The Benthic Trophic Corner Stone Compartment in POPs Transfer from Abiotic Environment to Higher Trophic Levels—Trichoptera and Ephemeroptera Pre-Alert Indicator Role

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Abstract: Persistent organic pollutants (POPs) have been at the forefront of environmental contamination research even before their ban in 2001 at the Stockholm Convention. Their relation to different compartments of the environment (biotic and abiotic) has been thoroughly investigated. This article aims to identify whether the benthos could represent a reliable indicator of environmental contamination with POPs and to highlight its potential transfer role between abiotic and upper trophic compartments—benthos feeders. In this regard, we determined that the Ephemeroptera samples have higher concentrations \((p < 0.05)\) of \(\Sigma PCB\), \(\Sigma HCH\), and \(\Sigma DDT\) than sediment samples while Trichoptera samples have higher concentrations \((p < 0.05)\) only in the case of \(\Sigma PCB\) and \(\Sigma DDT\). This, along with the fact that the frequency of detection for POPs is similar between the sample types (sediments, Trichoptera, and Ephemeroptera), makes the benthos samples valuable indicators of contamination with sediment samples working as complementary information about how recent the contamination is.

Keywords: lotic ecosystem; organochlorine pesticides; polychlorinated biphenyls; sediments; benthos

1. Introduction

In spite of the fact that in the Lower Danube Basin the footprints of \(Homo sapiens\) actions date back to 180,000 BC, and over time the adverse effects of these activities on habitats and biodiversity have become more and more aggressive and complicated [1–7], the human-dominated management of streams and rivers in the Anthropocene is not impossible if the management approach is integrated and adapted on-site [8–10]. One of the main second-order tributaries of the Danube River is the Mureș River, its upper and middle watershed being placed in the amphitheater-shape Transylvanian Depression, ringed by the South-Eastern Carpathians, and inhabited by over seven million people [11]; it is a space with variable and important human impact [12–15]. The different types of human impact presence here are more or less addressed by scientific studies; however, studies on the presence and effects of the xenobiotic chemical compounds, namely persistent organic pollutants (POPs) [16], a relatively new threat around the globe [17], in the studied area on aquatic life and their environment are scarce.

POPs, which were recognized as one of the main threats due to their toxicity and diverse environmental and human health risks [18], are mainly manmade composite substances characterized by high resistance to photolytic, chemic, biologic, and mechanic deterioration [19–21]. Being volatile, POPs are carried through air at far-reaching distances from the emission sources [22]; moreover, POPs pollute lotic systems by the way of groundwater or by direct discharge [18,23] and are accumulated in sediment [24], which plays a major role in the water quality of rivers and the quality of biota [4,25,26], adsorbed in suspended particles and on/into the aquatic organisms [27–29]. Owing to the hydrophobic
nature of POPs, they concentrate in the sediment, thus changing this abiotic compartment into a pool for pollutants \[24,27,30\]. This is the argument for why sediment is the main source of POPs and a risk to the benthic organisms \[31,32\] that bioaccumulate them throughout their lifecycle by simple contact with sediment and water, and through food intake \[33\]. The contamination with POPs is granted an unequivocal mechanism due to their hydrophobic quality and reciprocal action of sediment with benthic organisms, considered to be the principal hazard path \[29\]. Through the feeding on insect larvae by higher food trophic net organisms, POPs begin to bioconcentrate from one level to another \[33,34\]. It is accepted that POPs bioaccumulate in the animals’ fatty tissues because of their lipophilic nature \[18\], including in edible organisms such as fish \[35,36\]. We suggest that as a rationale effect, the trophic niches created by benthic organisms represent a compartment with a functional-transfer role of POPs between the abiotic and higher-level trophic compartments ending on the human plates, the insect larvae representing, due to their relatively high biomass and turn-over, a very important taxonomic group for the circulation of POPs in nature.

Until their ban in 2001 through the Stockholm Convention \[18\], POPs were considered a success in the field of insecticide use. This was the case for organochlorine pesticides (OCPs), in the electric industry, and polychlorobiphenyls (PCBs). Because of their long period of use, POPs became ubiquitous compounds in air, water, and soil \[34,37\]. Due to this, POPs were deemed to be a major threat, while the research in this field has acknowledged their impact and secondary negative effects on the environment and on human health \[38–42\]. It has been determined that the presence of POPs in the environment leads to its degradation which in turn affects the health and integrity of every organism through body anomalies, physiological imbalances, disturbances in sex ratios, impaired reproduction, low fertility, and cancer \[39,42\].

