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## 1 Insights on the particle-attached riverine archaeome at different

## 2 seasons and in response to multipollution during a Mediterranean

## 3 extreme storm event

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#### 25 Abstract

Even if Archaea deliver important ecosystem services and are major players in global 26 27 biogeochemical cycles, they remain poorly understood in freshwater ecosystems. To our 28 knowledge, no studies specifically address the direct impact of xenobiotics on the riverine 29 archaeome. Using environmental DNA metabarcoding of the 16S ribosomal gene, we 30 previously demonstrated bacteriome significant responses to pollutant mixtures during an 31 extreme flood in a typical Mediterranean coastal watercourse. Here, using the same 32 methodology, we sought to determine whether archaeal community shifts were also driven by 33 environmental stressors during the same flood. Further, we wanted to determine how archaea 34 taxa compared at different seasons. In contrast to the bacteriome, the archaeome showed a 35 specific community in summer compared to winter and autumn. We also identified a significant 36 relationship between *in situ* archaeome shifts and changes in physicochemical parameters 37 along the flood, but a less marked response to river hydrodynamics than bacteria. New urban-38 specific archaeal taxa, which were significantly related to multiple stressors, were identified. 39 Through statistical modeling of both domains, our results demonstrate that Archaea, seldom 40 considered as bioindicators of water quality, could provide a rapid assessment of microbial 41 pollution risk and thus have the potential to improve monitoring methods of watersheds. 42

43 Keywords: microbial ecotoxicology; water quality; extreme storm event; coastal
44 Mediterranean rivers; sewer overflow; environmental DNA; metabarcoding.

#### 45 **1. Introduction**

46 In the 1970s, Woese and collaborators highlighted a new domain of life, the Archaea, distinct 47 from Bacteria and Eukaryota domains (Woese and Fox 1977; Woese et al. 1978). With the 48 advent of Next Generation Sequencing (NGS), archaeal research and knowledge are expanding 49 (Adam et al. 2017; Bang and Schmitz 2018). Archaea are recognized as major players in the 50 global biogeochemical cycles of carbon, nitrogen, hydrogen and sulfur (Casamayor and 51 Borrego 2009; Offre et al. 2013; Castelle et al. 2015) and there are at least two metabolisms, 52 essential for nutrient cycling, which are carried out exclusively by archaea: methanogenesis 53 and anaerobic methane oxidation (Joye 2012). Despite their importance for ecosystem 54 functioning and their ubiquitous presence (Chaban et al. 2006; Casamayor and Borrego 2009; 55 Herfort et al. 2009; Auguet et al. 2010), the environmental archaeome has been overlooked for 56 decades. Most studies have concentrated on marine archaea (Zinger et al. 2012; Zeglin 2015), 57 and the diversity and importance of archaea in other environments have been largely 58 disregarded (Casamayor and Borrego 2009; Zinger et al. 2012; Zeglin 2015; Adam et al. 2017). 59 Only recently, continental freshwater habitats have emerged as one of the largest reservoirs of 60 archaeal diversity (Auguet et al. 2010; Zinger et al. 2012). Some studies have linked archaeal 61 communities to water physicochemical properties such as pH, temperature and nutrients 62 (Herfort et al. 2009; Wang et al. 2018; Lei et al. 2020). Methanobrevibacter smithii, for 63 example, was proposed as a potential indicator of human-specific sewage pollution (Johnston 64 et al. 2010; McLellan and Eren 2014). To assess water quality, several studies have focused on 65 freshwater archaea (Cannon et al. 2017; Dila et al. 2018), notably for their potential remediation 66 role in heavily contaminated urban rivers (Samson et al. 2019; Lei et al. 2020). Cannon et al. 67 (2017) found that a rain event induced changes in the structure of microbial communities, 68 including archaea, from environmental DNA (eDNA) and stressed the importance of 69 considering hydrological conditions when studying riverine microbiomes. Both chemical and 70 microbial pollutants reach surface waters via point sources (such as urban sewage) or diffuse 71 sources (linked to runoff) of pollution. It is well known that during rainfall events, particles 72 from terrestrial soils and river basin sediments remobilize, and affect water quality (Garcia-73 Esteves et al. 2007; Dumas et al. 2015; Faure et al. 2015) because soils and sediments store 74 pathogens, nutrients and pollutants. Thus, suspended particles have a crucial role in the transfer 75 of contaminants to surface waters through runoff and in the resuspension of river sediments 76 during storms (Turner and Millward 2002; Amalfitano et al. 2017). Moreover, some studies 77 have reported that riverine archaea tend to be associated with particles and derive only from

78 allochthonous inputs (Crump and Baross 2000; Casamayor and Borrego 2009; Hu et al. 2018). 79 This is all the more important in Mediterranean regions, where extreme hydrological events 80 are expected to become more intense and frequent due to climate change (Cowling et al. 2005; 81 Blanchet et al. 2016). Furthermore, first-flush events during rainfalls lead to Combined Sewer 82 Overflows (CSOs), which carry large loads of contaminant mixtures over surface waters 83 through in-sewer solids resuspension (Ashley et al. 1992; Osorio et al. 2012; Oursel et al. 2014; 84 Reoyo-Prats et al. 2017, 2018). We previously demonstrated that an extreme Mediterranean 85 flood produced shifts in the particle-attached bacterial compartment from eDNA, severely 86 affecting resident riverine communities (Noyer et al. 2020). In this study, we went a step further 87 by using metabarcoding of the 16S ribosomal RNA gene (rDNA) to explore how the diversity 88 of the particle-attached archaeome changed within the same environmental samples, and to 89 compare the responses between the bacteriome and the archaeome. Next, we modeled shifts in 90 both communities using physical parameters as well as several families of chemical parameters 91 as environmental forces, which is a first in environmental microbiology. This study addressed 92 the following questions: how did the fluviatile particle-attached archaeal community change 93 between seasons and how did it evolve over the course of a heavy rain event? How our findings 94 compare to other studies of archaeal alpha and beta diversity in lotic ecosystems? Did 95 environmental parameters drive structural shifts in the archaeome as they did in the 96 bacteriome? Was there a strong relationship between key taxa and environmental dynamics? 97 How did the seasonal and temporal succession of bacterial and archaeal communities compare? 98 The answer to these questions could, in general, provide insights on the use of microorganisms 99 in water quality assessment and, in particular, help on the rapid risk assessment of multiple 100 pollutants in aquatic ecosystems.

#### 101 **2. Materials and Methods**

#### 102 **2.1. Study site, sampling information**

Sampling took place in the Têt River, a watercourse representative of Mediterranean coastal watercourses with a torrential regime that discharges into the Gulf of Lion (Southeast of France) (Dumas et al. 2015; Reoyo-Prats et al. 2017) downstream from the Perpignan city wastewater treatment plant, the main threat to the water quality of this river (Fig. 1a, Conseil Général des Pyrénées Orientales 2009, 2012; Reoyo-Prats et al. 2017). The Vinça dam, situated 40 km upstream from the sampling station, controls the downstream river flow, particularly during floods. For sampling methodology and sampling site see Reoyo-Prats *et al.* (2017). In
short, ten liters of river water were collected in summer, winter, and autumn, before, during
and after a 5-year flood that we followed 24h/24h during more than 4 days (see Fig. 1b for
further details).

113 First flushes during the flood brought the highest levels of *E. coli* and *Enterococci* ever detected 114 in this river (Reoyo-Prats et al. 2017) as well as relative higher abundance of other typical 115 sewer bacteria (Nover et al. 2020) and higher levels of dissolved pharmaceuticals (Reovo-Prats 116 et al. 2018), which indicated the moment at which sewer overflows occurred (Fig. 1b). 117 Environmental DNA sampling was previously described by Noyer et al. (2020). In short, a liter 118 of mixed-water sample was entirely (or until clogged) filtered through cellulose acetate MF-119 Millipore membrane filters with 3 µm porosity (Merck Darmstadt, Germany) and repeated 120 three times to obtain three replicates per sample.

