive hyporheic community comprising early larval stages of Coleoptera and Diptera,
374 as well as, more representatives of the permanent meiofauna, including Rotifera, Copepoda and
375 Acari. This community collectively contributed 27% of the whole-stream production, stressing
376 the need to include the hyporheos to yield accurate whole-stream production budgets

377 Our estimates of hyporheic productivity in the forested stream can be compared with the
378 study of Majdi et al. 2017 performed in two sandy forested streams in north-western Germany

379 (i.e., Furlbach and the Ems Table 4). This study sampled both the benthic and hyporheic
380 compartments simultaneously, collected the entire invertebrate distribution, and assessed each
381 compartment's relative contribution to production. Compared to this study, the productivity of
382 the hyporheic community in our forested stream was within the range of values observed in the
383 Furlbach and the Ems (Table 4). However, the relative contribution of the hyporheic
384 compartment to whole-stream invertebrate production (27%) was approximately half compared
385 to Furlbach and the Ems (48-51%) (Table 4). We attribute these differences to 2 main reasons.
386 First, the differing substrate types present in the streams. The sediment of the forested stream
387 was predominantly coarse gravel and cobble, in contrast to the sandy substrates of the Furlbach
388 and the Ems. The presence of sand and the absence of a riffle-pool sequence have been found to
389 favour meiofaunal size invertebrates (Poff et al. 1993). The differing substrate may therefore
390 explain the lower relative contribution of benthic macroinvertebrates, and therefore of the
391 benthic compartment to whole-stream production in the sandy streams, in comparison to the
392 results obtained in this study. Furthermore, differences in the sampling strategy in the studies
393 could have contributed to exacerbate observed discrepancies. While we employed both a Surber
394 sampler and a corer, Majdi et al. (2017) solely employed a sediment corer which might have
395 overlooked the contribution of larger and more mobile invertebrates such as Ephemeroptera and
396 gammarids.

397 Finally, our study is among the few that directly compares hyporheic versus benthic
398 production by including both the macroinvertebrates and the permanent meiofauna. The first
399 pioneer study which quantified stream meiofauna production was conducted by Stead et al.
400 (2005) which demonstrated the meiofauna can contribute significantly, comprising 52% of
401 benthic production. Despite this study was conducted in the benthic sediment (0-5 cm depth), its

402 findings stimulated subsequent studies to expand the sampling boundaries and include the
403 hyporheic zone in secondary production quantification. Nevertheless, since Stead et al. (2005)
404 there has been a distinction between studies focusing on the hyporheic compartment and
405 sampling the meiofauna and those sampling the benthic compartment and the macrofauna.
406 However, both fractions are present in the compartments, thus sampling strategies should aim to
407 include both in both compartments (Schmid-Araya et al. 2020). By applying combined sampling
408 strategies, we showed that in both streams the majority of production occurs in the benthic
409 compartment and is sustained by the benthic macrofauna. However, in permeable sediment, the
410 meiofauna still contribute substantially to benthic and hyporheic production and must therefore
411 be included in secondary production estimates (Tod and Schmid-Araya 2009, Majdi et al. 2017,
412 Schmid-Araya et al. 2020, Pasqualini et al. 2023). Consequently, we advocate for a more
413 comprehensive understanding of the entire stream ecosystem, through a holistic approach that
414 encompasses not only the two compartments, but also the entire length distribution.

415

416 **Broader implication**

417 Our results bring further evidence that while resource availability is the major driver of
418 production differences, sediment characteristics are the dominant factor controlling secondary
419 production repartition between compartments within streams. We demonstrate that fine sediment
420 deposition derived from agricultural land use, by strongly changing the environmental conditions
421 in the hyporheic compartment, reduces the hyporheic compartment functionality. Given that fine
422 sediment deposition and enhanced nutrients often co-occur in agricultural streams, and that
423 agricultural streams are becoming increasingly common (Tank et al. 2021), it is likely that an
424 increasing number of streams will experience a decline in hyporheic functionality coupled with

425 an increase in benthic functionality, leading to unnaturally high production levels. Therefore, it is
426 of the great importance to take immediate action to preserve both the functional and habitat role
427 of both compartments by improving current agricultural practices. Land-use practices in the
428 catchment exert a fundamental influence on the lotic ecosystem. Consequently, to improve
429 stream's conditions, it is necessary to implement actions at the catchment scale. Nevertheless,
430 local actions such as the establishment of riparian buffer zones, which simultaneously reduce
431 nutrients (Hoffmann et al. 2009) and fine sediment inputs (Dunn et al. 2022), should also be
432 implemented. These actions will not only improve habitat conditions for biodiversity, but will
433 also improve streams functionality.

