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9 **Environmental characteristics of a tundra river system in Svalbard. Part 2: chemical**  
10 **stress factors**

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23

24 **Abstract:** Bacterial communities in the Arctic environment are subject to multiple stress  
25 factors, including contaminants, although typically their concentrations are small. The Arctic  
26 contamination research has focused on persistent organic pollutants (POPs) because they are  
27 bioaccumulative, resistant to degradation and toxic for all organisms. Pollutants have entered  
28 the Arctic predominantly by atmospheric and oceanic long-range transport, and this was  
29 facilitated by their volatile or semi-volatile properties, while their chemical stability extended  
30 their lifetimes following emission. Chemicals present in the Arctic at detectable and  
31 quantifiable concentrations testify to their global impact. Chemical contamination may induce  
32 serious disorders in the integrity of polar ecosystems influencing the growth of bacterial

33 communities. In this study, the abundance and the types of bacteria in the Arctic freshwater  
34 were examined and the microbial characteristics were compared to the amount of potentially  
35 harmful chemical compounds in particular elements of the Arctic catchment. The highest  
36 concentrations of all determined PAHs were observed in two samples in the vicinity of the  
37 estuary both in June and September 2016 and were 1964 ng L<sup>-1</sup> (R12) and 3901 ng L<sup>-1</sup> (R13)  
38 in June, and 2179 ng L<sup>-1</sup> (R12) and 1349 ng L<sup>-1</sup> (R13) in September. Remarkable  
39 concentrations of the sum of phenols and formaldehyde were detected also at the outflow of  
40 the Revelva river into the sea (R12) and were 0.24 mg L<sup>-1</sup> in June and 0.35 mg L<sup>-1</sup> in  
41 September 2016. The elevated concentrations of chemical compounds near the estuary  
42 suggest a potential impact of the water from the lower tributaries (including the glacier-fed  
43 stream measured at R13) or the sea currents and the sea aerosol as pollutant sources. The  
44 POPs' degradation at low temperature is not well understood but bacteria capable to  
45 degrading such compounds were noted in each sampling point.

46 **Keywords:** Arctic, Freshwater contamination, POPs, Bacterial abundance, Bacterial diversity,  
47 Environmental changes

## 48 **1. Introduction**

49 The Arctic contains some highly productive ecosystems despite its extreme environmental  
50 conditions, strong seasonal changes in irradiance and snow cover, and the primary  
51 productivity concentrated in the short summer (Nguyen et al. 2015). Bacterial extremophiles  
52 are among the dominant life forms in the Arctic. They are able to survive in the harsh polar  
53 conditions and have developed mechanisms that allow them to cope with a variety of stress  
54 factors, e.g. temperature fluctuations, repeated freeze-thaw cycles, high or low levels of  
55 salinity or pH, UV light and desiccation (Sahay et al. 2013; Hoover and Pikuta 2013;  
56 Ntougias et al. 2016). These environmental stresses are yet enhanced by the increasing



57 concentrations of harmful chemical compounds, including persistent organic pollutants  
58 (POPs). There are a few local sources of contaminants in the Arctic, such as military  
59 installations, industrial outlets and waste from the old mines, settlements and ships, or the use  
60 of insecticides for insect control. However, the majority of Arctic pollution problem arises  
61 from a combination of long-range transport of pollutants and the Arctic haze phenomenon,  
62 locking the contaminated air in the area for months.

63 The concentrations of chemical compounds, including contaminants, differ in various aqueous  
64 reservoirs: lakes, river and tributaries (Kosek et al. 2018). Pollutants in the environment are  
65 exposed to degradative forces. Among them biotic degradation or metabolic processes are  
66 known to play a vital role in deciding overall fates of organic pollutants. They not only  
67 contribute to the disappearance of the original form of pollutants but also change their  
68 physicochemical properties and due to it, their transport and distribution behavior among  
69 various compartments in the environment. Physical and chemical factors may render a given  
70 contaminant more or less susceptible to bacterial degradation (Matsumura 1989). On the other  
71 hand, in aquatic environment, there are some bacterial communities incapable of degrading  
72 pollutants, and in such areas the concentration levels of pollutants increase remarkably (Ma et  
73 al. 2016; Nadal et al. 2015). Lakes in remote areas such as the Arctic have been of particular  
74 interest over the last decade for investigating the fate and dynamics of POPs (Evenset et al.  
75 2004; Evenset et al. 2007; Ahrens et al. 2016). Increasing trends in contamination levels  
76 suggest that these areas are significant trapping sites of persistent toxic pollutants (Jiao et al.  
77 2009). Due to the low temperature in the Arctic, mineralization of POPs is extremely slow in  
78 cold habitats and they likely bioaccumulate in the adipose tissue and then biomagnify in  
79 species inhabiting the polar regions (Kosek et al. 2007). Bacteria inhabiting the Arctic, are  
80 more strained and susceptible to the adverse effects of POPs than the bacteria living in other  
81 regions due to their long life and slow detoxifying. Moreover, in the nutrient-limited

82 environments, aromatic compounds may serve as a carbon source, also under sulfate-reducing  
83 and nitrate-reducing conditions. However, very little is known about their anaerobic  
84 degradation pathways (Foght 2008; Mallick et al. 2011), particularly in polar regions. Low  
85 temperature catabolic genes/enzymes activity is of a great interest due to the their  
86 biotechnological applications.

87 The main purpose of this article was to study the interactions between the pollutants and  
88 bacterial abundance. Selected xenobiotics, such as polycyclic aromatic hydrocarbons (PAHs),  
89 phenolic compounds and formaldehyde, and several potentially toxic metals, were determined  
90 in the Revelva catchment, as were the bacterial volume and the total number of bacteria. In  
91 the selected samples from this river system, metagenomic research was conducted to examine  
92 the bacterial community composition and its adaptation to this environment.

## 93 **2. Materials and Methods**

### 94 2.1. Fieldwork

95 The study was conducted in the Revelva catchment (Wedel-Jarlsberg Land, southwestern  
96 Spitsbergen, near the Polish Polar Station Hornsund). A detailed map of the sampling area is  
97 shown in Part 1 of this article (Kosek et al. submitted) and our former work (Kosek et al.  
98 2018). In brief, the samples were taken from 14 locations in the river, its tributaries and lakes  
99 through which it flows, from mountain streams filling rocky beds to its estuary at the  
100 Hornsund fjord bay Ariebukta (Table 1). Among the tributaries, the largest one was fed by  
101 glacier melt (Ariebekken). Each place was sampled twice, in June and September 2016,  
102 reflecting a shift from melting snow patches to permafrost thaw and rainwater as main water  
103 sources over the summer season. Furthermore, three points (R4, R8 and R14) were checked  
104 for the bacterial taxonomy. The tested points differed remarkably in terms of geological  
105 substratum, vegetation and water flow velocity, which influence chemical constituent sources

106 and the potential of self-cleaning for these environments. Points R4 and R8 were located in  
107 the areas of no or limited biological soil crust, while the point R14 was surrounded by boggy  
108 vegetation, composed of a mixture of mats formed by cyanobacteria and bryophytes, as well  
109 as by small lichens and saxifrages in varying proportions (Kumar et al. 2017). Additionally, it  
110 should be noted that water was flowing in the points R8 and R14 (most rapidly in R8), while  
111 in the point R4 (lake) it was relatively stagnant. Separate aliquots were prepared for chemical  
112 composition analysis (in pre-cleaned 1 L HDPE bottles, stored at 4°C), microbiological  
113 parameters quantification (50 mL, preserved with 2% formaldehyde, stored at 4°C) and  
114 metagenomics (1.5 L, stored frozen).

115 **Table 1.** Location of the sampling points in the Revelva catchment in Svalbard.

## 116 2.2. Chemical Analysis

117 The concentrations of PAHs were determined in freshwater samples using Gas  
118 Chromatography coupled with Mass Spectrometry Technique, while formaldehyde and the  
119 sum of phenols have been determined using Spectrophotometry Method. Trace elements have  
120 been determined with Inductively Coupled Plasma Mass Spectrometry. Further technical  
121 specifications of the analytical equipment and method, including basic validation parameters  
122 of the analytical procedures, are given in Table 2. All blanks were prepared with Milli-Q  
123 deionised water. A further chemical description of these samples (inorganic ions, electrical  
124 conductivity) can be found in Part 1 of this article (Kosek et al. submitted).

125 **Table 2.** Validation parameters and technical specifications used in the applied analytical  
126 procedures.