## 121 2.2. Nucleic acid extraction, 16S rRNA gene sequencing, and sequence analyses to 122 obtain Operational Taxonomic Units (OTUs) contingency table

123 Nucleic acid extraction and bacterial sequencing from eDNA were described in a 124 previous study (Noyer et al. 2020) and were performed in triplicate per sample. For archaea 125 sequencing, a single set of DNA replicates from each sample was sent to the Research and 126 Testing Laboratory (RTL, Texas, USA). Two more replicates were later sent to the Genome 127 Quebec laboratory (GQ, Montreal, Canada), except for samples t61 and t68 which were only 128 sequenced once again for technical reasons. DNA from samples t19 and t23 from the first 129 replicate, which was already sequenced at RTL, was also sent to GQ to be sequenced again for 130 comparative purposes. Sequencing targeted V3 and V4 hypervariable regions of the 16S rDNA 131 by using 519wF (5'-CAGCMGCCGCGGTAA-3') and 1017R (5'-132 GGCCATGCACCWCCTCTC-3') universal archaeal primers (Borrego et al. 2020) and was 133 performed on an Illumina MiSeq sequencer using a 2x300bp paired-end protocol. Libraries 134 were generated by pooling equimolar ratios of amplicons before sequencing. In contrast to RTL 135 sequences, those provided by GQ contained primers. In order to pool sequences together, we 136 eliminated primers from GQ sequences using Cutadapt version 1.8.3 for Unix (Martin, 2011). 137 Default options were used with the exception of sequences treatment as paired with -g and -G 138 options for forward and reverse primers respectively, taking wildcards into account, discarding 139 untrimmed sequences and setting an overlap of 14 bp, a quality-base of 33 and an error-rate of 140 0.1. Harmonized sequences were pooled together and archived before being uploaded to the

141 Galaxy platform (Afgan et al., 2016): http://sigenae-workbench.toulouse.inra.fr. FROGS 142 pipeline (v. r3.0-3.0, Escudié et al. 2018) was used to process sequences to form OTUs and to 143 taxonomically affiliate them as described in Noyer et al. (2020) except that VSEARCH (v2.6.2, Rognes et al. 2016) was used as a read pair assembler, which allows a higher number of 144 145 sequences to be conserved when amplicon sizes are highly variable, as was the case for our 146 dataset. Read length was set to 300pb and the amplicon size was set to 450pb for the minimum 147 and optimized to 545pb for the maximum. We assigned each OTU up to the species taxonomic 148 level based on blast alignment using the affiliation tool (v. r3.0-2.0) and the Silva 138.1 149 database with a pintail score of 100, which allowed for the most accurate affiliation possible. No archaea hits were removed hereafter. To resolve the taxonomic ambiguity of OTUs that 150 151 were multi-affiliated within the FROGS pipeline when using blast against the curated pintail 152 100 score silva database, we blasted these OTUs against the NCBI 16S nucleotide collection

153 database using the megablast algorithm.



154 Fig. 1 Têt River archaeome sampling sites and environmental parameters measured in the same samples. (a) 155 Watershed of the river with sampling site (black arrow), located after wastewater treatment plan (WWTP) of the 156 city of Perpignan, combined sewers (black crosses) and water reservoirs indicated as grey rectangles (adapted 157 from Reovo-Prats et al. 2017). (b) Environmental parameter dynamics in the Têt River at different seasons and 158 159 along an autumn flood. For sample names, see table below figure. Autumn sample names are followed by a number that indicates the sampling time in hours after t0, which was sampled at autumn basal level water 160 discharges. Sampling took part at crucial moments of the flood that occurred thereafter: at first flushes (t17-t19-161 t23), before the flow peak (t32-t37), during the flow peak (t38-t41), following the release of water from the 162 upstream Vinça reservoir (t44-t61) and during the return to basal level (t68-t86-t109). ProxyDyn1 corresponds to 163 the dynamics of particulate organic carbon (/20 mg/l), which represented the dynamics of water flow, also 164 represented in figure, total suspended solids, total organic carbon, total nitrogen, and terbuthylazine parameters. 165 ProxyDyn2 corresponds to aminomethyl phosphonic acid (AMPA, µg/l), which represented glyphosate, 166 phosphate, copper, temperature, E. coli, enterococci, diclofenac, sulfamethoxazole and carbamazepine 167 parameters. ProxyDyn3 corresponds to lead ( $/150 \mu g/g$ ) in the representation of the dynamics of cadmium, zinc, 168 and conductivity parameters. ProxyDyn4 corresponds to pH (/70), which represented cobalt, nickel, and chrome 169 parameters. Three parameters, Diuron, 2.4D and NO<sup>3-</sup>, had a unique dynamic. For further details on statistical 170 analyses for environmental parameters, see Noyer et al. (2020).

#### 171 **2.3. Archaeal diversity analyses**

172 Diversity analyses were performed using the output OTUs contingency table, tree and 173 dissimilarity matrices calculated using FROGS as input for *Phyloseq* within R package 1.24.2 (McMurdie 174 Holmes 2013) and a collection of additional R functions and 175 (https://github.com/mahendra-mariadassou/phyloseq-extended). Trimming rare OTUs affects 176 alpha-diversity measurements sensitive to rare OTUs such as Chao1, Observed richness and 177 Shannon indices, while rarefaction is controverted as well when concerning some alpha diversity indices (McMurdie and Holmes 2014; Cameron et al. 2021). Alpha diversity was 178 therefore calculated using non-filtered and non-normalized replicates from the GQ laboratory 179 180 only, because the lower sequence depth of RTL sequenced replicates precluded comparison. 181 Fisher, Simpson, Shannon, and Pielou alpha diversity indices were used, together with the 182 nonparametric Chao1 species richness estimator. These indices provide complementary 183 information regarding evenness and richness aspects of alpha diversity that are interesting to take into account (see, for instance, Walters and Martiny 2020). Kruskal-Wallis test (KW) 184 185 followed by a post hoc Dunn test with R software (v. 3.5.1, R Core Team 2018) were applied to evaluate diversity changes through time. Beta diversity was assessed on all replicates, 186 187 independently of platform origin, because sequencing depth is not relevant in this case. 188 Singletons were filtered out and then abundance was normalized to the sample with the lowest 189 number of sequence reads. Using this dataset, relative abundances by phylum and class were 190 plotted. To detect potential outliers in the dataset we proceeded by (i) checking the number of 191 sequences of each replicate, (ii) checking OTU abundance distribution among replicates of the 192 same sample and (iii) calculating qualitative Jaccard and quantitative Bray-Curtis 193 dissimilarities from replicates separately and using Principal Coordinates Analysis (PCoA) to

194 visualize replicate dissimilarities. Once outliers checked, beta dissimilarities were recalculated 195 on the averaged OTU abundances. To this end, the number of OTUs decreased to less than 196 10,000 after the dataset was normalized, thus allowing qualitative Unifrac and quantitative 197 Weighted-Unifrac (W-U) dissimilarities, which also consider phylogeny of OTUs, to be 198 included within FROGS. PCoA and hierarchical clustering Ward.D2 methods were used to 199 visualize archaeal community dissimilarities among samples. A one-way analysis of similarity 200 (ANOSIM, Clarke 1993) was performed to test significant differences between sample groups 201 resulting from hierarchical clustering. To further check for significant differences in archaeal 202 community shifts at the class level, the Mann-Whitney test (MW) was implemented with R 203 software.

