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PRIMARY RESEARCH PAPER

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Response of biofilm-dwelling nematodes to habitat changes 2 in the Garonne River, France: influence of hydrodynamics 3 and microalgal availability 4

Nabil Majdi · Walter Traunspurger · Stéphanie Boyer · Benoît Mialet · Michèle Tackx · Robert Fernandez · Stefanie Gehner · Loïc Ten-Hage · Evelyne Buffan-Dubau

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10 Abstract Lotic epilithic biofilms are submitted to seasonal disturbances (e.g. flood events, self-detach-11 12 ment), which influence the biomass, diversity and 13 viability of their algal and bacterial communities. The 14 objective of this study is to examine whether (1) 15 biofilm-dwelling nematodes respond to such seasonal 16 changes in terms of diversity and community struc-17 ture, (2) nematode species and feeding-types distri-18 bution respond to the varied trophic situations within 19 the biofilm, since variations in biofilm microalgal 20 composition may represent a variation in available 21 food. The biofilm-dwelling nematode community was 22 monitored in a temperate river over an 18 month 23 period with a high sampling frequency. These data 24 were linked to environmental abiotic and biofilm 25 biotic factors. Nematode density was positively 26 correlated to biofilm and microalgal biomass, but

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was dampened by floods. A clear seasonal pattern of 27 the community was detected (summer shift), so that 28 two nematode groups stand out: (1) the epistrate-29 feeders Chromadorina bioculata (Schultze in Carus, 30 1857) and Chromadorina viridis (Linstow, 1876) 31 were primarily related to diatom availability, and 32 dominated the nematode assemblage most of the 33 time, (2) seven species from various feeding types 34 (deposit-feeders, suction-feeders and chewers) grew 35 mainly under summer conditions concomitantly to a 36 change of biofilm trophic status and microalgal 37 composition. Overall, the results suggested that, in 38 addition to abiotic disturbances, the availability of 39 potential preys in the biofilm might represent an 40 important driver of nematode community patterns. 41

Keywords Nematodes · Periphyton · Diversity · 44 Feeding types · Algae · Environmental factors 43

Introduction

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In rivers, any hard submerged substrate can be coated 46 by a complex assemblage of organisms (e.g. bacteria, 47 fungi, algae, heterotrophic protozoans, meiofauna 48 and macrofauna) embedded in a mucous matrix of 49 exopolymeric substances (Costerton, 2000; Leflaive 50 51 et al., 2008). This organic layer which is named either epilithic biofilm, epilithon, 'Aufwuchs' or periphyton 52 can comprise more than 30% of microalgae in terms 53 of biomass (Peterson, 1996). Consequently, epilithic 54

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55 biofilms can constitute the main site of primary production in shallow water rivers harbouring hard 56 substrates such as the Garonne in its middle part 57 (Ameziane et al., 2003). These biofilms contribute 58 59 substantially to benthic food web functioning (Liess 60 & Hillebrand, 2004) and to biogeochemical processes such as decomposition and nutrient retention (e.g. 61 Ford & Lock, 1987; Battin et al., 2003; Teissier et al., 62 2007). However, epilithic biofilms are unstable 63 64 habitats, well-exposed to environmental perturba-65 tions. Hence they are strongly influenced by seasonal disturbances such as floods (Biggs & Close, 1989) 66 67 and self-detachment, a temperature-dependent bacterial degradation of the mat (Biggs, 1996; Boulêtreau 68 69 et al., 2006). These disturbances are recognized to shape the biomass, diversity and viability of the algal 70 and bacterial communities inhabiting the mat (e.g. 71 72 Peterson & Stevenson, 1992; Lyautey et al., 2010), 73 implying important consequences on the functioning 74 of biofilm processes (Cardinale, 2011).

75 Free-living nematodes are important protagonists 76 within biofilm communities: on the one hand, epilithic 77 biofilms represent both a habitat and a probable important food resource for them (e.g. Peters & 78 79 Traunspurger, 2005; Gaudes et al., 2006; Traunspurger 80 et al., 2006; Caramujo et al., 2008). On the other hand, 81 it has been suggested that nematode activity (e.g. 82 through bioturbation and grazing) could affect key 83 biofilm processes: for instance, Mathieu et al. (2007) 84 indicate that nematodes influence the oxygen turnover 85 of artificial diatom biofilms, and Sabater et al. (2003) 86 and Gaudes et al. (2006) highlight that meiofauna (mainly nematodes) can influence the release of 87 88 unpleasant odorous metabolites (e.g. geosmin) by cyanobacterial biofilms, implying high economic 89 90 relevance for fishing industry and drinking water 91 production.