In this study, we focused on the benthic functional compartment, namely certain components: the insect larvae from Trichoptera and Ephemeroptera orders as these organisms are common in the trophic nets of benthic lotic habitats. Trichoptera and Ephemeroptera are representative in the studied area and have long been considered as water pollution indicators \[43,44\]; they have been used as indicators of environmental contamination with trace elements \[45,46\]. Furthermore, insects are considered to be one of the main groups used in biodiversity studies due to their high individual and taxonomic group count and their complexity \[47\]. Their importance is derived from their trophic net position, being the link between collectors of organic matter from the substrate, filter feeders, herbivores, detritivores, and predators \[48,49\]. It can also be noted that functional feeding groups (FFGs) can influence contaminant accumulation \[50\] due to their preferred feeding style.

All Trichoptera species are found in freshwater on every continent except Antarctica \[51\]. Trichoptera can be used to obtain information about pollution because they are susceptible to environmental changes. The capacity of their larvae to accumulate and bioamplify POPs from the sediments can be used to indicate the presence and level of pollution. In the larval stage, Trichoptera species are aquatic and benthic with a period of two months to two years of development. Because of this long larval stage, the larger part of the nutrients gathered by the individual occurs during this period. Trichoptera larvae, depending on the species, are designed as different types of feeders, such as collectors of organic materials from the substrate, filter feeders, herbivores, detritivores, or predators. Because collecting feeders concentrate the organic matter inside their bodies and because the diversity and number of individuals of this group are high, they constitute the main source of nutrients for other aquatic organisms \[51,52\].

The food demands for the species of the Ephemeroptera are similar to those of the Trichoptera. Ephemeroptera species also can be used to obtain data about pollution because they are susceptible to such environmental modifications. Their larvae capacity to accumulate and bioamplify POPs from the sediments can be used to indicate the presence and level of pollution. The lack of certain species can offer information about the nature of pollution if the characteristics of the pollutants are known \[53,54\]. This can be further expanded
upon when considering that studies involving sediment samples can be accompanied by studies that identify benthos contamination with POPs for a better understanding of the underlining processes. The individuals from this order can have a larval period of up to two years and are found in aquatic habitats around the globe, with the exception of the Antarctic [55]. Most species from the Ephemeroptera order are collecting and grazing feeders and can even consume detritus [56,57].

Based on a few key studies [32,58], it has been reported that bottom surface sediment samples confer good data with regards to river environment pollution, being a good indicator of pollutants.

The aim of our study was to determine if Trichoptera and Ephemeroptera larvae can be good indicators of habitat contamination with POPs because of their already established role in water quality assessment for other bioaccumulating types of pollutants [36] and negatively influence in time the fish [59–65] traditionally used as food by humans in the studied area [66–68]. We expected that these indicators, through their bioaccumulation and bioamplification capacity, can reveal the aquatic ecosystem contamination with POPs. The advantage of working with these larvae is that they are relatively easy to be sampled, and because of their relatively low mobility, they offer the chance to relate the results of POP contamination to specific river sectors.

2. Materials and Methods

2.1. Study Area and Sample Collection

The Mureș River is the largest tributary of the Tisza River, constituting an important sub-basin of the Danube Basin, with a total length of 716 km and a multiannual flow of 177 m$^3$ s$^{-1}$, in the Romanian segment [69]. The Mureș River basin contains over seven million people and accumulates the majority of the human communities’ pollutants [11,12]. The origins here of potential POPs pollution are the chemical industry, intensive livestock farms, waste warehouses, water treatment plants, metallurgical factories, electrical transformer plant facilities, etc. [70].

Sediment, Ephemeroptera, and Trichoptera samples were collected on 7–16 August 2016 from the following sampling sites: M1, M2, M3, M4, M6, M7, M11, M12, and M14 on the Mureș River (Figure 1).

The establishment of the sampling stations was based on the following criteria: the presence of point sources of pollution with POPs (industrial sources, landfills, wastewater treatment plants, untreated municipal wastewater discharges, direct discharges of industrial wastewater, animal husbandry pollution sources); areas with intensive agriculture; areas with high density of domestic animals.

M1–645 m altitude, average width of the minor riverbed 25 m (max. 30 m, min. 20 m), The river has a mountainous appearance, with a slope in the center of the riverbed; the substratum consists of boulders and medium-sized stones; gravel and sandy areas appear toward the banks. There is a strand of willows on the banks. The benthic macroinvertebrate community consists of Ephemeroptera (39.41%), Plecoptera (25.29%), Trichoptera (12.94%), Amphipoda (7.06%), Planariidae (6.47%), Gastropoda (4.12%, Ancylus fluviatilis), Chironomidae (4.71%).