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#### 2.4. Statistical analyses for inference

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#### 2.4.1. Constrained (canonical) ordination analyses by environmental parameters dynamics

206 Physicochemical environmental parameters were previously measured (Reovo-Prats et al. 207 2017) in the same samples in which nucleic acid extractions were performed. Measured 208 parameters included pH, temperature, conductivity, flow, total suspended solids, particulate 209 organic carbon, total organic carbon, nitrogen and 250 pesticide molecules, 90 210 pharmaceutically active compounds, polycyclic aromatic hydrocarbons and polychlorinated 211 biphenyls, nutrients, trace metals in the particulate fraction and fecal indicators (load in Escherichia coli and Enterococci). Collinearity issues resolution and the seven major 212 213 environmental dynamics retained for further analyses are described in Noyer et al. (2020). For 214 clarity, the retained variables that will be used for further analyses as proxies of correlated 215 environmental variables are summarized in Fig. 1b. Constraint-based ordination analyses were 216 then used to evaluate the relationships between the normalized OTU contingency table and 217 retained environmental parameter dynamics using vegan R package version 2.5-3 (Oksanen et 218 al. 2018). Detrended correspondence analysis (DCA) performed on the OTU dataset rendered 219 a first axis gradient length of 3.7, so both canonical correspondence (CCA) and redundancy 220 analyses (RDA) were performed (ter Braak 1988). We also performed a Hellinger transformed-221 based RDA (tbRDA) (Legendre and Gallagher 2001) and a distance-based RDA (dbRDA) 222 using all beta dissimilarities from the previous section. Permutation analyses of variance were 223 used to evaluate the significance of constraint-based models, axes, and variables. Variables 224 were tested by adding each of them independently and the number of permutations was set to 225 10,000. To further determine which OTUs best responded to environmental variables, we 226 reduced the OTU matrix to a percentage of abundance so that CCA/RDA modeling became

significant. Instead of 0.005%, as reported by Bokulich *et al.* (2012) and Noyer *et al.* (2020) for bacteria, we found a percentage of 0.05% for the archaeal dataset. RDA modeling of OTUs with  $\ge$  0.05% of the total read number (i.e. keeping 2,688,328 reads), allowed for the OTU goodness of fit (GOF) to be calculated. As for bacteria, OTUs retained for further analyses had a GOF  $\ge$  GOF<sub>average</sub>, which is considered a conservative approach to OTU selection.

232 2.4.2. Network construction via module eigengene analysis

233 We used the Molecular Ecological Network Analyses Pipeline (MENAP: 234 http://ieg4.rccc.ou.edu/MENA/) to build the relationships among OTUs following the 235 developer's recommendations (Deng et al. 2012; Tao et al. 2018) but sample-specific OTUs 236 were kept for network construction by changing "OTUs present at least in one sample". An 237 automatically generated similarity threshold value (0.32) was obtained with Random Matrix 238 Theory (RMT)-based method, which allowed network construction ensuring that the 239 connections between microorganisms were non-random ( $R^2$  of power-law = 0.28). The network 240 was separated into modules via the short random walk method (Pons and Latapy 2005), which 241 had the highest modularity (0.078) (Newman 2004). Module higher-order organization was 242 then performed via eigengene analysis (Langfelder and Horvath 2007) using default parameters 243 to obtain the correlation significance between modules and environmental parameters 244 dynamics. Cytoscape software (v. 3.7.1, Shannon et al. 2003) was used to visualize this 245 constraint-based network. We also used MENAP to explore the relationship between OTUs 246 from Bacteria (from the previous study by Noyer et al., 2020) and Archaea domains and flood 247 environmental dynamics. The network was constructed with the same parameters as before except zero counts were replaced "by 0.01" instead of "on paired values", to avoid losing an 248 249 important number of OTUs when using the matrix of joined OTUs from both domains. The 250 highest modularity was obtained via the leading eigenvector method (0.520). The RMT 251 threshold was 0.940 and the  $R^2$  of power law = 0.863.

#### 252 **3. Results**

### 253 254

### 3.1. Changes in particle-attached archaeal diversity along a storm event in the Têt River

To study changes in an urban archaeome along a storm event, a high-resolution environmental
DNA sampling from river water was performed during a major flood that occurred in autumn
2013. Two other seasons were also sampled during the summer and winter drought periods.

258 All reported sequence data passed standard quality controls of certified sequencing companies.

A total of 1,804,302 reads corresponding to 350,106 operational taxonomic units (OTUs) were

260 identified. Most OTUs (98%) were assigned to the archaeal domain, the rest were eliminated

261 from further analysis.

262 Alpha diversity statistical analyses showed significant differences among samples for Shannon (KW = 0.03, Fig. 2) and Pielou (KW = 0.03, Table S1) indices only. These two indices changed 263 264 similarly along seasons and during the flood, with equivalent significant differences between 265 samples except for the Pielou index for the summer drought, which was significantly higher than the flow peak (t38-t41) indicating a greater evenness in OTU abundances in this sample. 266 In general, both indices decreased significantly at t41, i.e. right after the flood peak, with 267 respect to t17 and t19. Three hours later, a significant increase in diversity was noticed (t44). 268 269 Even if the observed OTU number, Chao1 and Fisher indices did not show significant 270 differences (Table S1), they had the same change pattern as the Shannon index (Fig. 2), except 271 for a much lower value in the summer drought sample. This result contrast with the Pielou and 272 Simpson indices (Table S1), which emphasize the evenness component of diversity, as opposed 273 to Fisher and Shannon indices, which appear more related here to the richness component of 274 diversity (Magurran 2004).



275

Fig. 2 Alpha diversity of the archaeome of the Têt river along time. Changes in observed OTU number (blue) and
Shannon index (red) along the flood (tX) and at summer and winter droughts (SD and WD respectively). For the

Shannon index, different letters indicate a significant difference between samples (dunn.test<0.025) and red arrows show major significant differences. Observed OTU number was not significantly different (KW=0.14). The dotted profile is the flow level at each sampling point (see Fig. 1, also for sample names). Even though the absence of replicates for t61 and t68 samples impeded statistical testing, they are represented through time for comparison.</p>

283

284 After normalization of OTU matrix once singletons were excluded, a total of 1,377 OTUs 285 (3,777 sequences/sample) were retained for further analyses. We noticed a great difference in 286 the taxonomic composition of the summer drought (SD) with respect to all other samples. SD 287 had a significantly lower amount of Nitrososphaeria and Thermoplasmata classes when 288 compared to all samples (Fig. 3, MW p=0.012 and p=0.021 respectively) and a higher amount 289 of Nanoarchaeia (Nanoarchaeota phylum, Fig. 3) and Bathyarchaeia (Crenarchaeota phylum) 290 classes when compared to most samples. These large differences in the summer sample community were confirmed with the beta diversity analyses (data not shown), as all samples 291 292 significantly separated from the SD sample with all beta dissimilarity indices, contrarily to 293 WD, which always clustered with t17, t19, and t32. Additionally, the taxonomic composition 294 of the WD sample was very similar to t17, t19, and t32 samples, mainly composed of 295 Nitrososphaeria class, followed by Nanoarchaeia, Thermoplasmata, and then three classes, 296 Methanobacteria (Euryarchaeota phylum), Methanomicrobia (Halobacterota phylum) and 297 Bathyarchaeia at variable smaller abundances (Fig. 3).





303

304 Given these results, only flood samples were used to further explore structural diversity 305 changes using beta dissimilarities. These samples comprised a total of 1,739 OTUs (19,355 306 sequences/sample) after eliminating singletons and normalization. Using different beta 307 dissimilarities allowed for a better assessment of which differences are responsible for the 308 community structure (either presence/absence or abundance and/or phylogeny of OTUs). 309 Using only OTU presence/absence with Jaccard qualitative dissimilarity, three communities 310 were significantly differentiated (Fig. S2.1a). The first community group included samples 311 collected at t0, t17, t19 and t32, the second included samples t23, t37, and t38, and the third included the rest of the samples (t41-t109), which was the most differentiated cluster. But when 312 313 phylogenetic relationships were considered using Unifrac qualitative dissimilarity, t41 sample 314 grouped with t23, t37 and t38 instead (Fig. S2.1b). With Unifrac, this last cluster was the most 315 differentiated. When Bray-Curtis quantitative dissimilarity, which considers OTU abundance, 316 was used, three community groups of samples were distinguished (Fig. S2.1c). The first 317 included from t17 to t38 samples, the second included t0, t41, and t68 samples, and the third 318 t44, t61, t86, and t109 samples. When considering phylogenetic relationships using W-U, 319 observed community groups coincided with those of Bray-Curtis dissimilarity (Fig. 4, bold 320 black line), but the most significantly differentiated communities were those from the group of 321 t23, t37 and t38 (axis 1) and then t41 and t0 samples (axis 2). Even though the first taxonomic 322 changes occurred from the first flood sample at t17, samples t23, t37 and t38 had a particular 323 significant increase in Methanobacteria (Euryarchaeota phylum) (Fig. 3, MW, p=0.001). 324 Sample to differentiated significatively from samples from the end of the flood (from t44 to 325 t109) only when considering OTUs presence/absence but not when abundance alone or with 326 phylogeny were considered. This sample had a significantly lower abundance of taxa from the 327 Thermoplasmata class with respect to that group of samples (Fig. 3b, MW p=0.003). Regarding 328 t41 sample community, it had a particular taxonomy, significantly dominated by 329 Nitrososphaeria class (MW p=0.002) and W-U dissimilarity significantly separated t41 from 330 the rest of the samples (Fig. 4). With regard to the group of samples at the end of the flood 331 event, from t44 to t109, we could notice a significant increase of Thermoplasmata class with 332 respect to all other flood samples (Thermoplasmatota phylum, MW p=0.002, Fig. 3b). 333