## 127 2.3. Quality assurance / Quality Control (QA/QC)

128 The analytical procedures used to determine individual components in the studied samples  
129 have been validated against certified reference materials (CRMs) concordant with ISO Guide  
130 34:2009 and ISO/IEC 17025:2005. The data obtained here were subject to strict QC  
131 procedures. The analysis of trace elements involved the application of Standard Reference  
132 Material (RM) NIST 1643e Trace Elements in Water, and RM Enviro MAT ES-L-2CRM,  
133 ES-H-2 CRM SCP SCIENCE. The calibration of the apparatus was based on RMs by  
134 Inorganic ventures ANALITYK: CCS-4, CCS-6, CCS-1, IV-ICPMS-71A. The sensitivity of  
135 the applied methods was tested by injecting standard mixtures of the analytes in the measured  
136 concentration range. Linear calibration curves of the peak area against standard concentration  
137 showed correlation coefficients ( $R^2$ ) in the range of 0.898–0.999 for all standards.  
138 Technically, each sample was analysed in triplicate. The instrumental background was  
139 checked by inserting Milli-Q water blanks once per every six samples. All the obtained values  
140 for organic compounds (PAHs) in CRMs were within the confidence interval. Reproducibility  
141 and recovery for both groups of organic compounds were high (85%–105%) with relative  
142 standard deviation (RSD) 4%–10%. Finally, the measurements of formaldehyde and the sum  
143 of phenols have been done in accordance with norms ISO 8466-1 and DIN 38402 A51,  
144 respectively.

#### 145 2.4. Bacterial Abundance Analysis

146 The microbial community parameters quantification has been thoroughly described in Part 1  
147 of this article (Kosek et al. submitted). Briefly, three parameters: total bacterial number,  
148 average bacterial cell volume and bacterial biomass (a product of the former two parameters),  
149 were quantified in the 28 water samples (14 samples collected in June and 14 samples  
150 collected in September). The method applied was epifluorescence microscopy, with DAPI  
151 stain, on filters with a pore diameter of 0.2  $\mu\text{m}$ . We used a Nikon Microscope 80i with NIS-

152 Elements BR 3.0, a MultiScan automated image analysis system, and a high resolution color  
153 digital camera (Nikon DS-5Mc-U2).

## 154 2.5. Bacterial Community Structure Analysis

155 In this study, we use the data obtained in Part 1 of this study (Kosek et al. submitted) to  
156 analyse another aspect of an Arctic tundra river system: the impact of chemical stress factors  
157 on abundance and bacterial community. As a background, we briefly describe the type of data  
158 used here and the methods applied in their acquisition. The following paragraph concerns 6  
159 samples in total, collected in points R-4, R-8 and R-14 in June and September. The data was  
160 then used with a special focus on bacterial genera which could decompose pollutants,  
161 especially from the PAHs group.

162 The bacterial community structure, i.e. percentage division into main phyla and the smaller  
163 taxonomic units (including genus, or even species level, if possible to determine  
164 unequivocally), was analysed using next generation sequencing (NGS) technology. This was  
165 conducted in 0.2-µm filter residue, from which microbial DNA was isolated and analysed  
166 using 16S microbial sequencing on a MiSeq platform (Illumina). Prior to this procedure, the  
167 DNA concentration was determined with an ND-1000 UV-Vis spectrophotometer. On the  
168 obtained DNA samples, PCR (polymerase chain reaction) was conducted using Q5 Hot Start  
169 High Fidelity 2X Master Mix (New England Biolabs), following amplification with the  
170 primers: 341F – CCTACGGGNGGCWGCAG and 785R –  
171 GACTACHVGGGTATCTAATCC. The results were processed using a set of bioinformatics  
172 tools (see Part 1, Kosek et al. submitted). The affinity of the bacterial communities found in  
173 the analysed samples was explored with cluster analysis, and these were used for the  
174 estimation of biodiversity indices.

## 175 3. Results and Discussion



### 176 3.1. Chemical stress factors occurring in the studied freshwater samples

#### 177 3.1.1. pH

178 In freshwater environments, pH has been shown to be a decisive environmental factor  
179 determining the bacterial community composition, often being the most important one  
180 compared to factors such as temperature, organic matter, water retention time, and nutrient  
181 concentrations (Lindström et al. 2005). pH is also an environmental factor that can vary  
182 greatly in aquatic ecosystems (Bååth and Kritzberg 2015). Lake, river and stream waters can  
183 have pH values below 4 and above 9 even within small geographical areas. In highly  
184 productive lakes, pH at the surface may be 2 units higher than in bottom waters. The variation  
185 of the values is driven by vertical differences in photosynthesis, respiration, and redox  
186 conditions (Wetzel 2001). pH can also fluctuate rapidly. For example, during snow melt and  
187 rain storms, pH values in streams can decrease several units, sometimes within a few hours  
188 (Lawrence 2002). On the other hand, sunny days can result in high photosynthetic activity  
189 with the increase of water pH values. Accordingly, changes of 2-3 pH units may be found in  
190 highly productive aquatic environments (Tank et al. 2009). During episodes of rapid pH  
191 changes, the bacterial community may not be optimally adapted to the new pH condition,  
192 resulting in impaired functions, and sometimes, inhibition of bacterial growth (Bååth and  
193 Kritzberg 2015). Freshwater pH values, and in particular their changes, may pose a big threat  
194 to bacterial development and play a key role as a stress factor. However, in our study, the pH  
195 values in the collected samples were differing only slightly and ranged from 7.0 to 8.0 both in  
196 June and September 2016 (Figure 1). Former hydrochemical studies of the Hornsund fjord  
197 area (including Revelva catchment) show high hydrochemical variability, with some values  
198 within similar ranges as described in Part 1 of this article (Kosek et al. submitted). Small pH  
199 values variation can be explained by the ability of bacteria to regulate the pH of water.  
200 Consequently, bacteria are able to survive and develop even in the most harsh conditions. The

201 interaction between the microbes may be set by how their metabolism change the  
202 environment and react to those changes. Furthermore, many biochemical reactions involve a  
203 turnover of protons and bacteria also alter the pH around them. When the pH modification is  
204 beneficial for the bacteria, there is a positive feedback on their growth. The more bacteria  
205 there are in the water, the stronger they can change the environment. At adverse pH  
206 conditions, a sufficiently high cell density may therefore be needed to survive at all (Ratzke  
207 and Gore 2018).

208 **Figure 1.** The pH values detected in freshwater samples collected in June and September  
209 2016.

### 210 3.1.2. Trace elements

211 Trace elements were also determined in the samples from both June and September 2016. The  
212 concentrations of the following trace elements were determined in them: Li, Be, Al, V, Cr,  
213 Mn, Co, Ni, Cu, Zn, Ga, As, Rb, Sr, Ba, Tl and U (Table 3). The CVs of the obtained results  
214 ranged from 0.5 to 1.5 %.

215 **Table 3.** Concentrations ( $\pm$ standard deviation, SD) of trace elements in the collected  
216 freshwater samples.

217 The Revelva catchment waters are enriched in trace elements due to the presence of ore-  
218 bearing veins and metamorphic rocks in the area (Wojciechowski 1964; Smulikowski 1965).  
219 This geological substratum is more exposed in the upper parts of the catchment. Furthermore,  
220 the spatial variability of the underlying rocks in this catchment allows for the more abundant  
221 occurrence of titanium, possibly also barium, caesium, lithium, rubidium, and zinc in the  
222 upper part of the catchment; of zirconium in the left tributaries of the middle part, and of  
223 chromium and vanadium in both these areas. The local rocks are relatively abundant in  
224 aluminum and manganese throughout the catchment (Smulikowski 1965). As for the ore-

225 bearing veins, in the area occur those with chalcopyrite, cuprite, malachite and azurite, which  
226 are copper minerals, as well as smaller concentrations of sphalerite (with zinc) and galena  
227 (with lead). The specific locations of these veins favour the occurrence of copper near  
228 Skoddefjellet mountain, in the Arie glacier valley, and in the left tributaries of the biggest and  
229 smallest lakes in the valley (the top nameless lake and Revvatnet), the occurrence of lead in  
230 the Arie glacier valley and of both lead and zinc in the left tributaries of the smallest lake  
231 (Wojciechowski, 1964).

232 The trace metal concentrations detected at the two sampling occasions markedly differed from  
233 each other, with an increase in September. This may be caused by the occurrence of  
234 groundwater associated with the active layer of permafrost, which gains more importance in  
235 the hydrological regime of Revelva as snow patches disappear in the catchment. Apart from  
236 the local natural occurrence of trace elements, they are assumed to be derived to the Arctic  
237 mostly from long-range atmospheric transport (AMAP 2009), and these may be additionally  
238 supplied by September rainfalls. The increase in concentration of trace elements in September  
239 2016 was most evident in the central part of the lake shore and in two points located near the  
240 river estuary. The water stagnating in the lake experiences longer contact with suspended  
241 mineral matter, hence probably the higher trace element concentrations there. A similarly  
242 longer time may have contributed to the higher concentrations near the river mouth.  
243 Differences in individual trace element concentrations can be explained qualitatively in terms  
244 of mineral surface reactions, complexation, chemical weathering and sorption to solid-phase  
245 soil organic matter (Colombo et al. 2018), yet the detailed extent of these processes cannot be  
246 determined with the limited data we obtained and it is outside the scope of the current paper.

### 247 3.1.3. Organic compounds

248 In the collected samples, we have determined polycyclic aromatic hydrocarbons. Their  
249 concentration levels, as well as those of formaldehyde and the sum of phenols, are reported in  
250 Table 4.

251 **Table 4.** Concentrations ( $\pm$ standard deviation, SD) of PAHs, formaldehyde and the sum of  
252 phenols in the collected freshwater samples.