M2–358 m altitude, average width of the minor riverbed 90 m (max. 110 m, min. 70 m), linear flow, relatively uniform. The substrate consists predominantly of medium-sized stones, boulders appear in the center of the riverbed, and toward the banks is gravel covered with bioderm and the presence of sandy surfaces. Riparian vegetation consists of willows. The benthic macroinvertebrate community consists of Ephemeroptera (36.28%), Trichoptera (28.84%), Chironomidae (14.88%), Oligochaeta (13.49%), Planariidae (4.19%), Gastropoda (2.32%, Ancylus fluviatilis).

M3–294 m altitude, an average width of the minor riverbed of 40 m (max. 60 m, min. 30 m) and a linear flow with ups and downs, the substratum consists mainly of medium stones on a bed of sand and mud. In the center of the riverbed are boulders in some places, and on the banks, there are surfaces with gravel and sand. The left bank
has grassy vegetation, and on the right bank, there is a strip of willows. The benthic macroinvertebrate community consists of Ephemeroptera (50.91%), Trichoptera (29.09%), Chironomidae (10.30%), Oligochaeta (7.27%), and Blepharoceridae (2.43%).

Figure 1. The sample sites with geographical coordinates are: M1 (46 94567–025 29396), M2 (46 73324–024 70280), M3 (46 50931–024 47209), M4 (46 44489–024 04081), M6 (46 42958–023 97672), M7 (46 38317–023 82367), M11 (46 38317–023 82367), M12 (46 04262–023 55594), M14 (45 9626–022 81669).

M4–277 m altitude, average width of the minor riverbed 60 m (max. 80 m, min. 40 m), slower linear flow, with unraveling, the substrate consists of mud, clay, fine sand, and toward the center of the riverbed there are surfaces with small, medium, and large stones (marls) on the muddy bed. The banks are steep; on the bank, there is grassy vegetation, and on the right bank, there are willows. The benthic macroinvertebrate community consists of Trichoptera (30.88%), Ephemeroptera (28.57%), Planaridae (18.89%), Chironomidae (11.52%), Oligochaeta (9.68%), Bivalvia (0.46%, Sinanodonta woodiana).

M6–268 m altitude, average width of the minor riverbed 65 m (max. 85 m, min. 45 m), slower smooth flow; in the riverbed, there is a bottom threshold, and the substrate is formed of large and medium stones on mud bed. The course is meandering and the riparian vegetation is grassy, but willows also appear in certain places. The benthic macroinvertebrate community is formed of Ephemeroptera (34.31%), Planaridae (20.59%), Trichoptera (19.61%), Chironomidae (15.69%), Oligochaeta (9.80%).

M7–259 m altitude, average width of the minor riverbed 65 m (max. 85 m, min. 45 m), the substratum consists of gravel on a bed of sand, and on the banks, there is grassy vegetation and willows. The benthic macroinvertebrate community consists of Ephemeroptera (40.71%), Chironomidae (30.71%), Oligochaeta (22.14%), Trichoptera (5%), Planaridae (0.72%), Gastropoda (0.72%, Ancylus fluviatilis).

M11–278 m altitude, average width of the minor riverbed 70 m (max. 90 m, min. 50 m), the substrate consists predominantly of gravel on a bed of sand and mud, and sandy
surfaces appear on the banks. The benthic macroinvertebrate community consists of Trichoptera (25.0%), Chironomidae (24.40%), Ephemeroptera (21.39%), Oligochaeta (20.48%), Amphipoda (5.72%), Tipulidae (2.11%), Gastropoda (0.90%, Physella acuta).

M12–218 m altitude, average width of the minor riverbed 75 m (max. 95 m, min. 55 m), the substrate consists of gravel on a bed of sand, and on the shore, the gravel is covered with bioderm. The benthic macroinvertebrate community consists of chironomids (30.52%), Trichoptera (22.09%), oligochaetes (21.29%), Ephemeroptera (12.45%), nematodes (9.63%), gastropods (2.01%, Physella acuta), beetles (2.01%).

M14–171 m altitude, average width of the minor riverbed 75 m (max. 90 m, min. 55 m), the substrate consists of gravel on a bed of sand and mud; in the bed, there are depressions/pots, on the banks, there is a stripe of willows, and the benthic macroinvertebrate community is formed from Chironomidae (29.09%), Trichoptera (25.45%), Oligochaeta (20.0%), Gastropoda (15.15%, Physella acuta), Odonata (10.31%).

The sediment (for the analysis of the amount of POPs) was collected manually from the surface of the riverbed substrate (at 50 cm water depth), with the help of a plastic vessel; the supernatant was removed, and 200 g of sediment were placed in the sample vessel, frozen in the field at $-20^\circ$C, then stored in the laboratory at $-50^\circ$C.