Fig. 4 Structure of archaeal communities averaged across replicates. (a) Principal Coordinate Analysis (PCoA)
 and (b) hierarchical clustering with Ward D2 linkage method using Weighted-Unifrac dissimilarity computed on
 OTU average abundance. Lines indicate ANOSIM significant groups. Sample names are followed by a number
 that indicates the sampling time in hours after t0 (see Fig. 1 for details). Significant codes \*\* and \*\*\* indicate p value < 0.01 and < 0.001, respectively.</li>

**Table 1** Summary of constraint-based multivariate statistical models on archaea OTU matrix averaged over341replicates and without singletons. (a) Permanova significance of the five models tested and the percentage of342biological variance that is explained by each model using permutation test with anova.cca function. (b) Axes and343modeled variables significance after permanova using anova.cca of significant models in (a). Axes not shown344were not significant. p-values significance codes: (\*\*\* < 0.001 < \*\* < 0.01 < \* < 0.05).</td>

a	OTUs matrix transformation	Model significance	Variance (%)		
	Jaccard	0.003**	63.38%		
	Unifrac	0.008**	66.35%		
<b>UDKDA</b>	<b>Bray-Curtis</b>	0.326	60.04%		
	Weighted-Unifrac	0.004**	76.58%		
CCA		0.765	56.44%		
RDA	Hellinger	0.134	61.4%		

b	<b>OTUs matrix</b>	Axes significance and variance explained (%)	Statistical significance of modeled variables						
	transformation	CAP1	ProxyDyn1	ProxyDyn2	ProxyDyn3	ProxyDyn4	2.4D	Diuron	NO <sub>3</sub> -
dbRDA	Jaccard	0.002** (22.74)	0.001***	0.002**	0.001***	0.105	0.095	0.001***	0.096
	Unifrac	0.040* (30.24)	0.001***	0.021*	0.046*	0.458	0.383	0.036*	0.014*
	Weighted-Unifrac	0.006** (42.89)	0.006**	0.009**	0.039*	0.050*	0.004**	0.002**	0.166

#### **346 3.2. Modeling archaeome diversity according to multiple contaminant dynamics**

Constrained multivariate analyses were performed to determine if the retained environmental 347 dynamics (Fig. 1b) could statistically explain the observed community structure and diversity 348 349 shifts through time. Significance and percentage of variance explained by all models tested 350 were summarized in Table 1a. Jaccard, Unifrac and Weighted-Unifrac dbRDAs supported 351 significantly (p-value < 0.01) a link between environmental parameters included in the model 352 and our community data (Fig. S2.2). Models using Jaccard and Unifrac dissimilarities 353 explained between 63 and 66% of the total variance, respectively. Only the first canonical axis 354 was significant in both models, with a higher proportion of the variance in the dissimilarity matrix explained when phylogeny was considered using Unifrac (33% vs 23% for Jaccard, Fig. 355 356 S2.2a-b). In both models, the same four environmental dynamics were significant, ProxyDyn1, ProxyDyn2, ProxyDyn3, and Diuron (Table 1b). W-U dbRDA model performed best, 357 358 explaining 78% of variance in the OTU matrix. The first axis was significant and explained 44% of the total variance. Four environmental dynamics were significant according to this 359 360 model, ProxyDyn1, ProxyDyn2, 2.4-Dichlorophenoxyacetic acid (2.4D), and Diuron (Table 361 1b, Fig. S2.2c). Finally, only when the raw matrix was reduced to  $OTUs \ge 0.05\%$  of total read 362 number (see section 2.4.1 for details) tbRDA became significant (p-value = 0.046, DCA first axis length < 3), with one significant axis and three significant dynamics, ProxyDyn1 (p-value 363 364 = 0.032), ProxyDyn2 (p-value = 0.007) and Diuron (p-value = 0.001) (Fig. 5). This matrix 365 included 53 OTUs, and 284,506 reads, i.e. 3% of total OTUs, representing 69% of the total 366 reads) which were conserved for further analyses. This model explained 66.32% of the total 367 variance, and the first axis was significant and explained 37.6% of the variance. Samples well 368 projected to ProxyDyn2 and Diuron were t23 and t37, followed by t17 and t19, and to a lesser 369 extent t38 (see Fig. 5), which was well projected to ProxyDyn1. Notice however that axis 2 370 was not significant.



372 Fig. 5 Redundancy analysis (RDA) biplot with scaling by sites on the normalized matrix of OTUs with an 373 374 375

abundance  $\geq 0.05\%$ . The model explained 66.32% of the variance (p<0.05). Significance for axes and environmental dynamics after permanova analyses are indicated, p-value significance codes: \*\*\*<0.001<\*\*<0.01<\*<0.05. Sample names are followed by a number that indicates the sampling time in hours 376 after the beginning of the flood at t0. For further details on sample names and retained environmental variable 377 dynamics, see Fig. 1b. Perpendicular grey lines represent the projection of the corresponding samples onto the 378 corresponding dynamics and approximate the value of that sample along the variable (Legendre and Legendre 379 2012).

#### 380

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### **3.3.** Constrained molecular ecological network analyses by environmental

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#### 3.3.1. Archaeal network

parameters dynamics

383 Network analysis using MENAP resulted in four modules containing all OTUs and 774 links. Module eigengene analysis allowed correlation of modules with environmental variables. Only 384 385 one module was significantly correlated to the environmental dynamics of the flood, particularly with ProxyDyn2 and Diuron (Fig. 1b). This module included 28 OTUs of which 386 387 17 had a significant module membership (Fig. 6a), comprised of 11 with positive (52,149 reads) and six with negative (18,573 reads) correlations (Table S3.1). Worth noting is that 388 389 OTUs with positive module membership (Fig. 6b) had positive interactions with each other 390 (Fig. 6a) and negative interactions with OTUs with negative module membership (Fig. 6c), and 391 OTUs with negative module membership had positive interactions with each other and negative 392 interactions with OTUs with positive module membership. All positively correlated OTUs 393 were abundant in samples t23 (20,246 reads, which represent 39% of reads along the flood

394 from these OTUs, Table S3.1), t37 (20%), t19 (16%) and t17 (13%). A total of 60% of reads from these OTUs belonged to the Methanobacteria class (Euryarchaeota phylum). Two OTUs 395 396 were particularly abundant, OTU3, matching Methanobrevibacter (Mbr.) smithii at 99% 397 similarity after blastn search against NCBI database, and OTU11 matching Methanobacterium 398 (Mba.) palustre (100%). The next most abundant OTUs also matched methane-related taxa: 399 OTU41, which matched Methanosaeta (Msa.) concilii, 100%, from Halobacterota phylum), 400 OTU19 (Methanogranum sp. 98.04%, from Thermoplamatota phylum), OTU26 (Mbr. 401 acidurans, 99%, from Euryarchaeota phylum) and OTU16 (Methanomethylophilacea, 99.61%, 402 from Thermoplamatota). The other positively correlated OTUs (not highlighted in bold in 403 Table S3.1) did not match any further than the family level after a blast search of the NCBI database. Finally, negatively correlated OTUs were abundant in samples from the end of the 404 405 flood (t44-t109) and belonged to Nitrososphaeria (Crenarchaeota phylum, representing 85% of the sequences of these OTUs) and Thermoplasmata (15%) classes (Thermoplasmatota 406 407 phylum).