253 PAHs are a group of environmentally persistent organic compounds of varied toxicity, usually  
254 formed during the incomplete combustion of fossil fuels, biomass, and through other  
255 industrial activities. They have been found in the Arctic environment, originating both from  
256 the long-range atmospheric transport (Wang et al. 2013) and the local sources. Both human  
257 activity and natural phenomena (forest fires, volcanic eruptions) can produce them. PAHs  
258 have been found widely in polar environmental media: the atmosphere, water, sediments and  
259 biota (Polkowska et al. 2011; Kozak et al. 2017). They can be deposited and accumulated in  
260 ice for a long period of time and released to the environment when temperature exceeds the  
261 melting point (Ge et al. 2016). The results of PAHs analysis are shown in Table 4. The  
262 highest concentrations of PAHs have been detected in the sampling point R13 in June and in  
263 the sampling point R8 in September, and these were  $1871\pm 59$  ng L<sup>-1</sup> (ANT) and  $991\pm 42$   
264 ng L<sup>-1</sup> (FLA), respectively. Comparing the results of PAHs concentrations determined in  
265 summer 2016 with those collected and determined in summer 2015, it can be seen that the  
266 highest concentrations were observed in the same sampling points (Kosek et al. 2018).  
267 Slightly higher concentrations of PAHs were observed in the samples collected in 2016, but  
268 the differences are not statistically significant (Kruskal-Wallis ANOVA, all p levels for PAH  
269 congeners were above 0.13).

270 The sampling point R13 is located at the outflow from the Arie glacier, thus such a high  
271 concentration of PAHs observed in June can be explained by releasing pollutants from the

272 melting snow cover of the glacier. The difference between PAHs composition in June and  
273 September in the catchment (Figure 2) reflects well the order of PAHs elution from melting  
274 snowpack and the preferential storage of the more hydrophobic PAHs in ice (Kozioł et al.  
275 2017).

276 **Figure 2.** A box - whisker plot of the mean PAHs concentrations in water samples collected  
277 in June and September. Significant differences between seasons are indicated with the non-  
278 parametric Kruskal-Wallis ANOVA p-levels below 0.05 and given in boldface. The box  
279 encompasses the mean value  $\pm$  SD, the whiskers show the full range of values noted (where  
280  $<$ LOD values were assigned a half of the LOD level).

### 281 3.2. Microbial community

282 Various toxic elements and compounds, depending on their concentration in the environment  
283 and simultaneous effects of their occurrence, may or may not be an effective inhibitor of  
284 bacterial growth in aquatic environments. In Part 1 of this article (Kosek et al. submitted) we  
285 report bacterial abundance indices (total bacterial number, average cell volume and bacterial  
286 biomass) in the collected samples. Both spatially and temporally, these indices were  
287 characterised by a pronounced variability.

288 For this reason, in this study in June and September 2016, three points (R4, R8 and R14) were  
289 checked for the bacterial taxonomy. The general structure of studied microbial communities  
290 (based on the relative abundance of classified sequences) was composed mainly of bacterial  
291 taxa, with 43-53% *Proteobacteria*, 9-23% *Actinobacteria*, 6-12% *Bacteroidetes*, and more  
292 than 2% of *Planctomycetes*, *Firmicutes* and *Verrucomicrobia* in all samples (Part 1, Kosek et  
293 al. submitted). Interestingly, *Proteobacteria*, *Bacteroidetes*, *Firmicutes*, and *Actinobacteria*  
294 were also identified as the core phyla in activated sludge of municipal and industrial  
295 wastewater treatment system (Ibarbalz et al. 2013), which are typically polluted waters.

296 Indeed, some of the bacterial strains found in the sampled waters may be capable of  
297 decomposing specific pollutants. The specific allochthonous organic compounds of potential  
298 toxicity may modify the taxonomical structure of the microbial community and lead to the  
299 selection of bacteria that are capable of metabolizing them. For example, we have detected  
300 bacteria from *Flavobacteriaceae* (Bacteroidetes phylum) family, which are linked to the  
301 degradation of PAHs at low temperature (Eriksson et al. 2003). However, in the  
302 environmental niches, to obtain complete degradation of organic compounds, usually the  
303 complex bacterial community is involved. Therefore, in future studies it would be valuable to  
304 combine the data obtained from NGS with the analyses of specific functional genes involved  
305 in particular PAHs degradation.

306 Up to now numerous unique metabolic pathways of PAHs biodegradation have been already  
307 documented (Peng et al. 2008; Mallick et al. 2011; Ghosal et al. 2016), but in this terms the  
308 knowledge on the bacterioplankton community inhabiting the inland water system of the polar  
309 area is limited. Additionally, many of the reported data were from incubation of single or  
310 mixed cultures in laboratory experiment, while for in situ consortia and the observed for them  
311 degradation potential may differ, due to the combined activities of whole community  
312 members.

313 Major PAHs degrade approaches are highly linked to the oxygen presence/absence. Under  
314 aerobic conditions the oxygen is both the final electron acceptor and co-substrate to activate  
315 and subsequently cleave the aromatic ring, catalyzed by oxygenase enzymes (monooxygenase  
316 or dioxygenase) (Foght 2008; Carmona et al. 2009). In the anoxic conditions, which is  
317 regarded as more common in natural environments (e.g. aquifers, aquatic sediments and  
318 submerged soils), the attack on the aromatic ring is primarily based on reductive reactions  
319 (Foght 2008; Carmona et al. 2009), where nitrate, sulfate or ferric ions are used as final  
320 electron acceptors (Foght 2008; Carmona et al. 2009). For instance, *Hyphomonas* detected in

321 each sampling point in this study (see Table 5), showed the degradation potential after the  
322 addition of sources of nitrogen and phosphate to hydrocarbon-contaminated harbour  
323 sediments (Yakimov et al. 2005). The metabolic pathways of sulfate-reducing and Fe(III)-  
324 reducing bacteria are also of interest due to the role they play in the biogeochemistry  
325 including degradation of organic contaminants (Meckenstock et al. 2000). The sulfate-  
326 reducing *Desulfovibrio*-like bacteria, forming up to 0.26% of the bacterial community  
327 composition in the studied samples, as well as the spore-forming *Desulfotomaculum* and  
328 *Desulfosporosinus* genera from *Firmicutes* phylum (forming up to 0.22 % of the bacterial  
329 community), are also potentially capable of decomposing the organic pollutants occurring in  
330 this catchment, since these bacterial groups may use a variety of aliphatic and aromatic  
331 compounds as a carbon source (Hansen 1994). Interesting is also high relative abundance of  
332 psychrotolerant *Rhodoferrax* genus (from 1.56% to 5.6%), already reported as potential  
333 phenanthrene degrader (Martin et al., 2012), and in this study represented mainly by *R.*  
334 *ferrireducens* (>85%), which is capable of dissimilatory Fe(III) reduction at low temperature.

335 Additionally low molecular weight (LMW) PAHs, as naphthalene, anthracene and  
336 phenanthrene, more volatile and soluble in water are also more susceptible to biodegradation  
337 than high molecular weight (HMW) PAHs, thus also select for different microbial consortia  
338 (Vila et al. 2010). Some reports indicated the catabolic versatility of some bacteria (e.g.  
339 *Pseudomonadales* and *Sphingomonadales* from *Gammaproteobacteria* and  
340 *Alphaproteobacteria* classes, respectively) is linked to the presence of plasmid-encoded  
341 aromatic degradative genes (Peng et al. 2008), which can be disseminated by horizontal gene  
342 transfer to phylogenetically diverse bacteria (Nojiri et al. 2004). The list of members of  
343 bacterial community detected in studied sampling points and possibly involved in the  
344 pollutants degradation is given in Table 5.

345 **Table 5.** The list of members of bacterial community possibly involved in the degradation of  
346 pollutants. A genus was considered only if it constituted  $\geq 0.01\%$  of the community from a  
347 single sample.

348 Expanding the information on the variety of bacterial phyla and metabolic pathways presented  
349 in Part 1 of this article (Kosek et al. submitted), we conclude that various organic compounds  
350 (including those considered pollutants) may be also decomposed by them. This is consistent  
351 with the detection of pollutant-decomposing bacteria in such remote parts of the Arctic as the  
352 surface of the Greenland Ice Sheet (Hauptmann et al. 2017).

### 353 3.3. Statistical analysis of bacterial abundance and chemical background in the Revelva 354 catchment

355 A Principal Component Analysis (PCA) was performed to encompass the wider set of  
356 chemical variables connected to potentially toxicity (PAHs, HCHO, phenols and trace  
357 elements such as Ni, Zn, Cu, Co, Be, As, Mn, as well as pH, the extreme values of which are  
358 also a sign of hospitable environments). The above-mentioned trace elements were chosen  
359 based on the literature review by (Kabata-Pendias and Pendias 2001), on the basis of their  
360 highest potential for toxicity in the general biosphere (especially for plants). The quantitative  
361 data on the bacterial community, such as the total bacterial number (TBN) and the average  
362 volume of bacterial cells (ACV) described in details in Part 1 of this article (Kosek et al.  
363 submitted), were included in the analysis. For brevity, each PAH is referred to by an  
364 abbreviation listed in Table 2 of this manuscript. The PCA for this study was performed  
365 (Figure 3, 4) using R v. 3.4.4, using the *prcomp* function, on a log-transformed dataset, except  
366 the pH value which is a logarithm.