The samples were collected manually: the insect larvae were washed to eliminate the sediment particles from their bodies, which can bring supplementary quantities of POPs, and frozen at $-20^\circ$C immediately after collection, then transported to the laboratory where they were stored at $-50^\circ$C. The identification of insect larvae was performed at the order level, based on the morphological characters analyzed using an Olympus (150×) stereomicroscope. Afterward, the samples were separated in Petri dishes that were washed beforehand with acetone:cyclohexane 1:1 and set to dry overnight in a desiccator at room temperature (25°C) before extraction of POPs. Ephemeroptera and Trichoptera larvae of relatively homogenous size, for each group, were selected in order to avoid influencing the POP accumulation results. The biomass, expressed as dry weight, of the Ephemeroptera samples from which POPs were extracted is as follows: M1–0.8416 g (68 individuals), M2–0.7281 g (120 ind.), M3–1.1966 g (91 ind.), M4–0.2339 g (59 ind.), M7–0.1927 g (70 ind.), M11–0.1229 g (69 ind.), and M12–0.0915 g (73 ind.). In case of Trichoptera samples, the biomass, expressed as dry weight, is as follows: M1–0.8802 g (60 ind.), M2–0.2625 g (38 ind.), M3–0.2022 g (34 ind.), M4–0.4545 g (83 ind.), M6–0.1966 g (41 ind.), M11–2.0734 g (57 ind.), M12–0.7264 g (48 ind.), and M14–1.1978 g (42 ind.).

2.2. Reagents and Standards

The solvents (acetone, cyclohexane, and ultrapure water) of HPLC grade purity were purchased from BioAqua (Târgu Mureş, Romania), and the NaCl (99.9%), green malachite (99.9%), and sulfuric acid (96%) were from Sigma-Aldrich. The PCB congeners (28, 29, 31, 47, 52, 56, 66, 74, 99, 101, 105, 110, 112, 114, 128, 136, 137, 138, 141, 149, 151, 153, 156, 157, 160, 180, 183, 187, 189, 194, 196, 199, 206, 207, 209), DDT (o, p'-DDE, p, p'-DDE, o, p'-DDD, p, p'-DDT, p, p'-DDT), HCH (α-HCH, β-HCH, γ-HCH, δ-HCH, ε-HCH), chlordane (oxychlordane, trans-chlordane, cis-chlordane), HCB, and mirex standards were obtained from LGC Standards (Sibiu, Romania).

2.3. Sample Preparation

For the extraction of OCPs and PCBs from sediment and from benthos (insect larvae), we used a method involving 3:2 cyclohexane:acetone and ultrasonication [71]. Sediment was dried overnight in a desiccator, ground up with a mortar and pestle, and 5 g was used for extraction. In the case of benthos samples, the entire biomass was used for extraction. Internal standards (PCB 29, 112, 207) were added, and then the powder was extracted twice by ultrasonication using a Q500 sonicator (QSonica) (Newtown, CT, USA) and cyclohexane:acetone (3:2). The extract was separated from the sediment using an NF 800 R (Nüve) (Ankara, Turkey) centrifuge for 10 min at 2000 rpm; the supernatant (liquid extract) was transferred and calibrated to volume (1 mL) by evaporation. The extract was...
purified using clean sulfuric acid (96%) (Sigma-Aldrich) (București, Romania), and the resulting solution was calibrated to volume (1 mL) by evaporation. A volume of 0.5 mL was transferred to vials and stored for a maximum of two weeks before injection.

2.4. Instrument Analysis

The quantification of OCPs and PCBs was performed on a 7890 B (Agilent) (București, Romania) gas chromatograph (GC) coupled to a 7010 A (Agilent) triple quadrupole mass spectrometry (MS) system (București, Romania). The GC column was DB-5ms 60 m, 0.25 mm, 0.25 µm (Agilent) with helium as the carrier and quench gas, while nitrogen was used as the collision gas. The OCPs and PCBs were identified by running separate standards and observing the specific retention time and by collision-induced dissociation and observing the specific pattern of ions [72–74]. The inlet ran in splitless mode with 54.3 mL min\(^{-1}\) carrier gas, the column carrier gas was set to 1.3 mL min\(^{-1}\), the quench gas was set to 2.25 mL min\(^{-1}\), and the collision gas was 1.5 mL min\(^{-1}\). The oven temperature was programmed to increase from 90 °C with a 2 min hold to 180 °C with a 2 min hold (25 °C−min\(^{-1}\)), 220 °C with a 2 min hold (1.5 °C−min\(^{-1}\)), 275 °C with a 2 min hold (3 °C−min\(^{-1}\)), and finally 300 °C with a 4 min hold (25 °C−min\(^{-1}\)).