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409 Fig. 6 (a) Molecular ecological network of the unique module significantly positively correlated with 410 environmental dynamics, particularly ProxyDyn2 and Diuron (see Fig. 1b for details and sample names). (b) 411 Histogram of total reads of OTUs with positive module membership in function of samples along the flood. (c) 412 Same for OTUs with negative module membership. Environmental dynamics are followed by module correlation 413 value between parenthesis and the p-value significant code as follows: \*\*\*<0.001<\*\*<0.01<\*<0.05. OTUs are 414 colored according to their phylum. The positive and negative connectivities between OTUs are indicated by blue 415 and red lines, respectively. Only OTUs with a significantly correlated abundance profile with module are 416 represented in this figure.

417

#### 418 3.3.2. Archaeal and bacterial joining network

419 To further explore the microbial relationships with environmental parameters, we performed a 420 second eigengene analyses by first constructing a network using the 53 retained archaeal OTUs 421 from this study, and the 260 retained bacterial OTUs from the previous study using the same 422 samples and methodology (Nover et al. 2020). The network we obtained before considering 423 environmental parameters consisted in 200 nodes (OTUs) and 967 links and was composed of 424 21 modules. When considering environmental parameters, the network was composed of 136 425 OTUs, 442 links and 13 modules (Fig. 7, Table S3.2). All retained environmental dynamics 426 (Fig. 1b) were positively or negatively correlated with one or more modules and there were 427 127 bacterial and 9 archaeal significant OTUs. More than half of the total reads (53%) were 428 represented in module 1 (Fig. 7). Archaeal OTUs were present in five modules (Fig. 7). Six 429 archaeal OTUs represented 41% of the total reads among significant OTUs in the network and were linked to ProxyDyn2 and/or Diuron. Notably, OTU3 and OTU41 in module 1, OTU16, 430 431 OTU26, and OTU61 (Mba. acidurans, 98.42%) in module 2, and OTU47 (Mba. formicicum, 100%) in module 5 (Table S3.2). They were mainly present in t17, t19, t23, and t37 samples. 432 433 Two archaeal OTUs, 94 and 44, in module 4 correlated to  $NO_3^-$  (4% of the total reads, Fig. 7), 434 and belonged to Nitrosotaleaceae family (Crenarchaeota phylum) and were mainly present 435 towards the end of the flood event. Of the 127 bacterial OTUs present in this joined network, 49 were present in the bacterial only network from the previous study (Noyer et al. 2020, in 436 437 purple in Table S3.2) and were significantly linked to the same environmental dynamics. Worth 438 noting is that the bacteria in modules which were positively correlated with ProxyDyn2 and 439 Diuron were also the most relatively abundant in t17, t19, t23, and t37 samples. The most 440 abundant of these bacterial OTUs was OTU5, which matched Arcobacter cryaerophilus and 441 represented 20% of the total reads included in the network.

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444 Fig. 7 Summary of joined bacterial and archaeal network analysis. Each colored rectangle represents a module 445 whose number appears at the bottom left corner of each rectangle together with the percentage of reads in the 446 module out of the total number of reads analyzed within the network. The number of OTUs and the proportion of 447 reads within each module for each domain: archaea (A) and bacteria (B) are also indicated in each rectangle. At 448 the top left of each rectangle the environmental dynamics are indicated with module eigengene correlation value 449 between parentheses and the p-value significant code as follows: \*\*\*<0.001<\*\*<0.01<\*<0.05. The rectangle on 450 the bottom right corner represents the six modules whose percentage of reads is less than 1% of the total number 451 of reads analyzed in the network.

#### 452 **4. Discussion**

### 453 454

# 4.1. Particle-attached archaeome diversity at three different seasons was higher and more even than in all other lotic ecosystems studied so far

455 Archaea have been studied in only a few river environments (Herfort et al. 2009; Hu et al. 456 2016, 2018; Cannon et al. 2017; Samson et al. 2019; Lei et al. 2020; Cao et al. 2020; Pinto et 457 al. 2020; Shen et al. 2021) and to the best of our knowledge, no study has characterized their 458 diversity over different seasons in lotic ecosystems. Here, eDNA was extracted from riverine 459 water samples from summer, autumn, and winter and characterized using high-throughput 460 Illumina sequencing of the archaeal rDNA in a coastal Mediterranean watercourse, the Têt River (Southeast France). Shannon alpha diversity (Fig. 2) from summer, winter and autumn 461 drought samples were not significantly different (SD = 6.22; t0 = 6.25 and WD = 7.39). This 462 was also the case for all alpha diversity indices (Table S1). Lei et al. (2020) obtained lower 463 464 values of Shannon index (ranging 4.07-5.72) through Illumina sequencing of a highly polluted 465 river in China. Along a river in India sampled during summer and sequenced by metagenomic 466 analyses, Samson et al. (2019) found a Shannon diversity varying from 3.12 to 4.79, depending 467 on the sampling site. Hu et al. (2018) studied microbial diversity in a high-elevation river in 468 China using universal primers and found even lower Shannon values for archaea along the river 469 (2-2.7) and a Pielou evenness index very high (0.77-0.82) compared to ours (ranging 0.44-0.51 470 in Têt drought samples along seasons). These authors used 97% identity to define OTUs, while 471 the other two studies did not specify how they defined the OTUs in their studies. Nevertheless, 472 given the lower number of OTUs retained in these studies, the differences in alpha diversity 473 with respect to our study could be due to less throughput sequencing. Borrego et al. (2020) 474 studied particle-attached archaea in a Mediterranean water reservoir in summer with the same 475 specific primers as in the present study, and they also found a lower Shannon alpha diversity 476 than ours  $(4.9 \pm 0.15)$ . Gobet *et al.* (2014), in a comparative study of different alpha diversity 477 indices between ARISA versus pyrosequencing methodologies, determined significant correlation results among methods, particularly with Shannon index, without a need for 478 479 correction for sequencing depth. In summary, although the Têt River archaeal diversity was 480 higher and less even (lower Pielou index) when compared to other riverine environments 481 studied so far, additional studies with higher throughput and standardized sequencing pipeline 482 analyses are needed to better understand the ecology of fluviatile archaea.

483 Studies on lotic ecosystems have revealed the presence of three major phyla, Crenarchaeota, 484 Euryarchaeota and Thaumarchaeota using the previous silva database (Hu et al. 2016, 2018; 485 Samson et al. 2019; Lei et al. 2020). When considering archaeal taxonomy changes since the 486 last silva database release, the major phyla observed in the Têt river corresponded to those 487 same phyla, but we also found the Nanoarchaeota phylum (Fig. 3a). In a recent study, which 488 used the same primers as those used in our study, Woesarchaeia class (now called 489 Nanoarchaeia) was found in a small water reservoir sampled in summer (Borrego et al. 2020). 490 In our study, the summer drought community showed, through beta diversity analyses, the 491 greatest difference with respect to communities present in autumn and winter and was also 492 dominated by this class of the Nanoarchaeota phylum, followed by Bathyarchaeia class 493 (Crenarchaeota phylum). Primer bias could thus be responsible for the absence of this class in 494 previous riverine studies. Nevertheless, winter drought and dry autumn weather (t0) samples 495 had a smaller proportion of these classes at the expense of Nitrososphaeria class 496 (Crenarachaeota phylum, Fig. 3). On the other hand, Herfort et al. (2009) sequenced five 497 DGGE fragments amplified from surface waters of the Têt River in June 2005 that matched 498 two classes, a marine benthic group Crenarchaeota and LDS/RCV from Halobacteriales order (currently at Halobacteria class from Halobacterota phylum). In this study, we also found taxa from the genus *Methanomicrobia*, which were present in all samples independently of season, and are classified within Halobacterota and thus closely related to Halobacteria. While we also found taxa from the marine benthic group A from Crenarchaeota phylum (not shown), they were absent in the summer drought. Given the technical limitations of the DGGE fingerprinting method, a biased result cannot be discarded.