434

435 **ACKNOWLEDGEMENTS**

436 Author contributions: JP, MB, NM and MW conceived of the study and designed the
437 field methods, JP conducted the fieldwork and analyzed the data. JP and MB led the writing of
438 the paper with contributions from all authors. The authors thank S. Bauth, S. Willige,
439 A. Kneur, H. Matthes, K. Reinmann, R. Degenhardt and S. Jolitz-Seif for their assistance with
440 field and laboratory work. A. Hoff and the GEWANA department for analysis of nutrients
441 in the laboratory. D. Graeber provided helpful comments on the sampling design.

442 This research was funded by the research program “Changing Earth – Sustaining our
443 Future” of the German Helmholtz Association and was part of the Integration Platform
444 ‘Freshwater resources’. The authors declare no conflict of interest.

445

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627

628 **FIGURE CAPTIONS**

629 Fig. 1. Nutrient concentrations measured bimonthly in the compartments of the forested (A-E)
630 and agricultural stream (F-L). Different lowercase letters indicate significantly different
631 values at the $p < 0.05$ level (Tukey's HSD post hoc test) within the same stream. Please
632 note the different scales on the y-axis. Values in the benthic compartment of the forested
633 stream are missing because of the infeasibility to install firmly a tube at a depth of 5 cm.

634 Fig. 2. NMDS ordination of macrofauna communities in the forested stream (A) and in the
635 agricultural stream (B). NMDS of permanent meiofauna communities in the forested
636 stream (C) and in the agricultural stream (D). Letters indicate the taxa identified with
637 SIMPER that contributed the most to differences among compartments. a *Agapetus*
638 *fuscipes*, b *Amphinemura* spp., c *Ancylus fluviatilis*, d *Baetis* spp., e Ceratopogonidae, f
639 Chironomidae, g *Chironomini* spp., h *Elmis* spp., i *Esolus* spp., j *Glossosoma* spp., k
640 *Ibisia marginata*, l *Leuctra* spp., m *Limnius* spp., n Nematoda, o *Nemoura* spp., p
641 Oligochaeta, q Orthocladiinae, r *Potamopyrgus antipodarum*, s *Prodiamesa olivacea*, t
642 *Sericostoma* sp., u Simuliidae, v *Tanytarsini* spp., z Tanypodinae. NMDS = nonmetric
643 multidimensional scaling.

644 Fig. 3. Median and 95 % CI of annual whole-stream invertebrate production (A). Mean relative
645 contribution of the compartments to annual whole-stream invertebrate production (B).

646 DM = dry mass.

647 Fig. 4. Cumulative percentage of taxa's contribution to whole-stream production in the benthic
648 and hyporheic compartments of the 2 streams.

649 Table 2 Secondary production, biomass and abundance of taxa collected in the forested stream. Values are annual medians and 95%
 650 CI. Values are rounded to nearest decimal except when <1. DM = dry mass.