2.5. Quality Control

The sediment samples were analyzed in three technical replicates in an analytical series that included six spiked samples (river sediment), with all the investigated analytes for recoveries, two blinds (river sediment), and four solvent blanks. The standard curve linearity (R\(^2\)) was above 0.993 for all analytes. The mean recovery percentage for sediment ranged from 81 to 119.5% for PCBs, 84.5 to 100.9% for DDTs, 81.8 to 100.4% for HCHs, 111.8 to 114.7% for chlordane, 87.3% for HCB, and 89.4% for mirex. The relative standard deviation values (RSD%) were situated between 0.9 and 18.3% for PCBs, 2.7 and 12.8% for DDTs, 0.6 and 2.7% for HCHs, 2 and 7.5% for chlordane, 2.7% for HCB, and 1.8% for mirex. No analytes were detected in the blanks. The data are reported as the arithmetic mean of the three technical replicates.

The benthos samples were separated by order into Ephemeroptera and Trichoptera and analyzed in an analytical series that included three spiked samples (Trichoptera samples), with all the investigated analytes for recoveries, one blind (Trichoptera sample), and four solvent blanks. The standard curve linearity (R\(^2\)) was above 0.996 for all analytes. The mean recovery percentage for benthos ranged from 76.8 to 116.4% for PCBs, 70 to 83.2% for DDTs, 70.9 to 72% for HCHs, 96.8 to 99.4% for chlordane, 105.2% for HCB, and 71.9% for mirex. The relative standard deviation values (RSD%) were situated between 0.5 and 7.5% for PCBs, 2.3 and 5.8% for DDTs, 0.3 and 3.2% for HCHs, 1.4 and 6.5% for chlordane, 2.4% for HCB, and 2.5% for mirex. No analytes were detected in the blanks.

All the reported concentrations were higher than the limit of quantitation (LOQ) [75].

2.6. Data Analysis

We investigated the ratios of p,p’-DDD/p,p’-DDE [76] and (p,p’-DDE + p,p’-DDD)/p,p’-DDT [77] and their differences between the analyzed sample types; the ratios were calculated for the samples where all the compounds were present. We analyzed the data distribution by employing the Shapiro–Wilk normality test implemented in the base R 4.1.0 package. Afterward, a Kruskal–Wallis test was done in the base R 4.1.0 package, followed by a check of the empirical cumulative distribution functions (ECDF) of the investigated groups using the latticeExtra package in R. Based on the findings, the Conover–Iman multiple comparisons test with Bonferroni correction was implemented using the DescTools package in R, taking into account the adjusted \(p\) values (\(p < 0.05\) was considered significant). Spearman’s rank correlation analysis was done in GraphPad Prism version 6.0 (\(p < 0.05\) was considered significant). The graphs were generated with the ggplot2 [78] package in R 4.1.0 and with GraphPad Prism version 6.0. The map (Figure 1) was generated using the QGIS 3.6 software [79] and the Natural Earth Data maps.
3. Results

Out of the nine investigated sites, sufficient Trichoptera and Ephemeroptera individuals could be sampled from only eight of them. Mirex and chlordanes were not found in any of the analyzed samples while HCB was found in only three Trichoptera samples (M11 with 0.63 ng g\(^{-1}\) dry weight, M12 with 0.89 ng g\(^{-1}\) dry weight, and M14 with 0.74 ng g\(^{-1}\) dry weight). The concentration of ΣPCB ranged from 1.27 to 15.04 ng g\(^{-1}\) dry weight for sediment samples, from 33.06 to 66.51 ng g\(^{-1}\) dry weight for Trichoptera samples, and from 23.99 to 68 ng g\(^{-1}\) dry weight for Ephemeroptera samples (Table 1). In the case of ΣHCH, the concentrations ranged from 0.19 to 40.02 ng g\(^{-1}\) dry weights for sediment samples, from 9.64 to 93.25 ng g\(^{-1}\) dry weight for Trichoptera samples, and from 59.65 to 239.39 ng g\(^{-1}\) dry weight for Ephemeroptera samples (Table 1). For ΣDDT, the concentrations ranged from 1.21 to 25.95 ng g\(^{-1}\) dry weight for sediment samples, from 26.73 to 57.17 ng g\(^{-1}\) dry weight for Trichoptera samples, and from 19.94 to 148.31 ng g\(^{-1}\) dry weight for Ephemeroptera samples (Table 1). We determined the frequency of detection of ΣPCB to be 31.3% in sediment samples, 50% in Trichoptera samples, and 62.5% in Ephemeroptera samples. For ΣHCH, the frequency of detection was 31.3% in sediment samples, 50% in Trichoptera samples, and 50% in Ephemeroptera samples. For ΣHCH, the frequency of detection was 31.3% in sediment samples, 75% in Trichoptera samples, and 100% in Ephemeroptera samples for ΣDDT (Table 1). For HCB, the frequency of detection was 37.5% for Trichoptera samples while the sediment and Ephemeroptera samples were not contaminated with HCB.