505 506

# 4.2. Particle-attached archaeome shifts gave evidence of allochthonous inputs into the Têt River along a Mediterranean extreme flood

507 In this manuscript, we characterized the temporal shift of the particle-attached riverine archaeal 508 community throughout an autumn storm event that included a 5-year flood and led to an instantaneous peak discharge of 270 m<sup>3</sup>/s. Cannon et al. (2017) is the only study that 509 510 characterized river archaeal composition in water samples from a contaminated urban river 511 before and after transient rainfall using Illumina sequencing of the rDNA. They found a weak 512 resolution in archaeal sequences of samples from before and after the rain event, which they 513 concluded might be due to a primer bias. In this study, we demonstrated major changes in 514 archaeal taxonomy, and therefore community shifts observed through beta diversity, occurred 515 at specific moments along the flood event and not when comparing samples from before and 516 after the flow peak. It is therefore possible that the absence of differences in Cannon et al. 517 (2017) was due to the resilience of archaeal communities at the end of the rainfall as was the 518 case at the Têt River. Communities from the end of the flood, from t44 to t109 were indeed 519 more similar to the t0 sample (sampled before the rainfall started, see Fig. 4). A first shift in 520 the structural diversity occurred after t17, particularly in samples t23 and t37. These samples corresponded to moments of multipollution peaks that occurred at storm first flushes CSOs and 521 522 inputs from in-sewer sediment resuspension (Fig. 1b, see also Reovo-Prats et al. 2017). 523 Methane-related classes such as Methanobacteria, Methanomicrobia and Methanosarcinia 524 became significantly abundant at these moments, and our results support these classes as major 525 components of riverine communities related to pollutants, organic matter inputs and/or hypoxic 526 sediments, as reported by Casamayor and Borrego (2009) with respect to the Euryarchaeaota 527 phylum. All these classes belonged indeed to the Euryarchaeaota before the last silva database 528 release. Lei et al. (2020) also found a predominance of different Euryarchaeota methanogens 529 in water samples from a black odor, highly polluted Chinese river and more particularly the 530 large presence of Methanobacterium genus (Methanobacteria class) in water as well as in hotter 531 and lower-oxygen, downstream sediments, what corroborates our hypothesis on the origin of 532 this river community shift. A second major community shift during the storm event occurred 533 at t41, just after the maximum water discharges (Fig. 3). Compared to Lei *et al.* (2020), who 534 did not observe a significant variation in alpha diversity along vertical and horizontal river 535 samples, we noticed a significant decrease in the archaea alpha diversity at t41 (Table S1). This 536 can be explained by the presence of three OTUs representing 90% of all reads in this sample 537 (not shown), which belonged to Nitrosopumilaceae and Nitrosophaeraceae families 538 (Nitrososphaeria class). Nitrososphaeria is the only known ammonia-oxidizing archaea (AOA) 539 class which accomplishes nitrification in all kinds of environments (Pinto et al. 2020) but has 540 been specifically found attached to terrestrial soil and freshwater sediments (Sonthiphand et al. 541 2013; Li et al. 2018). The AOA predominated assemblage at t41 is therefore derived from soil 542 runoff or sediments from upstream environments or from the resuspension of deep river 543 sediments, as they became predominant over polluted in-sewer sediments resuspension and 544 urban runoff-related assemblages that predominated until t38. The last community shift 545 included a significant enrichment of Thermoplamata class in samples t44 to t109, that 546 coincided with the start of the second peak of flow, which occurred due to the release of waters 547 from the upstream Vinca reservoir (Reovo-Prats et al. 2017). This class has been associated 548 with anoxic, sulphide-rich lentic sediments (Fillol et al. 2015; Compte-Port et al. 2017). The 549 release of water from this reservoir during regular floods is performed via a valve situated at 550 the bottom of the dam. As the bottom reservoir waters are anoxic (Fovet et al. 2020) and flow 551 increase at the bottom level of the dam could potentially lead to sediment release as well, both 552 are potential causes for the increase of Themoplasmata when dam discharges became 553 predominant at t44.

# 4.3. Major environmental forces were linked to particle-attached archaeome shifts during an extreme event through comprehensive modelling analyses

Among the few papers in the literature that have linked archaeal diversity to environmental parameters, all addressed the effect of one family of parameters on changes in archaeal communities, such as nutrients (Herfort et al. 2009; Hu et al. 2016, 2018; Lei et al. 2020; Cao et al. 2020) or metals (Mahamoud Ahmed et al. 2020; Shen et al. 2021). This is, therefore, the first study on the Archaea domain where both diversity and a large panoply of physicochemical parameters (notably nutrients, trace metals, pharmaceuticals, and pesticides) have been analyzed for the same samples. To analyze such complex datasets, we used a comprehensive 563 analysis as such performed on bacteria (Noyer et al. 2020). On one side, Bokulich et al. (2012) 564 suggested quality-filtering strategies to eliminate artifacts before interpretation of results and 565 recommended a conservative OTU threshold of 0.005% for bacteria. On the other side, it has 566 been suggested to use multivariate cut-offs instead of arbitrary thresholds to delineate abundant 567 versus rare OTUs, as the latest are largely dependent on sequence coverage (Jia et al. 2018). 568 Furthermore, multivariate cut-offs can be set considering environmental parameters when 569 available (Gobet et al. 2010). Based on these studies, we designed a strategy with the purpose 570 to define this cut-off at the point where the constraint-based model which considers OTU 571 relative abundance (tbRDA model) became significant. The environmental response of the 572 archaeal community was obtained with a tenfold higher threshold (0.05%) than bacteria 573 (0.005%, Nover et al. 2020). This result indicated that archaeal taxa that responded to 574 environmental pollutants were ten times more abundant than bacteria. Less abundant archaeal 575 taxa were therefore less crucial than those same bacterial taxa in the response of their respective 576 communities to environmental parameters. We believe that was the reason why the constraint-577 based model obtained when using Bray-Curtis dissimilarity i.e., that considers OTU abundance 578 only, turned out non-significant for the archaeome (Table 1a) in contrast to the bacteriome. The 579 other constrained multivariate analyses models that were significant, as well as the constrained 580 network analyses, gave evidence of two dynamics, ProxyDyn2 and Diuron, as responsible for 581 major structural diversity shifts observed on riverine particle-attached archaea during the 582 extreme flood event (Fig. 5 and 6b). These specific dynamics were discharged in the watershed 583 by CSOs as well as by urban runoff (Reoyo-Prats et al. 2017) from t17 to t38 and represented 584 several environmental pollutants including pesticides, as well as copper and dissolved 585 pharmaceutical products and a contaminant, phosphate (see Fig. 1b for further details). These 586 parameters have been found to affect bacterial communities in freshwater ecosystems, and 587 metals and nutrients also affect archaeal communities in freshwaters (Hu et al. 2016; Lei et al. 588 2020; Shen et al. 2021). Nevertheless, to our knowledge, the effect of xenobiotics such as 589 pesticides or pharmaceutical products on archaeal communities from freshwater ecosystems 590 has not yet been specifically reported. But, the pesticide glyphosate, for instance, interferes 591 with the aromatic-acids pathway in microorganisms, including archaea, and alters microbial 592 communities (van Bruggen et al. 2021). On the other hand, eutrophication by phosphate has 593 been found to decrease glyphosate degradation by biofilms and increase AMPA accumulation 594 in surface waters (Carles et al. 2019), what could therefore indirectly affect microbial 595 communities. The third significant dynamics according to the tbRDA, ProxyDyn1, projected 596 on axis 2, was mainly linked to the flow peak sample at t38, but axis 2 turned out not significant 597 (Fig. 5). ProxyDyn1 was neither correlated to the significant module of network analyses (Fig.598 6).