367 In the analysis conducted for the whole summer season, the scree plot shape suggested that  
368 the PCs 1, 2 and 3 were likely significant factors. We also conducted further analyses for June





369 and September separately, showing only the first two PCs in each (also as suggested by the  
370 scree plot)

371 **Figure 3.** PCA conducted for the potentially toxic chemicals and indices of the bacterial  
372 community structure detected in the hydrological environment of the Revelva catchment.  
373 Top: The space defined by the PCs 1 and 2, with a division by the month of sampling  
374 (colours). Bottom: Same analysis, space defined by the PCs 2 and 3, division by the type of  
375 the hydrological environment. Numbers 1-14 depict samples from locations R1-14 in June,  
376 numbers 15-28 denotes samples from the same locations in September, in the same order (e.g.  
377 number 28 is the sample from point R14 in September).

378 A clear division between sampling times was found to be depicted by the two main variability  
379 components (PC 1 and 2; Figure 3 Top), which accounted for approximately 34% of the total  
380 variability in the dataset. In the beginning of the summer season, elevated concentrations of  
381 NAP, ACE, ACY (lower molecular weight PAHs, also more water soluble), HCHO and Zn  
382 were noted, as well as a higher ACV bacterial community index. These factors may be linked  
383 to the still melting snow patches in the catchment (water-soluble PAHs elute from snowpack  
384 earlier, while the other PAHs may be stored as particle-bound and even incorporated into ice  
385 by refreezing – Meyer et al. 2009; Koziół et al. 2017). HCHO may even be produced in  
386 snowpack by photochemical reactions of more complex organic compounds, including those  
387 occurring in particulate forms (Sumner and Shepson 1999; Grannas et al. 2004). Snow is a  
388 medium poor in nutrients, hence the specialised k-strategist bacteria may predominate there,  
389 growing but not multiplying rapidly (compare Part 1, Kosek et al. submitted).

390 On the other hand, the late season was characterised by a higher concentration of most other  
391 toxic elements and compounds and the TBN index, which shows that the r-strategists were not  
392 limited by these toxic chemicals in their reproduction. This can be interpreted in terms of the

393 concentrations being too low to have a limiting impact on the bacterial community.  
394 Furthermore, some of the detected trace elements (such as Cu, Mn and Zn) have ample local  
395 sources in the geological substratum and therefore the local bacterial community is well  
396 adjusted to their presence. It is interesting to notice, however, that the most abundant presence  
397 of zinc deviates from its naturally enriched area in the upper part of the catchment (with  
398 higher concentrations in some of the samples located in the lower parts of the catchment,  
399 especially points R10 and R13), which may suggest its pollution origin. It is also the only  
400 trace metal in this analysis to correlate closer with ACV than TBN.

401 Multiple further trace elements, as well as phenols and higher molecular weight PAHs, tend to  
402 increase in concentration over the course of the summer season, which may be linked to the  
403 shift in the hydrological regime from snow-fed to permafrost thaw and occasional rainfall  
404 (Pulina et al. 1984). Among the trace elements analysed here, As, Co, Cu, Mn, Ni and Zn  
405 have been found by Kozak et al. (2015) in the local rainfall composition, likely representing  
406 both the local and distant sources, including rock dust, sea spray and human-activity-related  
407 emissions. Especially the Zn concentrations in rainfall may be very high in this region (mean  
408 concentration in Kozak et al's study reaching  $28.99 \mu\text{gL}^{-1}$ ), and hence it can be treated as a  
409 pollutant. Phenols may originate from local plant tissue decomposition (Grannas et al. 2004),  
410 which agrees well with their high concentrations in the samples from the lower part of the  
411 catchment, where lush tundra vegetation grows. The high-molecular-weight PAHs distinguish  
412 the upper part of the catchment, where they could have been stored longer from the snowpack  
413 sources, e.g. frozen in the ground ice.

414 The type of hydrological environment was best distinguished on the PC 2 and 3 graph (Figure  
415 3. Bottom), depicting approximately 27% of the total variability in the dataset. The river  
416 samples may have been grouped due to their location in the lower part of the catchment rather  
417 than by their difference from the smaller streams, since they are distinguished by the presence

418 of the earlier mentioned phenols, which may come from tundra vegetation, and Cu, Ni and  
419 Co, of which at least Cu should occur abundantly in the left tributaries of the lower part of the  
420 catchment. The lake samples clustered around such characteristics as higher molecular weight  
421 PAHs, some trace metals, HCHO and ACV, which indicates that in certain circumstances the  
422 stagnant water may gain more toxic characteristics and be less habitable to the generalist  
423 bacteria population, although the effect is not consistent across all samples.

424 **Figure 4.** PCA conducted for the potentially toxic chemicals detected in the different  
425 hydrological environments of the Revelva catchment (denoted by colour coding) and indices  
426 of the bacterial community structure in these waters. Each graph for a separate sampling  
427 occasion, the space defined by the PCs 1 and 2. Top: June. Bottom: September. Numbers 1-14  
428 depict samples from locations R1-14 in that particular month.

429 In June, a similar separate graph was prepared, concentrating on 47% of the total data  
430 variability depicted by PCs 1 and 2 (Figure 4. Top). The first PC, explaining almost 32% of  
431 the variability, differentiated strongly between the samples with the high and the low  
432 concentrations of PAHs, highlighting especially their elevated concentration in the glacier-fed  
433 stream at R13. It also maintained the clear division between environments with the high ACV  
434 (characteristic for k-strategists) and high TBN (r-strategy indicator; compare Part 1, Kosek et  
435 al. submitted). TBN was correlated relatively closely (and positively) with PAHs  
436 concentrations in June, hence probably this type of POPs was not counteracting the  
437 multiplication of bacteria, and perhaps these compounds could even be used as an organic  
438 substrate by some of the organisms present there. However, in September (Figure 4. Bottom),  
439 the concentrations of selected PAHs (ANT, FL) and formaldehyde showed a close affinity to  
440 ACV, as did Be and As. These compounds may have been limiting factors to bacterial  
441 multiplication in the samples taken in the upper part of the catchment then. However, TBN  
442 remained in a positive correlation with NAP and phenols.

#### 443 **4. Final remarks and conclusions**

444 The Arctic environment, although it seems to be free from anthropogenic pollution, is not as  
445 pristine as it might seem. Remarkable concentration levels of persistent organic pollutants  
446 have been detected in the freshwater samples collected from the Revelva catchment. Globally  
447 emitted contaminants accumulate in the Arctic and can be stored in this cold environment for  
448 a long period of time (Mackay and Wania, 1995; Friedman and Selin, 2016). Moreover,  
449 climate change influences the release of these contaminants through elevated melt rates,  
450 resulting in increased contamination locally (Blais et al. 2001; Miner et al. 2018). The  
451 microbial community interacts with contamination in the Arctic (e.g. Hauptmann et al. 2017),  
452 however it is yet unknown how universal such interactions are. The important issue is to  
453 know whether contaminants present in the environment are a toxic factor for bacteria, or  
454 whether they show the ability to deal with these pollutants and reproduce in spite of their  
455 presence. Described research shows that the catchment chosen for this study constitutes a  
456 place of accumulation of persistent organic pollutants and also some trace elements that may  
457 be toxic for the bacteria. The determined pollutants may pose a serious threat to the  
458 development of bacteria in the Revelva catchment. Depending on their concentration in the  
459 environment and simultaneous effects of their occurrence, they may or may not be an  
460 effective inhibitor of bacterial growth in aquatic environments. However, despite the presence  
461 of contaminants and the limited nutrient supply (described in Part 1, Kosek et al. submitted),  
462 the Revelva catchment is characterised by a great biodiversity. The general structure of these  
463 microbial communities was composed mainly of bacterial taxa, such as *Proteobacteria*,  
464 *Actinobacteria*, *Bacteroidetes*, *Planctomycetes*, *Firmicutes* and *Verrucomicrobia*. The  
465 bacterial ability to degrade toxic compounds depends on numerous factors, which were not  
466 studied in this research, but there is a possibility that the bacteria present there decompose the  
467 described contaminants. For example, *Bacteroidetes* are linked to the degradation of PAHs at

468 low temperature. Denitrifiers, as relatively abundant in this study *Flavobacterium*, sulphur  
469 (*Desulfovibrio*, *Desulfotomaculum*, *Desulfosporosinus*) or psychrotolerant Fe-reducing  
470 bacteria (*Rhodoferrax*) are also potentially capable of decomposing of the persistent organic  
471 pollutants occurring in this catchment (Martin et al. 2012; Kappell et al. 2014). The potential  
472 for biodegradation of polycyclic aromatic hydrocarbons (PAHs) at low temperature is not  
473 well understood, but such biodegradation would be very useful for remediation of polluted  
474 sites. Bacteria inhabiting the Revelva catchment have adapted to live in difficult conditions. It  
475 can be hypothesised that they show the potential for decomposing persistent organic  
476 pollutants, but in order to confirm it, it is necessary to conduct more thorough research at the  
477 molecular level.