| Sample | Sediment | Trichoptera | Ephemeroptera |
|--------|----------|-------------|---------------|
| POPs   | ΣPCB     | ΣHCH        | ΣDDT          | ΣPCB     | ΣHCH        | ΣDDT          | ΣPCB     | ΣHCH        | ΣDDT          |
| Min (ng g\(^{-1}\) dry weight) | 1.27     | 0.19        | 1.21          | 33.06    | 9.64        | 26.73         | 23.99    | 59.65       | 19.94         |
| Max (ng g\(^{-1}\) dry weight) | 15.04    | 40.02       | 25.95         | 66.51    | 93.25       | 57.17         | 68.56    | 239.39      | 148.31        |
| Mean (ng g\(^{-1}\) dry weight) | 5.01     | 11.39       | 11.23         | 46.03    | 33.31       | 43.27         | 41.21    | 131.83      | 91.86         |
| Median (ng g\(^{-1}\) dry weight) | 2.20     | 2.34        | 10.29         | 42.27    | 15.18       | 47.71         | 35.17    | 114.14      | 114.86        |
| Number of occurrences | 5 | 5 | 5 | 4 | 4 | 6 | 5 | 4 | 8 |
| Frequency of detection * | 31.3% | 31.3% | 31.3% | 50.0% | 50.0% | 75.0% | 62.5% | 50.0% | 100.0% |

* Frequency of detection for POPs based on analyzed samples, not total number of sites.

The Spearman rank correlation coefficients between sediment and sampled benthos were the following: −0.5 (p > 0.9999) when comparing sediment to Trichoptera samples for ΣPCB, 0.5 (p > 0.9999) when comparing sediment to Trichoptera samples for ΣHCH and 1 (p = 0.333) when comparing sediment to Ephemeroptera samples for ΣHCH, and 0.5 (p > 0.9999) when comparing sediment to Trichoptera samples for ΣDDT and 0.9 (p = 0.083) when comparing sediment to Ephemeroptera samples for ΣDDT.

The p,p'-DDD/p,p'-DDE ratio was calculated for six of the nine sediment samples (with a minimum value of 0.377 and a maximum value of 2.672), for three of the eight Trichoptera samples (0.219, 0.174, and 0.244), and for two of the eight Ephemeroptera samples (0.465 and 0.526). The (p,p'-DDE + p,p'-DDD)/p,p'-DDT ratio was calculated for five out of the nine sediment samples (with a minimum of 0.195 and a maximum of 1.694) and for three of the Trichoptera samples (3.360, 5.0393, and 10.115), while the ratio for Ephemeroptera samples was calculated for just one of the sampling sites (10.575).

In the case of ΣDDT, the most frequent compound detected in all the sample types was p,p'-DDE (sediment 55.6%, Trichoptera 75%, and Ephemeroptera 100%). In the case of sediment samples, p,p'-DDE was followed by p,p'-DDD (44.4%), and then on equal footing by o,p'-DDD, o,p'-DDT, and p,p'-DDT (33.3%), while o,p'-DDE was not found in any sediment samples. For Trichoptera samples, p,p'-DDE was followed by all the other compounds in equal frequency: p,p'-DDD, o,p'-DDD, o,p'-DDT, p,p'-DDT, and o,p'-DDE (37.5%). In the case of the Ephemeroptera samples, p,p'-DDE was followed by p,p'-DDD, o,p'-DDD, and o,p'-DDT with an equal detection frequency of 25% and then by p,p'-DDT (12.5%), while o,p'-DDE was not quantified in any of the sample sites. The median concentration decreased as follows for sediment samples: p,p'-DDT > p,p'-DDD.
> p,p'-DDE > o,p'-DDD > o,p'-DDT > o,p'-DDE; in the following order for Trichoptera samples: p,p'-DDE > p,p'-DDT > p,p'-DDD > o,p'-DDT > o,p'-DDD > o,p'-DDE; and for Ephemeroptera samples in the following order: p,p'-DDE > p,p'-DDD > o,p'-DDD > o,p'-DDT > p,p'-DDT > o,p'-DDE.