#### 599 4.4. Key players in the response of the riverine archaeome to multiple stressors

600 One of the major objectives of this study was to identify OTUs which could play a key role in 601 the archaeal community response to environmental changes derived from the delivery of 602 xenobiotics into the river by different pollutant sources. OTUs with positive membership to 603 the significant module from constraint-based network analysis were affiliated with 604 Euryarchaeota, Halobacterota and Thermoplasmatota (Fig. 6a, Table S3.1). These major OTUs 605 played a significant role in the response of the archaeome to multiple stressors derived from 606 point sources of pollution, as they were most abundant in samples t17-t23, t37, and t38 (Fig. 607 6b, Table S3.1) and were linked to ProxyDyn2 and Diuron but not to ProxyDyn1, which is a 608 proxy for diffuse sources of pollution. The predominant OTU11, affiliated with Mba. palustre, 609 is a species isolated for the first time in peat bogs that has the ability to use secondary alcohols 610 to produce methane (Zellner et al. 1988; Chaban et al. 2006). Mbt. smithii (OTU3) was the 611 second most abundant key player, which is known to be derived from the human 612 gastrointestinal tract (Miller et al. 1982; Oliveira et al. 2016) and identified as a potential 613 indicator of sewage (Johnston et al. 2010; McLellan and Eren 2014). The third most abundant 614 OTU41 was affiliated with *Msa. concilii*, which is involved in methane production from acetate 615 (Zwain et al. 2017) and is abundant in-sewer biofilms (Sun et al. 2014). What was striking in 616 the present study was that other OTUs, which had not yet been linked to CSOs and/or pollutant 617 inputs in the literature, acted as major OTUs in the response of the archaeome to pollutant 618 mixtures (Fig. 6a, Table S3.1). Three of them matched Methanobacteria (OTU26) or 619 Thermoplasmata (OTUs 16 and 19) classes and were also related to methane; and the other 620 three major OTUs were affiliated to Nanoarchaeia and Bathyarchaeia classes (OTUs 39, 66 621 and 34). Lastly, several OTUs affiliated to Thermoplasmata and Nitrososphaeria classes, had 622 a significant negative membership to the module of network analysis linked to ProxyDyn2 and 623 Diuron (Fig. 6c). These key players were instead linked to samples from the end of the flood 624 event, from t44 to t109. As evidenced above, they were taxa from allochthonous origins. All 625 these urban taxa, which were identified as either positively or negatively correlated with major 626 stressors, could be used as alternative bioindicators for rapid risk assessment of the impact of 627 multiple stressors on aquatic ecosystems, as proposed for bacteria (McLellan and Eren 2014; 628 Dila et al. 2018; Noyer et al. 2020).

# 629 4.5. Comparison between particle-attached archaeal and bacterial diversity at 630 seasons and along the flood event

631 Few studies so far have compared archaeal and bacterial structural diversity in lotic ecosystems 632 (Cannon et al. 2017; Hu et al. 2018) and, to the best of our knowledge, the differences in the 633 responses of both domains to seasons have not yet been studied. Fluviatile archaeal response 634 to environmental dynamics differed from the bacterial response in three ways. First, the 635 archaeome showed a specific community in summer compared to winter and autumn samples, 636 while bacterial communities from these three seasons clustered together when compared to 637 communities along the flood (Nover et al. 2020). Second, environmental forces structured the 638 archaeome diversity differently depending on beta diversity distance-based statistical models, 639 particularly when considering qualitative vs. quantitative dissimilarities. These models were 640 not always significant (Table 1), while for bacteria all models were significant. Third, while 641 the bacterial community structured differently at the two multipollution events of this extreme 642 flood, which were derived from CSOs and the flow peak (Reoyo-Prats et al. 2017), archaeal 643 structural shifts could only be interpreted after further statistical modeling of OTU abundances 644 and extreme event dynamics. On the one hand, the archaeal riverine resident communities 645 shifted significantly according to two environmental dynamics only, ProxyDyn2 and Diuron 646 (Fig. 5 and 6). These dynamics were derived from point sources of pollution, such as CSOs, 647 in-sewer sediments resuspension and urban runoff conducted through CSOs. On the other hand, 648 bacteria also responded significantly to other environmental parameters linked to diffuse 649 sources of pollution. Notably, at the highest water discharge at t38 and t41, bacteria showed a 650 specific community that was correlated to ProxyDyn1 and was thus attributed to allochthonous 651 inputs from runoff (Reoyo-Prats et al. 2017). The archaeome shifted at t41 into an anoxic 652 community, most likely related to the resuspension of deep sediments, and was not 653 significantly related to ProxyDyn1 (Fig. 3 and 5) and therefore the flow peak. In conclusion, 654 archaea can help to better understand the origin of watershed sediments and can thus be of 655 interest for quality assessments of suspended matter.

To further understand the different responses of archaea and bacteria domains of life to river contaminants and hydrodynamics, we used a network analysis combined with a module eigengene analysis of bacterial and archaeal taxa best fitted to significant beta diversity models. One remarkable finding was the joint presence of archaea *Mbr. smithii* and *Msa. concilii* with bacteria *Arcobacter cryaerophilus, Bacteroides graminisolvens, Cloacibacterium normanense* and *Macellibacteroides fermentans* in Module 1, which was significantly correlated with 662 ProxyDyn2 and Diuron. All of these species have already been identified as potential indicators of human-specific sewage pollution (Dick and Field 2004; Johnston et al. 2010; McLellan and 663 664 Eren 2014; McLellan and Roguet 2019) and A. cryaerophilus is a known human pathogen 665 (Collado et al. 2010). While no archaea has yet shown pathogenic effects on humans 666 (Cavicchioli et al. 2003; Bang and Schmitz 2018), archaea and bacteria can share genes through 667 horizontal gene transfer, which is particularly enhanced in anaerobic environments (Fuchsman 668 et al. 2017). Biofilms that develop in urban pipes are anaerobic (Guisasola et al. 2008), and the 669 co-habitation of archaea with bacterial pathogens in urban systems can increase the risk of 670 antibiotic resistance gene transfer. Sewers have indeed been identified as reservoirs of 671 antibiotic-resistance bacteria carried by human pathogens (Millar and Raghavan 2017; Auguet 672 et al. 2017). Similarly, resistance to pollutants could be enhanced between both domains 673 through the same mechanisms. Furthermore, the presence of both domains in the same habitat 674 pointed out to potentially common as well as complementary metabolic and physiological 675 functions. Another interesting finding that emerged from this analysis was the presence of 676 archaea together with bacteria OTUs in module 4, which significantly correlated with NO3<sup>-</sup> 677 (Fig. 7, Table S3.2). Notably, two Nitrososphaeria class OTUs, known to be dominant in 678 particle- attached ammonia-oxidizing archaeal communities (Cai et al. 2019; Pinto et al. 2020) 679 were linked to module 4. Wang et al. (2018) also demonstrated a significant influence of 680 dissolved inorganic nitrogen on the composition of bacterial and archaeal communities along 681 an urban river. In the present study, the addition of bacterial OTUs in the network analysis strengthened the importance of nitrates to drive shifts in archaeal assemblages, otherwise 682 683 mitigated when archaea were modeled alone, which enhance the interest of studying different 684 domains of life to better understand environmental drivers of community structure in natural 685 ecosystems.

#### 686 **5.** Conclusion

687 Our study provides the first overview of archaeal community shifts along an extreme storm 688 event that led to multiple pollutions in a typical coastal Mediterranean watercourse. Shifts were 689 also compared with changes in archaeal diversity during three seasons. Archaea from the Têt 690 river showed a specific community in summer compared to winter and autumn samples, and a 691 higher alpha diversity and lower evenness could be observed in this river compared to other, 692 yet less-thorough studies on riverine archaeal communities. Further studies on the spatio-693 temporal shifts in archaeal assemblages in these ecosystems are therefore urgently needed to 694 better understand seasonal shifts as well as their ecological diversity. For the first time, a 695 comparison of the response of archaea alone and together with bacteria in a fluviatile ecosystem 696 has shed light on the similarities and differences in their responses to seasons and when facing 697 multiple stressors derived from an extreme event. The fluviatile bacteriome and archaeome did 698 not respond in the same way to environmental forces. Extreme events were stronger at 699 structuring bacterial communities than seasons, while the opposite was observed in archaeal 700 communities. In contrast to bacteria, which responded quickly and significantly to both sewage 701 overflow and river hydrodynamics and associated environmental parameter changes, the 702 archaeal community shifted in response to multipollution derived from point sources and from 703 the resuspension of deep anoxic sediments but not so clearly at the river flow peak. Archaeal 704 taxa already known as urban-specific, as well as new archaea, mainly methane-related and 705 never identified as urban-specific taxa, predominated assemblages during multiple stress 706 events and were confirmed through statistical modeling of archaeal alone and together with 707 bacteria. These taxa could be used as bioindicators of point sources of pollution. Our results 708 highlight fluviatile archaea, seldom considered as bioindicators of water quality, could provide 709 a rapid risk assessment of multiple pollutants in aquatic ecosystems, as is the case for bacteria. 710 Furthermore, a better understanding of parallel shifts in assemblages from both domains of life 711 when confronted with multiple stressors could help to improve how urban watersheds are 712 monitored and would thus be extremely helpful in the management of pollution risk.