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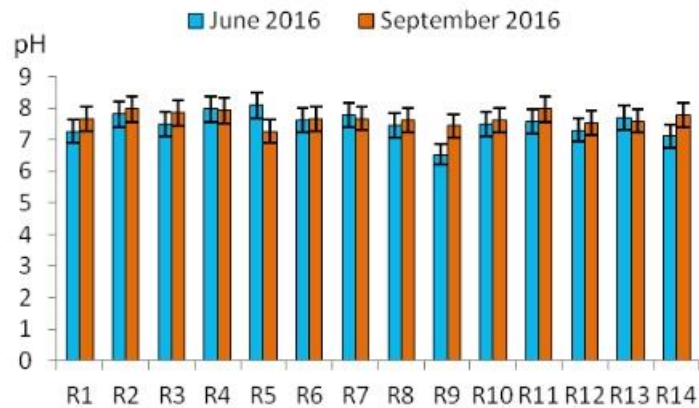
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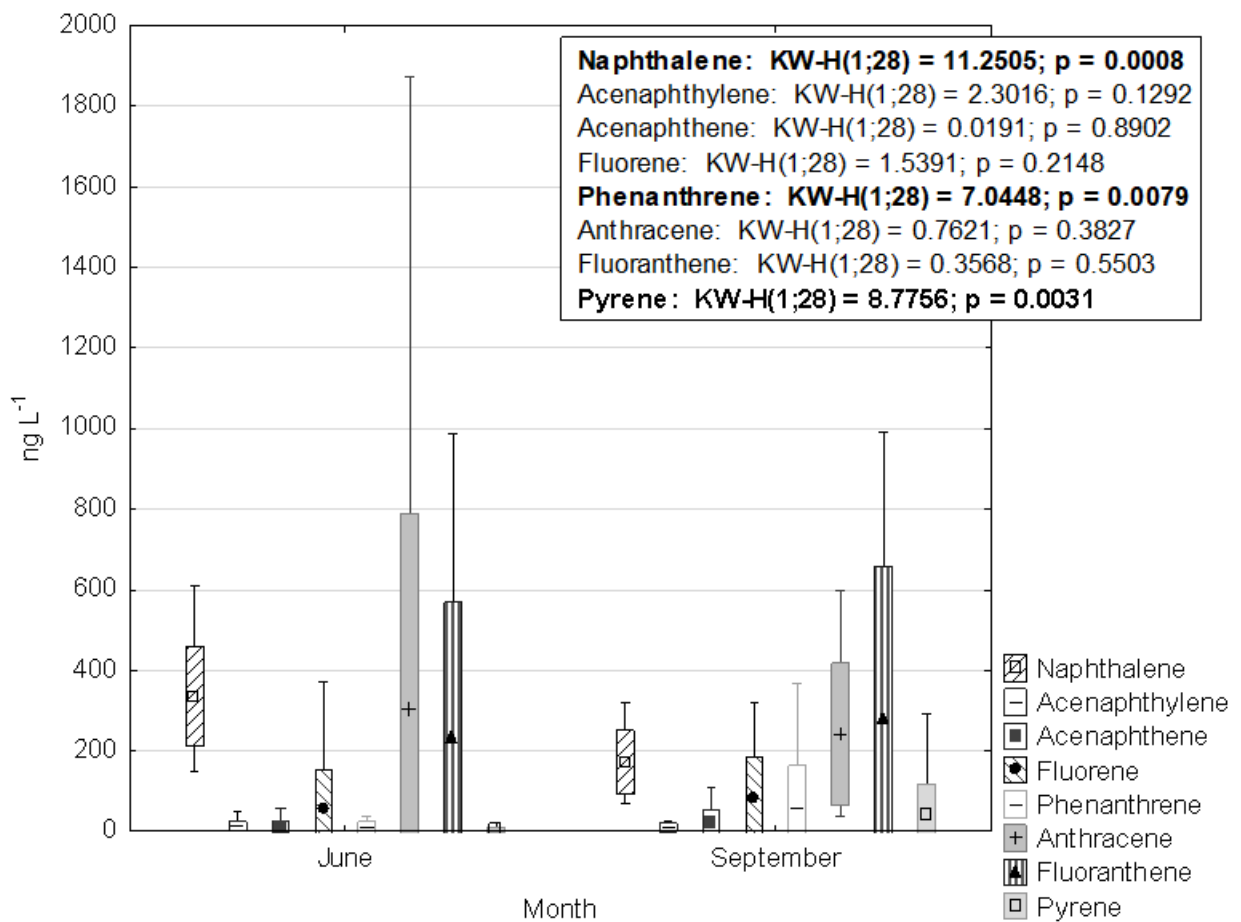
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868 **Figure 1.** The pH values detected in freshwater samples collected in June and September