Taxon	Abundance (ind./ m ²)		Biomass (mg/m ²)		Production (mg DM m ⁻² y ⁻¹)	
	Benthic	Hyporheic	Benthic	Hyporheic	Benthic	Hyporheic
<i>Elmis</i> sp.	70 (4 - 162)	0	2.4 (0.058 - 6)	0	33 (0 - 103)	60 (0 - 245)
<i>Esolus</i> sp.	388 (156 - 631)	665 (197 - 1295)	25.1 (6.7 - 47.2)	22.3 (6.5 - 39.9)	0	0
<i>Limnius</i> sp.	153 (43 - 339)	162 (19 - 319)	26.2 (9 - 47.3)	7.9 (0.047 - 19.9)	0	0
<i>Oulimnius</i> sp.	0	112 (0 - 562)	0	88.8 (0 - 444)	0	0
Ceratopogonidae	157 (13 - 422)	199 (55 - 369)	0.786 (0.035 - 2.7)	2 (0.138 - 4.3)	1220 (579 - 1941)	234 (36 - 453)
Chironomidae	3603 (1711 - 6979)	2057 (629 - 4953)	63.2 (22.6 - 125.1)	30.1 (6.1 - 83.1)	748 (220 - 1646)	520 (151 - 1273)
<i>Chironomini</i> sp.	539 (72 - 1552)	299 (38 - 771)	2.6 (0.103 - 9)	2.5 (0.082 - 7.7)	1069 (100 - 3117)	33 (0 - 92)
<i>Ibisia marginata</i>	80 (21 - 157)	0	41.2 (8.9 - 94.9)	0	0	0
Orthoclaadiinae	509 (279 - 777)	139 (0 - 565)	37.4 (12.5 - 79)	0.921 (0 - 3)	263 (131 - 424)	30 (0 - 74)
Simuliidae	102 (11 - 246)	0	20 (0.915 - 53.8)	0	0	0
Tanypodinae	874 (226 - 2246)	313 (42 - 697)	24.7 (6.8 - 57.6)	9.9 (0.897 - 28.4)	417 (97 - 921)	88 (0 - 215)
Tanytarsini	1581 (409 - 3342)	470 (162 - 804)	18.4 (3.8 - 42.8)	2.7 (0.322 - 5.5)	3628 (181 - 9550)	263 (4 - 1017)
<i>Baetis</i> sp.	295 (100 - 564)	104 (0 - 358)	34.7 (7.4 - 76.1)	0.95 (0 - 3.5)	0	0
<i>Ephemerella mucronata</i>	164 (22 - 392)	73 (0 - 310)	18.1 (0.825 - 42.7)	3.9 (0 - 15)	0	0
<i>Habroleptoides confusa</i>	78 (16 - 181)	0	34.3 (6.3 - 78.8)	0	0	0
<i>Ancylus fluviatilis</i>	277 (45 - 496)	0	174.1 (36.2 - 371.2)	0	0	0
Veliger (<i>Ancylus fluviatilis</i>)	308 (86 - 652)	76 (10 - 207)	30.3 (4.7 - 103.7)	6.9 (0.536 - 20.7)	0	0
Oligochaeta	137 (48 - 264)	0	41.7 (4.7 - 103)	0	339 (40 - 835)	0
<i>Agapetus fuscipes</i>	434 (111 - 1276)	64 (0 - 141)	29 (8.8 - 75)	4.4 (0 - 10)	0	0
<i>Amphinemura</i> sp.	406 (68 - 1210)	0	22.2 (5.2 - 45.7)	0	0	0
<i>Anomalopterygella chauviniana</i>	94 (5 - 259)	0	35.3 (5.5 - 85.3)	0	0	0
Drusinae	116 (0 - 533)	0	9.1 (0 - 39.8)	0	0	0
<i>Glossosoma</i> sp.	84 (19 - 163)	0	2.6 (0.447 - 5.5)	0	0	0