713

#### 714 Data Availability

715 Sequencing data are deposited on NCBI under BioProject ID PRJNA602803.

716

#### 717 Supplementary Information

Additional Supporting Information may be found in the online version of this article at thepublisher's website.

720

#### 721 Supplementary Information S1

Table S1. Alpha diversity indices at the Têt river with Kruskal-Wallis test results and graphical
 representation for each index.

724

#### 725 Supplementary Information S2

- 726 **Fig. S2.1.** Beta diversity based on additional dissimilarities.
- 727 Fig. S2.2. Jaccard, Unifrac and Weighted-Unifrac distance-based RDA triplot.

728	Supplementary Information S3
729	Table S3.1. Key player significant archaeal OTUs in module eigengene analysis significantly
730	related to environmental dynamics and graphical representation of their total reads along time.
731	Table S3.2. Key player significant bacterial or archaea OTUs in modules derived from module
732	eigengene analysis significantly related to environmental dynamics.
733	
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740	
741	Competing Interests
742	The authors have no relevant financial or non-financial interests to disclose.
743	
744	Contributions
745	Megane Noyer and Carmen Palacios contributed to conception and design of the study,
746	organized the database, performed the statistical analyses and wrote the first draft of the
747	manuscript. Maria Bernard helped with metabarcoding analyses. All authors contributed to
748	manuscript revision, read and approved the submitted version.
749	
750	Ethical Approval
751	Not applicable.
752	
753	Consent to Participate
754	All authors are informed and agree to the study.
755	
756	Consent to Publish
757	The authors declare no competing interests.

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Fig. 1 Têt River archaeome sampling sites and environmental parameters measured in the same samples. (a) Watershed of the river with sampling site (black arrow), located after wastewater treatment plan (WWTP) of the city of Perpignan, combined sewers (black crosses) and water reservoirs indicated as grey rectangles (adapted from Reoyo-Prats et al. 2017). (b) Environmental parameter dynamics in the Têt River at different seasons and along an autumn flood. For sample names, see table below figure. Autumn sample names are followed by a number that indicates the sampling time in hours after t0, which was sampled at autumn basal level water discharges. Sampling took part at crucial moments of the flood that occurred thereafter: at first flushes (t17-t19-t23), before the flow peak (t32-t37), during the flow peak (t38-t41), following the release of water from the upstream Vinça reservoir (t44-t61) and during the return to basal level (t68-t86-t109). ProxyDyn1 corresponds to the dynamics of particulate organic carbon (/20 mg/l), which represented the dynamics of water flow, also represented in figure, total suspended solids, total organic carbon, total nitrogen, and terbuthylazine parameters. ProxyDyn2 corresponds to aminomethyl phosphonic acid (AMPA,  $\mu g/l$ ), which represented glyphosate, phosphate, copper, temperature, E. coli, enterococci, diclofenac, sulfamethoxazole and carbamazepine parameters. ProxyDyn3 corresponds to lead  $(/150 \mu g/g)$  in the representation of the dynamics of cadmium, zinc, and conductivity parameters. ProxyDyn4 corresponds to pH (/70), which represented cobalt, nickel, and chrome parameters. Three parameters, Diuron, 2.4D and NO3-, had a unique dynamic. For further details on statistical analyses for environmental parameters, see Noyer et al. (2020).

**Fig. 2** Alpha diversity of the archaeome of the Têt river along time. Changes in observed OTU number (blue) and Shannon index (red) along the flood (tX) and at summer and winter droughts (SD and WD respectively). For the Shannon index, different letters indicate a significant difference between samples (dunn.test<0.025) and red arrows show major significant differences. Observed OTU number was not significantly different (KW=0.14). The dotted profile is the flow level at each sampling point (see Fig. 1, also for sample names). Even though the absence of replicates for t61 and t68 samples impeded statistical testing, they are represented through time for comparison.

**Fig. 3** Composition of archaeal communities averaged across replicates. Histogram of relative abundances (a) of the six major phyla and (b) of the ten major classes. Samples are organized according to sampling time from left to right: summer drought (SD), autumn flood (sample names are followed by a number that indicates the sampling time in hours after the beginning of the flood at t0), and winter drought (WD). The dotted profile is the flow level at each sampling point (see Fig. 1 for further details).

**Fig. 4** Structure of archaeal communities averaged across replicates. (a) Principal Coordinate Analysis (PCoA) and (b) hierarchical clustering with Ward D2 linkage method using Weighted-Unifrac dissimilarity computed on OTU average abundance. Lines indicate ANOSIM significant groups. Sample names are followed by a number that indicates the sampling time in hours after t0 (see Fig. 1 for details). Significant codes \*\* and \*\*\* indicate p-value < 0.01 and < 0.001, respectively.

**Fig. 5** Redundancy analysis (RDA) biplot with scaling by sites on the normalized matrix of OTUs with an abundance  $\geq 0.05\%$ . The model explained 66.32% of the variance (p<0.05). Significance for axes and environmental dynamics after permanova analyses are indicated, p-value significance codes: \*\*\*<0.001<\*\*<0.01<\*\*<0.01<\*\*<0.05. Sample names are followed by a number that indicates the sampling time in hours after the beginning of the flood at t0. For further details on sample names and retained environmental variable dynamics, see Fig. 1b. Perpendicular grey lines represent the projection of the corresponding samples onto the corresponding dynamics and approximate the value of that sample along the variable (Legendre and Legendre 2012).

**Fig. 6** (a) Molecular ecological network of the unique module significantly positively correlated with environmental dynamics, particularly ProxyDyn2 and Diuron (see Fig. 1b for details and sample names). (b) Histogram of total reads of OTUs with positive module membership in function of samples along the flood. (c) Same for OTUs with negative module membership. Environmental dynamics are followed by module correlation value between parenthesis and the p-value significant code as follows: \*\*\*<0.001<\*\*<0.01<\*\*<0.01<\*<0.05. OTUs are colored according to their phylum. The positive and negative connectivities between OTUs are indicated by blue and red lines, respectively. Only OTUs with a significantly correlated abundance profile with module are represented in this figure.

**Fig. 7** Summary of joined bacterial and archaeal network analysis. Each colored rectangle represents a module whose number appears at the bottom left corner of each rectangle together with the percentage of reads in the module out of the total number of reads analyzed within the network. The number of OTUs and the proportion of

reads within each module for each domain: archaea (A) and bacteria (B) are also indicated in each rectangle. At the top left of each rectangle the environmental dynamics are indicated with module eigengene correlation value between parentheses and the p-value significant code as follows: \*\*\*<0.001<\*\*<0.01<\*<0.05. The rectangle on the bottom right corner represents the six modules whose percentage of reads is less than 1% of the total number of reads analyzed in the network.

**Table 1** Summary of constraint-based multivariate statistical models on archaea OTU matrix averaged overreplicates and without singletons. (a) Permanova significance of the five models tested and the percentage ofbiological variance that is explained by each model using permutation test with anova.cca function. (b) Axes andmodeled variables significance after permanova using anova.cca of significant models in (a). p-values significancecodes: (\*\*\* < 0.001 < \*\* < 0.01 < \* < 0.05).</td>

Variance (%)		
66.35%		

b	OTUs matrix	Axes significance and variance explained (%)	Statistical significance of modeled variables						
	transformation	CAP1	ProxyDyn1	ProxyDyn2	ProxyDyn3	ProxyDyn4	2.4D	Diuron	NO <sub>3</sub> -
dbRDA	Jaccard	0.002** (22.74)	0.001***	0.002**	0.001***	0.105	0.095	0.001***	0.096
	Unifrac	0.040* (30.24)	0.001***	0.021*	0.046*	0.458	0.383	0.036*	0.014*
	Weighted-Unifrac	0.006** (42.89)	0.006**	0.009**	0.039*	0.050*	0.004**	0.002**	0.166

Supplementary Material 1

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