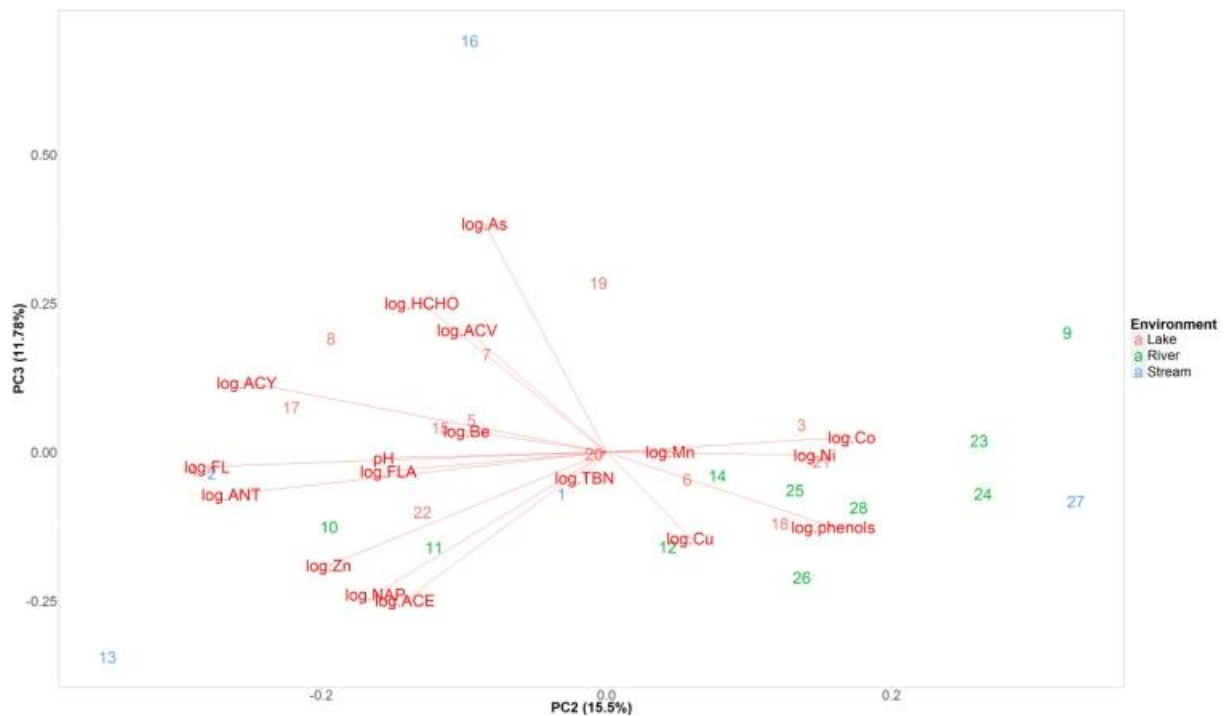
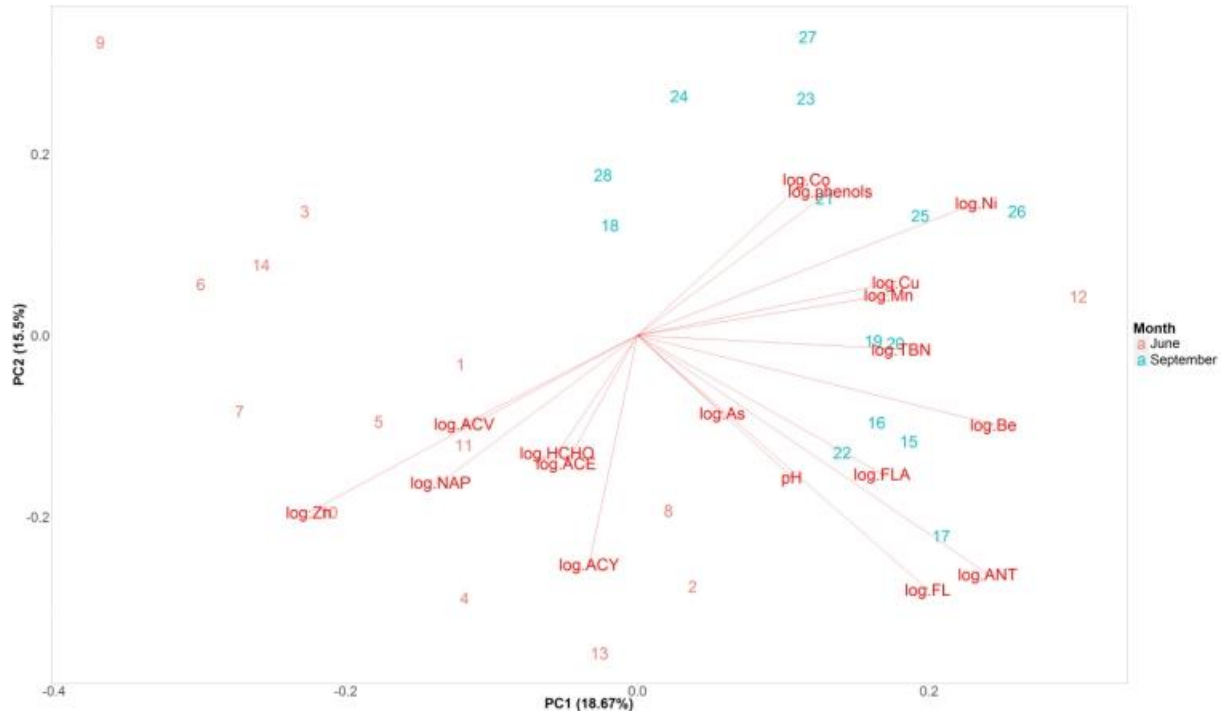
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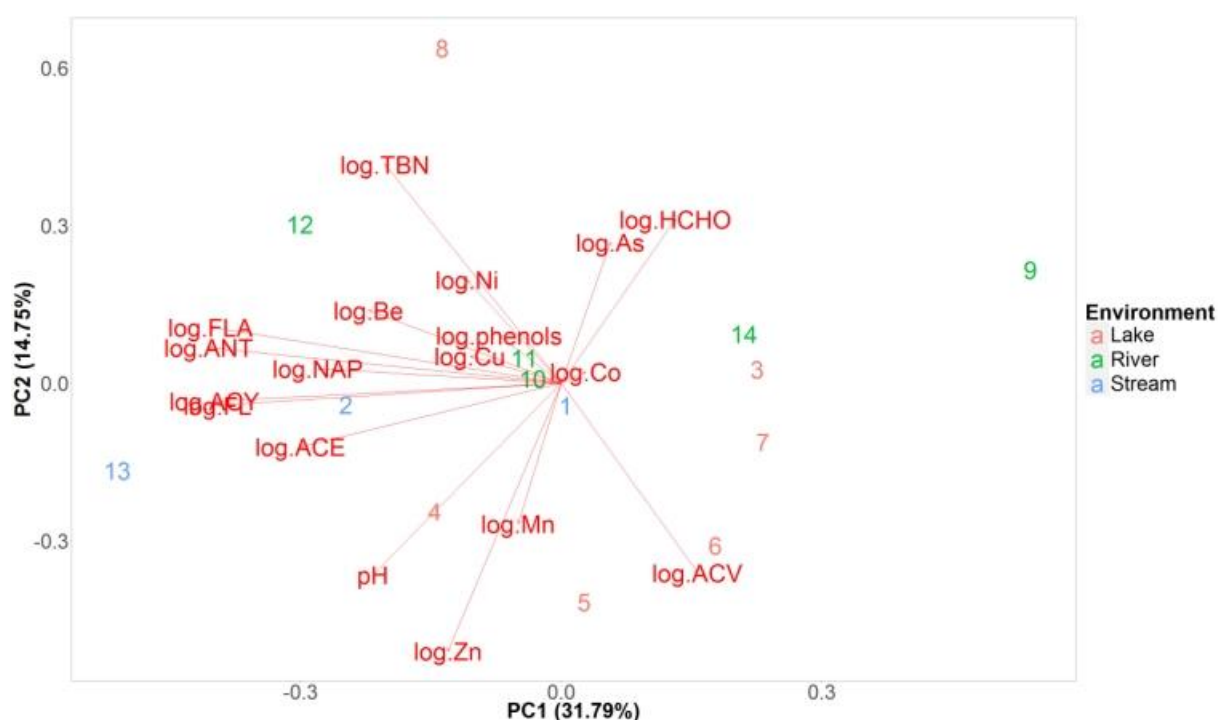
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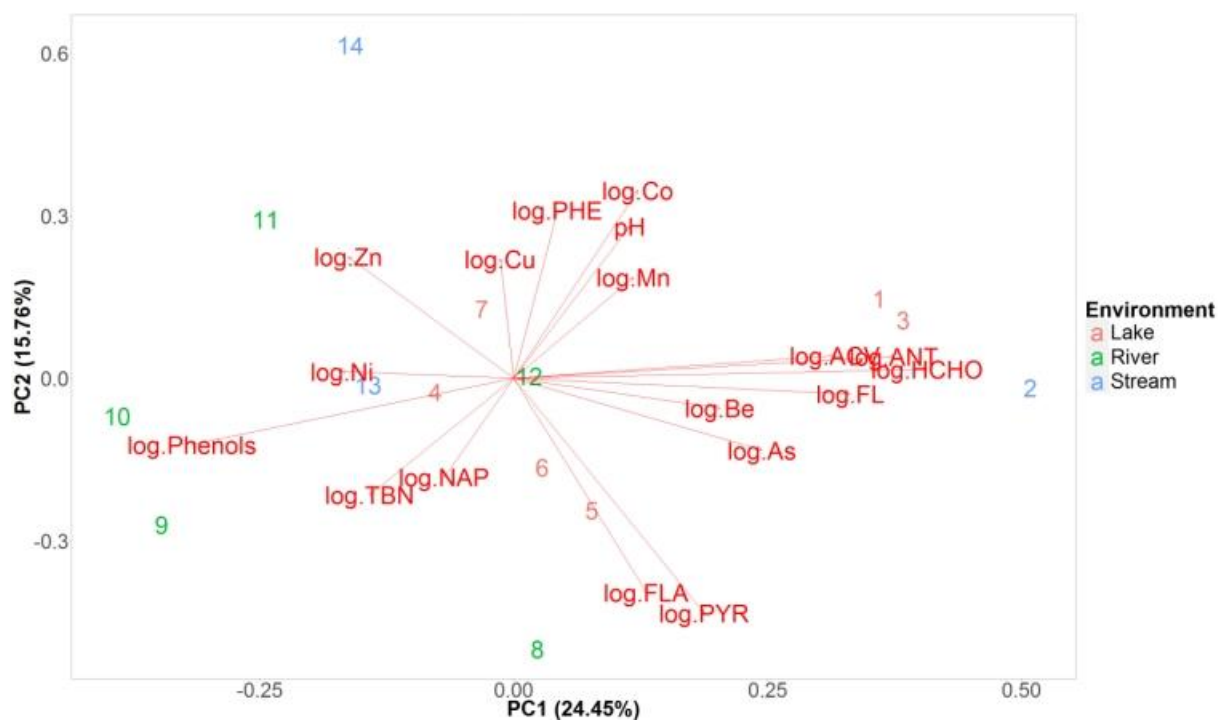
871 **Figure 2.** A box - whisker plot of the mean PAHs concentrations in water samples collected  
 872 in June and September 2016. Significant differences between seasons are indicated with the  
 873 non-parametric Kruskal-Wallis ANOVA p-levels below 0.05 and given in boldface. The box  
 874 encompasses the mean value  $\pm$  SD, the whiskers show the full range of values noted (where  
 875 <LOD values were assigned a half of the LOD level).



878 **Figure 3.** PCA conducted for the potentially toxic chemicals and indices of the bacterial  
 879 community structure detected in the hydrological environment of the Revelva catchment.  
 880 Top: The space defined by the PCs 1 and 2, with a division by the month of sampling  
 881 (colours). Bottom: Same analysis, space defined by the PCs 2 and 3, division by the type of  
 882 the hydrological environment. Numbers 1-14 depict samples from locations R1-14 in June,  
 883 numbers 15-28 denotes samples from the same locations in September, in the same order (e.g.  
 884 number 28 is the sample from point R14 in September).



885



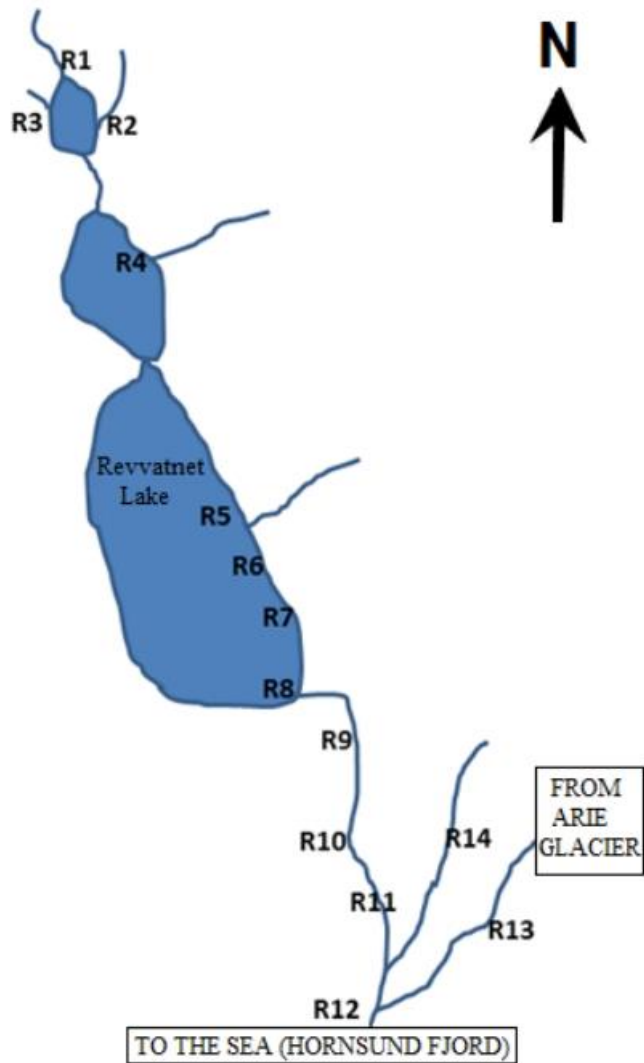
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887 **Figure 4.** PCA conducted for the potentially toxic chemicals detected in the different  
 888 hydrological environments of the Revelva catchment (denoted by colour coding) and indices  
 889 of the bacterial community structure in these waters. Each graph for a separate sampling  
 890 occasion, the space defined by the PCs 1 and 2. Top: June. Bottom: September. Numbers 1-14  
 891 depict samples from locations R1-14 in that particular month.