For ΣPCB, the most frequent compound detected was PCB 28 in sediment and Trichoptera samples (sediment 33.3% and Trichoptera 50.0%) and PCB 66 in Ephemeroptera samples with 50.0% detection. In sediment samples, PCB 28 was followed by PCB31 and PCB66 (22.2%) and then equally by PCB 52, PCB 56, PCB 149, PCB 170, and PCB 180 (11.1%), while the rest of PCBs were not identified. For Trichoptera samples, PCB 28 was followed by PCB 31, PCB 47, PCB 52, PCB 56, PCB 66, PCB 74, PCB 99, PCB 101, PCB 110, PCB 118, PCB 138, PCB 149, PCB 151, PCB 153, and PCB 180 (37.5%) then by PCB 170 and PCB 187 (25%) and PCB 105, PCB 114, and PCB 141 (12.5%) while the rest of compounds were not detected. In the case of the Ephemeroptera samples, PCB 66 was followed by PCB 28, PCB 52, and PCB 74 (37.5%), by PCB 31 with 25%, and then by PCB 52 and PCB 101 with an equal detection frequency of 12.5% while the other PCBs were not quantified in any of the sample sites. The median concentration decreased in the following order for sediment samples: PCB 28 > PCB 31 > PCB 66 > PCB 56 > PCB 180 > PCB 52 > PCB 170; in the following order for Trichoptera samples: PCB 28 > PCB 66 > PCB 31 > PCB 56 > PCB 52 > PCB 56 > PCB 74 > PCB 153; and for Ephemeroptera samples in the following order: PCB 28 > PCB 31 > PCB 52 > PCB 66 > PCB 56 > PCB 74 > PCB 101.

In the case of ΣHCH, the most frequent isomers detected were β-HCH and γ-HCH in sediment samples with a detection frequency of 22.2% and α-HCH and β-HCH in Trichoptera and Ephemeroptera samples with an equal detection frequency of 50%. In the case of sediment samples, β-HCH and γ-HCH were followed by α-HCH (11.1%), while δ-HCH and ε-HCH were not identified in any sediment samples. For Trichoptera samples, α-HCH and β-HCH were followed by γ-HCH and δ-HCH with an equal detection frequency of 37.5% and then by ε-HCH (12.5%). In the case of Ephemeroptera samples, α-HCH and β-HCH were followed by γ-HCH and δ-HCH with an equal detection frequency of 25% and then by ε-HCH (12.5%). The median concentration decreased in the following order for sediment samples: α-HCH > β-HCH > γ-HCH; in the following order for Trichoptera samples: β-HCH > α-HCH > γ-HCH > δ-HCH > ε-HCH; and for Ephemeroptera samples in the following order: α-HCH > β-HCH > δ-HCH > γ-HCH > ε-HCH.

As can be seen in Figure 2, the concentration of ΣPCB, ΣHCH, and ΣDDT is higher in the invertebrate samples when it is detected than in the sediment samples in all the cases. For ΣPCB and ΣDDT, there are significantly higher concentrations detected in both Ephemeroptera and Trichoptera than in the sediment samples (p < 0.05). Meanwhile, for ΣHCH and ΣDDT, only the Ephemeroptera group has a significantly higher concentration than the sediment samples, with the Trichoptera group having no difference in concentrations of POPs with regards to the other two analyzed groups (Figure 3).

In order to determine if the samples we analyzed (sediment, Trichoptera, and Ephemeroptera) all share the same source of contamination, we determined the ratios of p,p'-DDD/p,p'-DDE [76] and (p,p'-DDE + p,p'-DDD)/p,p'-DDT [77] between sample types (Figure 4).
Figure 2. Concentration of \( \Sigma \text{PCB} \), \( \Sigma \text{HCH} \), and \( \Sigma \text{DDT} \) in sediment (○), Ephemeroptera (△), and Trichoptera (□) samples from the investigated sites.

Figure 3. Significant differences in concentration between sediment, Trichoptera, and Ephemeroptera. Box plots with whiskers at the minimum and maximum values while the box is delimited by the 25th and 75th percentile with the median shown as a line and the mean as a plus sign. Different letters above the boxes indicate significant differences (\( p < 0.05 \)).
4. Discussion

The detection of Trichoptera and Ephemeroptera individuals in only eight out of nine sampling sites is due to the different habitats and habitat qualities encountered during this research along the Mureș River. Some of the sites where sediment was sampled were not populated with individuals from the Trichoptera (M7 site, Figure 1) or Ephemeroptera (M14 site, Figure 1) orders or the populations were not developed enough for a representative sample size.

Mirex was not directly used in Romania which is why we did not find its presence in sediment or benthos samples, and we could argue that the trans-boundary contamination with mirex is low in Romania.