<i>Hydropsyche</i> sp.	86 (8 - 241)	0	77.6 (5.1 - 252.3)	0	0	0
<i>Leuctra</i> sp.	372 (116 - 903)	192 (9 - 508)	13.8 (5.9 - 22.3)	3.8 (0.015 - 12.8)	0	0
<i>Nemoura</i> sp.	382 (72 - 823)	89 (0 - 255)	5.5 (1.3 - 10.3)	0.632 (0 - 1.7)	0	0
<i>Sericostoma</i> sp.	89 (37 - 157)	0	156.7 (66.4 - 278.6)	0	0	0
Acari	970 (582 - 1807)	1408 (620 - 3017)	10.1 (3 - 24.5)	11.6 (1.8 - 30.8)	92 (1 - 391)	2 (2 - 4)
Alona	179 (9 - 543)	64 (9 - 130)	0.233 (0.012 - 0.638)	0.071 (0.004 - 0.178)	43 (12 - 99)	0
Copepoda	3939 (1543 - 8486)	5363 (2246 - 1.1e+04)	4.8 (1.8 - 11.7)	5.6 (2 - 11.6)	547 (207 - 1109)	114 (29 - 252)
Copepoda: Naupli	625 (214 - 1133)	1614 (719 - 3263)	0.002 (0.001 - 0.003)	0.004 (0.001 - 0.01)	6 (2 - 14)	2 (0.1 - 5)
Nematoda	8.2e+04 (3.5e+04 - 1.5e+05)	1.2e+05 (5.6e+04 - 2e+05)	2.2 (0.647 - 5.9)	2.1 (0.734 - 3.6)	2619 (1302 - 4322)	898 (472 - 1464)
Ostracoda	110 (14 - 280)	285 (21 - 614)	0.176 (0.007 - 0.396)	0.264 (0.005 - 0.554)	169 (38 - 373)	59 (3 - 130)
Rotifera	1.3e+05 (6.2e+04 - 2.3e+05)	2.1e+05 (1e+05 - 3.5e+05)	2.3 (1.1 - 3.5)	3.6 (1.6 - 5.8)	312 (68 - 904)	98 (38 - 186)
Tardigrada	6336 (2311 - 1.1e+04)	1.5e+04 (5307 - 4.3e+04)	0.387 (0.136 - 0.651)	0.831 (0.291 - 2.2)	4 (0.7 - 10)	6 (1 - 13)

651

652 Table 3 Secondary production, biomass and abundance of taxa collected in the agricultural stream. Values are annual medians and
 653 95% CI. Values are rounded to nearest decimal except when <1. DM = dry mass.

Taxon	Abundance (ind./ m ²)		Biomass (mg/m ²)		Production (mg DM m ⁻² y ⁻¹)	
	Benthic	Hyporheic	Benthic	Hyporheic	Benthic	Hyporheic
<i>A. aquaticus</i>	138 (0 – 358)	0	96 (0 – 220)	0	185 (0 – 405)	0
<i>Gammarus pulex</i>	94 (19 – 200)	0	210 (20 – 477)	0	543 (85 – 1166)	0
<i>Elmis</i> sp.	256 (0 – 893)	104 (0 – 437)	4 (0 – 12)	6 (0 – 31)	33 (0 – 103)	60 (0 – 245)
<i>Apsectrotanypus trifascipennis</i>	58 (19 – 107)	0	27 (11 – 43)	0	119 (49 – 180)	0
Ceratopogonidae	7218 (3581 – 1.1e+04)	1385 (169 – 3128)	95 (38 – 161)	16 (1 – 33)	1220 (579 – 1941)	234 (36 – 453)
Chironomidae	2471 (633 – 5336)	1715 (483 – 3300)	79 (18 – 186)	52 (14 – 136)	748 (220 – 1646)	520 (151 – 1273)
<i>Chironomini</i> sp.	3542 (1192 – 6841)	418 (0 – 957)	139 (10 – 423)	2 (0 – 6)	1069 (100 – 3117)	33 (0 – 92)
Orthoclaadiinae	242 (53 – 631)	163 (0 – 500)	52 (24 – 85)	2 (0 – 6)	263 (131 – 424)	30 (0 – 74)
<i>Prodiamesa olivacea</i>	201 (0 – 718)	135 (0 – 409)	75 (0 – 347)	55 (0 – 191)	280 (0 – 1276)	240 (0 – 768)
Tanypodinae	971 (208 – 2193)	202 (0 – 598)	47 (8 – 111.5)	10 (0 – 26)	417 (97 – 921)	88 (0 – 215)
Tanytarsini	4777 (1001 – 1e+04)	1256 (0 – 3838)	336 (10 – 1098)	15 (0 – 63)	3628 (181 – 9550)	263 (4 – 1017)
<i>P. antipodarum</i>	1092 (117 – 3384)	0	9574 (1323 – 2.8e+04)	0	14957 (2705 – 41319)	0
Naididae	142 (0 – 368)	0	624 (0 – 1628)	0	1294 (0 – 3101)	0
Oligochaeta	158 (24 – 339)	0	80 (5 – 217)	0	339 (40 – 835)	0
<i>Pisidium</i> sp.	213 (5 – 496)	0	528 (30 – 1284)	0	1143 (86 – 2641)	0
<i>Limnephilus</i> sp.	103 (3 – 275)	0	12 (0.662 – 31)	0	55 (4 – 131)	0
Acari	238 (53 – 596)	157 (111 – 282)	4 (0.008 – 11)	0.04 (0.03 – 0.06)	92 (1 – 391)	2 (2 – 4)
Alona	926 (166 – 2362)	0	1 (0.23 – 4)	0	43 (12 – 99)	0 (0 – 0)
Copepoda	5998 (2728 – 1e+04)	1512 (262 – 3342)	22 (8 – 52)	4 (0.4 – 10)	547 (207 – 1109)	114 (29 – 252)
Copepoda: Naupli	2813 (991 – 5630)	805 (162 – 1986)	0.035 (0.009 – 0.08)	0.009 (0 – 0.024)	6 (2 – 14)	2 (0.1 – 5)
Nematoda	8.4e+05 (3.6e+05 – 1.4e+06)	2.7e+05 (1.3e+05 – 4.6e+05)	21 (10 – 34)	7 (4 – 12)	2619 (1302 – 4322)	898 (472 – 1464)
Ostracoda	2082 (568 – 4158)	602 (160 – 1622)	5 (0.95 – 12)	2 (0.03 – 4)	169 (38 – 373)	59 (3 – 130)