892

894 **Table 1.** Location of the sampling points in the Revvatnet catchment in Svalbard.

ID	Latitude	Longitude
R1	77° 02,174' N	15° 20,391' E
R2	77° 02,170' N	15° 21,021' E
R3	77° 02,113' N	15° 20,391' E
R4	77° 01,960' N	15° 21,282' E
R5	77° 01,437' N	15° 22,505' E
R6	77° 01,218' N	15° 23,385' E
R7	77° 01,122' N	15° 23,690' E
R8	77° 01,022' N	15° 24,077' E
R9	77° 00,841' N	15° 25,028' E
R10	77° 00,949' N	15° 24,686' E
R11	77° 00,640' N	15° 25,905' E
R12	77° 00,040' N	15° 26,675' E
R13	77° 00,332' N	15° 27,209' E
R14	77° 00,179' N	15° 26,902' E



896 **Table 2.** Validation parameters and technical specifications used in the applied analytical procedures.

<b>Determined compounds/parameters</b>	<b>Measurement range</b>	<b>LOD<sup>4</sup></b>	<b>LOQ<sup>4</sup></b>	<b>Measurement method/technique</b>
<b>pH</b>	-	-	-	Electrochemical method: microcomputer pH-meter(Elmetron), electrode type EPS-1
<b>∑ Phenols<sup>1</sup></b>	0.025-5.00	0.001	0.003	Spectrophotometry method;
<b>Formaldehyde<sup>1</sup></b>	0.020-8.00	0.005	0.015	Spectrophotometer 6300, Jenway
<b>PAHs<sup>2</sup></b>				Gas Chromatography technique coupled with Mass Spectrometry;
<b>Naphthalene (NAP)</b>	1.02-3500	0.034	1.02	Gas Chromatograph 7890A (Agilent Technologies) with the application of Mass Spectrometer (5975C inert MSD Agilent Technologies), detector (Agilent Technologies 5975C) with electron ionization
<b>Acenaphthylene (ACY)</b>	0.012-1000	0.004	0.012	
<b>Acenaphthene (ACE)</b>	0.012-1000	0.004	0.012	
<b>Fluorene (FL)</b>	0.005-1000	0.002	0.005	
<b>Phenanthrene (PHE)</b>	0.008-1000	0.003	0.008	
<b>Anthracene (ANT)</b>	0.023-1000	0.008	0.023	
<b>Fluoranthene (FLA)</b>	0.042-1000	0.014	0.042	



	<b>Pyrene (PYR)</b>	0.084-1000	0.028	0.084	
<b>Trace elements</b> 3	<b>Li, Be, Ga, Rb, U, Tl,</b>	0.010-1000	0.010	0.030	Inductively Coupled Plasma Mass Spectrometry technique;  (Thermo Scientific XSERIES 2 ICP-MS)
	<b>V, Cr, Mn, Co, Ni</b>				
	<b>Al, Cu, Zn, As, Ba</b>	0.100-1000	0.100	0.300	
	<b>Sr</b>	1.00-1000	1.00	3.00	

897 <sup>1</sup>[mg L<sup>-1</sup>], <sup>2</sup>[ng L<sup>-1</sup>], <sup>3</sup>[μg L<sup>-1</sup>], <sup>4</sup>the limit of detection (LOD) and the limit of quantification (LOQ) were calculated based on the standard deviation  
898 of the response (s) and the slope of the calibration curve (b), according to the formulas: LOD=3.3(s/b), LOQ=10(s/b)

899

900 **Table 3.** Concentrations ( $\pm$ standard deviation, SD) of trace elements in the collected  
 901 freshwater samples.

	June 2016	September 2016
<b>Trace elements</b>	<b>Li</b> 0.054 $\pm$ 0.011 – 0.529 $\pm$ 0.022	0.1120 $\pm$ 0.0020 – 0.379 $\pm$ 0.017
<b>[<math>\mu\text{gL}^{-1}</math>]</b>	<b>Be</b> 0.0020 $\pm$ 0.0010 – 0.0110 $\pm$ 0.0070	0.0020 $\pm$ 0.0010 – 0.0130 $\pm$ 0.0060
	<b>Al</b> 0.429 $\pm$ 0.021 – 3.456 $\pm$ 0.041	1.233 $\pm$ 0.053 – 5.92 $\pm$ 0.12
	<b>V</b> 0.0300 $\pm$ 0.0070 – 0.099 $\pm$ 0.013	0.0130 $\pm$ 0.0030 – 0.1190 $\pm$ 0.0040
	<b>Cr</b> 0.0180 $\pm$ 0.0070 – 0.25 $\pm$ 0.10	0.0070 $\pm$ 0.0010 – 0.176 $\pm$ 0.082
	<b>Mn</b> 0.0020 $\pm$ 0.0010 – 0.0250 $\pm$ 0.0030	0.0120 $\pm$ 0.0020 – 0.318 $\pm$ 0.014
	<b>Co</b> 0.0040 $\pm$ 0.0010 – 0.0380 $\pm$ 0.0020	0.0100 $\pm$ 0.0010 – 0.0290 $\pm$ 0.0050
	<b>Ni</b> 0.073 $\pm$ 0.021 – 0.305 $\pm$ 0.025	0.124 $\pm$ 0.029 – 0.322 $\pm$ 0.020
	<b>Cu</b> 0.091 $\pm$ 0.011 – 2.33 $\pm$ 0.30	0.170 $\pm$ 0.019 – 0.819 $\pm$ 0.059
	<b>Zn</b> 0.049 $\pm$ 0.012 – 2.04 $\pm$ 0.57	0.029 $\pm$ 0.011 – 0.128 $\pm$ 0.017
	<b>Ga</b> 0.0380 $\pm$ 0.0070 – 0.183 $\pm$ 0.017	0.0790 $\pm$ 0.0070 – 0.2720 $\pm$ 0.0070
	<b>As</b> 0.136 $\pm$ 0.035 – 0.451 $\pm$ 0.062	0.108 $\pm$ 0.013 – 2.40 $\pm$ 0.13
	<b>Rb</b> 0.1900 $\pm$ 0.0060 – 0.415 $\pm$ 0.016	0.2050 $\pm$ 0.0060 – 0.6020 $\pm$ 0.0090
	<b>Sr</b> 4.156 $\pm$ 0.032 – 33.01 $\pm$ 0.20	16.13 $\pm$ 0.26 – 43.44 $\pm$ 0.78
	<b>Ba</b> 1.855 $\pm$ 0.014 – 9.241 $\pm$ 0.053	3.384 $\pm$ 0.076 – 15.79 $\pm$ 0.29
	<b>Tl</b> 0.0110 $\pm$ 0.0020 – 0.0160 $\pm$ 0.0030	0.0120 $\pm$ 0.0030 – 0.095 $\pm$ 0.019
	<b>U</b> 0.0100 $\pm$ 0.0010 – 0.50 $\pm$ 0.22	0.0100 $\pm$ 0.0020 – 1.168 $\pm$ 0.031

902  
 903 **Table 4.** Concentrations ( $\pm$ standard deviation, SD) of PAHs, formaldehyde and the sum of  
 904 phenols in the collected freshwater samples.

		June 2016	September 2016	
<b>PAHs</b> [ng L <sup>-1</sup> ]	<b>Naphthalene</b> (NAP)	150±23– 611±40	87±10 – 318±22	
	<b>Acenaphthylene</b> (ACY)	1.06±0.24 – 47±12	0.43±0.12– 27.3±8.3	
	<b>Acenaphthene</b> (ACE)	2.9±1.2 – 57±16	2.1±1.0–111±12	
	<b>Fluorene</b> (FL)	3.0±1.1–371±29	6.6±1.9–318±21	
	<b>Phenanthrene</b> (PHE)	24.1±7.6 – 30.2±7.6	3.6±1.2 – 368±34	
	<b>Anthracene</b> (ANT)	8.9±2.1 – 1871±45	38.0±9.3 – 599±31	
	<b>Fluoranthene</b> (FLA)	7.4±1.9 – 985±41	21.9±6.6 – 991±48	
	<b>Pyrene</b> (PYR)	3.3±1.2 – 21.3±4.2	2.64±0.89 – 293±20	
	<b>Phenolic compounds,</b>	<b>∑ Phenols</b>	0.0120±0.008 – 0.078±0.019	0.031±0.010 – 0.085±0.020
		<b>HCHO</b>	0.160±0.066– 0.53±0.19	0.130±0.054 – 0.29±0.17
	<b>HCHO</b> [mg L <sup>-1</sup> ]			

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906 **Table 5.** The list of members of bacterial community possibly involved in the degradation of pollutants. A genus was considered only if it  
 907 constituted  $\geq 0.01\%$  of the community from a single sample.