The Spearman rank correlation results come as no surprise for the situation as few sites are compared, and between these sites, only some have concentrations of POPs in all three sampling categories investigated (sediment, Trichoptera larvae, and Ephemeroptera larvae). The ratio analysis showed (Figure 4) that there are too few samples that are contaminated with all three compounds of interest that could be used in a reliable test. Contrary to these results, the frequency of detection of ΣDDT compounds shows a consistent decreasing trend from p,p'-DDE to p,p'-DDD that characterizes all of the sample types. In the case of ΣPCB, the decreasing trend starts with PCB28 for all the investigated sample types and is similar between them while a similar situation is brought forth from the analysis of ΣHCH isomers which shows that β-HCH is most commonly found with a higher frequency than the rest (when not of equal frequency). From these last observations, we can propose that the contamination of sediment, Trichoptera, and Ephemeroptera samples is consistent with a single source of pollution, which could very well be the sediment that is considered by many as a reservoir for POPs [24,27].
The similar frequency of detection for the different compounds of DDT between sample types is an indication that there is no metabolization of a specific compound to the detriment of others and that if there is metabolization, it probably happens at a proportional rate to what is happening in the sediment samples. Another possibility could be due to a combination of the latter with a different bioaccumulation pattern of the investigated POPs which could very well be the situation for ΣPCB and ΣHCH in which the frequency of detection varies. When comparing the frequency of detection with the median concentrations, we see that the pollution with DDT in sediment shows recent contamination (higher concentration of p,p’-DDT) while the median concentrations for benthos samples show contamination which is not so recent (higher concentration of p,p’-DDE). This could be due to the recent sediment contamination which the benthos samples have not had time to bioaccumulate. Another possibility is that the benthos invertebrates (or lower trophic levels) break down DDT and become the source for metabolites in higher trophic levels [80]. This furthers the notion that there is a need for biomonitoring groups and that the analysis of sediment is still necessary and complementary to benthos [32,57].

One reason that the Trichoptera samples do not have a significantly higher concentration than the sediment samples could be the fact that some Trichoptera larvae create small houses that protect them from the surrounding sediment which could inhibit the capacity of bioaccumulation through direct contact [81]. The fact that the concentration of POPs is higher in the Ephemeroptera samples than in the sediment samples (Figure 1) coupled with the higher frequency of detection (Table 1) in both the Trichoptera and Ephemeroptera samples makes the latter group a better candidate for further analysis in determining the environment’s POP contamination.

Our results support the idea of using Trichoptera and especially Ephemeroptera samples for biomonitoring purposes. Using benthos samples is usually a better decision than using other animals such as some fish because invertebrate larvae have restricted mobility [82] and are thus capable of better representing the study area. The use of invertebrate samples for biomonitoring has been proposed in the past for Leunereis culveri in estuarine environments [83] and for benthos samples separated in FFGs [50]; therefore, we consider that the benthos samples we analyzed in this study are suitable for determining the contamination of POPs in the lotic environment.

In completion of many classification systems of monitoring parameters that are used in European countries, but not only, and up-to-date more uniform and continual classification proposals [84], we suggest the possibility of use of the sediment and Ephemeroptera and Trichoptera complex in pre-alert assessing and monitoring of the presence and level of the highly toxic POPs (OCPs and PCBs) in aquatic ecosystems.

5. Conclusions

In this study, we determined that the benthos samples have higher concentrations of POPs (OCPs and PCBs) than sediment samples. This, along with the fact that the frequency of detection for POPs is similar between the sample types makes the sediment–Ephemeroptera–Trichoptera (SET) complex a suitable indicator of habitat contamination. The relatively short-medium-term gradient of pollution with POPs can be identified due to sediment and Ephemeroptera and Trichoptera assessment and monitoring, based on their short to medium to longer time of pollutant adsorption in sediments and bioaccumulation in Ephemeroptera and Trichoptera. This SET complex can be used to rapidly determine the appearance and longer time monitoring presence and concentration of POPs before they have time to bioaccumulate in edible organisms such as fish. Consequently, fish that live more than two years can be used in this respect as a much longer-term pollution indicator [36]. Therefore, the SET complex can constitute a pre-alert and alert group of indicators that reveal the presence and levels of POPs in aquatic systems before there is a threat to human food resources and to humans themselves.

Finally, it can be highlighted that the fact that the trophic niches created by these benthic organisms represent a compartment with an active functional-transfer role of POPs
between the abiotic and higher-level trophic compartments ending on the human plates, the insect larvae representing, due to their relatively high biomass and turn-over, a very important taxonomic group for POP circulation in nature, together with sediments at an initial point and fish at the end of such analysis.

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**Institutional Review Board Statement:** Ethical review and approval were waived for this study, since the biological material was formed by invertebrates. More than that these invertebrates are common, highly abundant in the environment, and of no national or international conservation value. Also, it must be added that no invertebrates were collected from protected areas.

**Informed Consent Statement:** This study did not involved humans.

**Data Availability Statement:** The data presented in this study are available on request from the corresponding author.

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