Rotifera	7.3e+04 (1.7e+04 – 1.8e+05)	2.8e+04 (7961 – 5.9e+04)	3 (0.56 – 9)	0.89 (0.29 – 2)	312 (68 – 904)	98 (38 – 186)
Tardigrada	665 (90 – 1968)	1081 (125 – 2892)	0.04 (0.002 – 0.13)	0.06 (0.01 – 0.16)	4 (0.7 – 10)	6 (1 – 13)

654

655 Table 4 Stream characteristics and mean secondary production estimates from studies conducted in headwater streams with sampling
 656 strategies comparable to those used in this study. Nutrient and temperature values are annual means (\pm SD), except temperature data
 657 from the Lone Oak and Lambourn, for which temperature range is reported. To convert production values expressed in Carbon to dry
 658 mass (DM) a factor of conversion of 2 had been applied DM = Carbon*2. – no data.

Stream	Environmental characteristics				Production		
	Sediment/ habitat	N-NO ₃ (mg/L)	SRP/P-PO ₄ (μ g/L)	Temperature ($^{\circ}$ C)	Benthic	Hyporheic	Total
					P (g DM m ⁻² y ⁻¹)		
Ems ¹	Sand	1.45 \pm 0.6	25.9 \pm 5.9	9.5 \pm 1.32	2.54	2.62	5.16
Furlbach ¹	Sand	3.69 \pm 2.82	5.32 \pm 4.08	7.88 \pm 2.11	5.7	5.22	10.92
Lone Oak ²	Gravel to large cobble	*Nutrient levels under limit of detection		4 - 15	5.22	–	5.22

Forested stream ³	Coarse gravel-cobble	1.9 ± 0.6	6.3 ± 2.9 ⁵	7.6 ± 3.2	5.66	2.05	7.7
Lambourn ⁴	Gravel	Hypernutrified		7 - 15	–	–	22.55
Agricultural stream ³	Fine sand	9.3 ± 2.3	31 ± 15 ⁵	9.3 ± 6.2	21.47	1.47	22.93
Lambourn ⁴	Gravel/macrophyte	Hypernutrified		7 - 15	–	–	64.99

659

660 ¹ Majdi et al. 2017, ² Stead 2005, ³ This study, ⁴ Tod and Schmid-Araya 2009 * NA in the study. Information available in Reiss and

661 Schmid-Araya 2008,⁵ values are SRP not P-PO4