Bacterial strains	R4-J	R4-S	R8-J	R8-S	R14-J	R14-S	Substrate	References
<i>Achromobacter</i>	0.12%	0.08%	0.12%	0.09%	0.20%	0.07%	PHE	Andreoni et al. 2004
<i>Acidovorax</i>	0.21%	0.30%	0.19%	0.38%	0.19%	0.34%	PHE	Eriksson et al. 2003; Martin et al. 2012
<i>Acinetobacter</i>	0.08%	0.05%	0.04%	0.02%	0.07%	0.10%	NAP, ANT, PHE, ACE, ACY	Ryu et al. 1989; Lal and Khanna 1996; Ghosal et al. 2013
<i>Actinocatenispora</i>	0.17%	0.22%	0.56%	0.21%	0.41%	0.17%	FL	Al-Mueini et al. 2007
<i>Arthrobacter</i>	0.05%	0.03%	0.03%	0.03%	0.08%	0.05%	FL, PHE	Grifoll et al. 1992; Casellas et al. 1997; Seo et al. 2006;
<i>Bacillus</i>	0.14%	0.16%	0.03%	0.03%	0.09%	0.13%	NAP, PYR	Samanta et al. 1999 Kumar et al. 2007; Kazunga and Aitken 2000
<i>Burkholderia</i>	0.21%	0.26%	0.07%	0.07%	0.17%	0.19%	NAP, PHE	Balashova et al. 1999; Seo et al. 2007; Laurie and Lloyd-Jones 1999a; 1999b
<i>Cycloclasticus</i>	0.02%	0.02%	0.01%	<0.01%	0.02%	0.02%	NAP, ANT, PHE, FL, PYR	Kasai et al. 2003; Geiselbrecht et al. 1998; Dyksterhouse et al. 1995; Wang et al. 2008; Kappell et al. 2014
<i>Dechloromonas</i>	0.01%	0.02%	0.01%	0.01%	0.02%	0.02%	NAP, ANT, PHE, FL, PYR	Coates et al. 2001a
<i>Desulfosporosinus</i>	0.22%	0.07%	0.01%	0.01%	0.05%	0.04%	Toluene	Sun et al. 2014
<i>Desulfotomaculum</i>	0.16%	0.13%	0.03%	0.02%	0.11%	0.03%	Biphenyl	<a href="#">Selesi</a> and <a href="#">Meckenstock</a> 2009
<i>Desulfovibrio</i>	0.26%	0.15%	0.10%	0.07%	0.22%	0.21%	NAP, ANT, PHE, FL, PYR	Villanueva et al. 2008



<i>Flavobacterium</i>	4.44%	2.75%	4.54%	2.19%	3.36%	1.55%	NAP	Widada et al. 2002; Kappell et al. 2014
<i>Geobacillus</i>	0.05%	0.02%	<0.01%	<0.01%	0.01%	0.01%	NAP	Bubinas et al. 2008
<i>Geobacter</i>	1.74%	0.96%	0.57%	0.16%	0.97%	0.92%	Benzoate	Coates et al. 2001b
<i>Janibacter</i>	0.01%	0.02%	<0.01%	0.01%	0.01%	0.01%	FL, PHE, ANT	Yamazoe et al. 2004
<i>Marinobacter</i>	0.01%	0.01%	<0.01%	0.01%	0.01%	0.02%	NAP, ANT, PHE	Al-Mailem et al. 2013; Kappell et al. 2014
<i>Marinobacterium</i>	0.01%	<0.01%	0.01%	<0.01%	0.02%	0.01%	NAP	Hedlund et al. 2001
<i>Methylobacterium</i>	0.03%	0.03%	0.01%	0.01%	0.02%	0.04%	PHE	Andreoni et al. 2004
<i>Micrococcus</i>	0.01%	<0.01%	0.01%	<0.01%	<0.01%	<0.01%	PHE	Ghosh and Mishra 1983
<i>Moraxella</i>	<0.01%	<0.01%	<0.01%	<0.01%	<0.01%	<0.01%	NAP	Tagger et al.. 1990
<i>Mycobacterium</i>	0.09%	0.07%	0.32%	0.21%	0.15%	0.12%	PYR, NAP, PHE, FLA, ANT, FL	Boldrin et al.. 1993; Schneider et al. 1996; Heitkamp et al. 1988; Churchill et al. 2008; Lee et al.. 2007; Van Herwijnen et al. 2003
<i>Mycobacterium vanbaalenii</i>	<0.01%	<0.01%	0.04%	0.02%	0.01%	0.01%	NAP, ANT, PHE, FLA, PYR	Kelley et al. 1990; Moody et al. 2001; Kelley et al. 1993; Kim et al. 2005
<i>Nocardia</i>	0.01%	<0.01%	0.01%	<0.01%	0.01%	0.12%	NAP, ANT, PHE	Zeinali et al. 2008a; 2008b
<i>Nocardioides</i>	0.05%	0.18%	0.01%	0.02%	0.02%	0.07%	PHE	Iwabuchi and Harayama 1997; Iwabuchi and Harayama 1998
<i>Novosphingobium</i>	0.27%	0.62%	0.05%	0.14%	0.34%	0.34%	NAP	Suzuki and Hiraishi 2007
<i>Ochrobactrum</i>	<0.01%	<0.01%	<0.01%	0.02%	0.01%	0.01%	PHE	Ghosal et al. 2010
<i>Paenibacillus</i>	0.07%	0.09%	0.03%	0.01%	0.18%	0.05%	NAP	Daane et al. 2001; 2002
<i>Paracoccus</i>	0.01%	0.01%	0.01%	0.13%	0.01%	0.07%	ANT, PHE, FL	Zhang et al. 2004
<i>Pasteurella</i>	0.07%	0.03%	0.41%	0.22%	0.22%	0.10%	FLA	Sepic and Leskovsek 1999
<i>Polaromonas</i>	5.08%	3.65%	2.40%	1.85%	2.49%	2.08%	NAP	Jeon et al. 2006
<i>Pseudoalteromonas</i>	0.18%	0.12%	0.06%	0.03%	0.12%	0.24%	NAP, PHE, FL	Hedlund and Staley 2006
<i>Pseudomonas</i>	0.19%	0.17%	0.09%	0.05%	0.25%	0.25%	PHE, NAP, PYR, ACE	Romero et al. 1998; Caldini et al. 1995;



								Weissenfels et al. 1990; Tian et al. 2003; Balashova et al. 1999; Kazunga and Aitken 2000; Prabhu and Phale 2003; Bosch et al. 1999
<b><i>Ralstonia</i></b>	0.06%	0.04%	0.03%	0.02%	0.05%	0.07%	NAP	Fuenmayor et al. 1998
<b><i>Rhizobium</i></b>	0.02%	0.01%	0.01%	<0.01%	0.01%	0.03%	ACE	Poonthrigpun et al. 2006
<b><i>Rhodococcus</i></b>	0.11%	0.19%	0.07%	0.09%	0.08%	0.10%	NAP, FL, ANT, FLA, PYR	Di Gennaro et al. 200; Dean-Ross et al. 2001; Dean-Ross et al. 2002; Walter et al. 1991
<b><i>Rhodoferrax</i></b>	2.68%	5.60%	1.57%	4.19%	1.56%	2.84%	PHE	Martin et al. 2012
<b><i>Shewanella</i></b>	0.06%	0.04%	0.02%	0.02%	0.04%	0.08%	NAP	Hilyard et al. 2008
<b><i>Sphingobium</i></b>	0.07%	0.03%	0.01%	<0.01%	0.02%	0.03%	NAP, PHE, ANT, FLA	Cavalca et al. 2007; Chadhain et al. 2007; Roy et al. 2012; Khara 2014; Keum et al. 2006
<b><i>Sphingomonas</i></b>	0.74%	0.80%	0.19%	0.44%	0.47%	0.98%	ACE, PHE, ANT, FLA, BaP, PYR	Pinyakong et al. 2004; Wattiau et al. 2001; Van Herwijnen et al. 2003b; Liu et al. 2004; Rentz et al. 2008; Kazunga and Aitken 2000
<b><i>Staphylococcus</i></b>	0.03%	<0.01%	0.04%	0.02%	0.01%	<0.01%	PHE	Mallick et al. 2007

908 NAP- naphthalene; ANT-anthracene; ACE-Acenaphthene; PHE-phenanthrene; FL-Fluorene; FLA-fluoranthene; PYR-pyrene; BaP-  
 909 benzo[*a*]pyrene; BaA-benz[*a*]anthracene.

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