92 Despite their important presence within these 93 habitats, biofilm-dwelling nematodes still remain 94 poorly considered as most nematological studies focus 95 rather on sediment-dwelling nematodes (Traunspurger 96 et al., 2006). As a matter of fact, most information on 97 biofilm-dwelling nematodes has issued from lentic 98 environments: e.g. spatial distributional patterns and 99 colonization pathways (Traunspurger, 1992; Peters & Traunspurger, 2005; Peters et al., 2005). So far, only 100 101 two previous studies have examined temporal distri-102 bution of biofilm-dwelling nematodes in running 103 waters during relatively short periods (Gaudes et al.,

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2006; Caramujo et al. 2008). But, long-term studies of104biofilm-dwelling nematodes are still lacking, which105hampers the assessment of how epilithic nematode106communities react and adapt to recurrent (seasonal)107abiotic disturbances and/or to fluctuations of food108resources over time.109

In this context, the questions put forward in this 110 study are: (1) In temperate areas, epilithic biofilms 111 are subject to seasonal temperature changes and 112 hydrological events, which, as mentioned above, 113 change their biomass and the composition of the algal 114 and bacterial communities. Is the biofilm-dwelling 115 nematode community influenced by such seasonal 116 changes of their habitat? (2) As variations in com-117 position of the microalgal community may represent 118 a variation in available food within the mat (in terms 119 of amount, availability and quality), do the nematode 120 species and feeding-types distribution match with the 121 biofilm trophic situation at a given time? With these 122 objectives, density, biomass, diversity, age, sex and 123 feeding types of the biofilm-dwelling nematode 124 community was monitored over an 18 month field 125 survey in a large temperate river: the Garonne (SW 126 France). These data were analysed to detect potential 127 seasonal changes, then the nematode species distri-128 bution was examined through the influence extent of 129 both environmental abiotic drivers and biofilm biotic 130 conditions. 131

Methods

Study site and sample collection	133
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The Garonne is the largest river of south-western 134 France with a drainage basin of 57,000 km² and a 135 length of 647 km. The Garonne River displays a 136 pluvio-nival flow regime with relatively short flash-137 floods caused by heavy rainfall (occurring mainly 138 between November and January) and a long annual 139 flood period due to snow-melt (April to June). In the 140 Garonne, alternate cobble bars are frequently found 141 even in channel up to the seventh-order. Between 142 floods (i.e. low-water periods), a high epilithic 143 biomass can grow on cobbles, being favoured by 144 low-water velocities on the river bed and low turbidity 145 (Boulêtreau et al., 2006). The study site was situated 146 on one of these cobble bars located at 36 km upstream 147

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Fig. 1 Location of the sampling site and cross-section view of the Garonne River at the sampling site

the city of Toulouse (01°17′53″E, 43°23′45″N; elevation 175 m a.s.l.), where the Garonne is of sixth-order
(Fig. 1).

151 Samplings (N = 51) were regularly performed from September 2008 to March 2010 when hydro-152 logical conditions permitted it (sampling was only 153 possible when discharge was lower than 175 m³ s⁻¹). 154 155 On each sampling occasion, 12 immerged cobbles (mean diameter: 10 cm) were collected underwater 156 157 using plastic bags to prevent any biofilm detachment 158 during removal. To consider water level changes and 159 depth where the biofilm typically develops (Amezi-160 ane et al., 2002), cobbles were collected on a cross-161 section from a reference point in the riverside so that 162 water height above cobbles remained between 30 and 163 50 cm. Collected cobbles were transported to the

laboratory within 2 h in cool boxes with minimal 164 disturbance. The biofilm was gathered by scraping the 165 upper surface of each cobble with a scalpel and a 166 toothbrush. Biofilm samples were finally suspended in 167 MilliQ water to obtain 12 biofilm suspensions (25 ml 168 each), in which algal aggregates were carefully crum-169 bled with scissors. These 12 biofilm suspensions were 170 used for the three following treatments: (1) nematode 171 species identification and density and biomass mea-172 surements, (2) HPLC analyses of microalgal pigments 173 and (3) epilithic ash-free dry mass (AFDM) measure-174 ments. Four replicate suspensions were used for each 175 treatment. Scraped cobbles were photographed, and 176 the surface of biofilm which had been removed 177 was clearly visible and measured using ImageJ soft-178 ware version 1.38 (Abramoff et al., 2004). Removed 179



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180 biofilm surfaces were then reported to corresponding

181 biofilm suspension volumes, so as densities, bio-

182 mass and pigment concentrations were quantitatively

183 expressed per area unit.

184 Nematode processing

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185 Nematodes were extracted from four replicate biofilm 186 suspensions using a modified gravity gradient centrifugation technique involving Ludox HS-40 after 187 188 Pfannkuche & Thiel (1988). Nematodes so extracted 189 were cleaned from Ludox by sieving through a 40 µm 190 sieve, then preserved in formaldehyde (5% final 191 concentration) and stained with 1% Rose Bengal. 192 Nematodes were counted in a Dolfuss cell (Elvetec 193 services, Clermont-Ferrand, France) under a Leica 194 MZ 9.5 stereomicroscope $(9 \times -90 \times)$ and their den-195 sity was expressed per cm². According to nematode 196 density, between 12 and 25 individuals were randomly picked up from each replicate while counting, 197 198 transferred to glycerol solution (Seinhorst, 1959), 199 mounted on slides and identified to the best species 200 level using a Leitz Dialux microscope at 1250× 201 magnification.

202 Nematodes were classified according to their age 203 (juveniles, fourth stage juveniles and adults), their 204 sexual category (females, gravid females and males), and their feeding type (epistrate-feeders, deposit-205 206 feeders, suction-feeders and chewers) after Traun-207 spurger (1997). The Maturity Index (MI) was 208 calculated on each sampling occasion as the weighted 209 mean frequency of individual colonizer-persister values (cp) after Bongers (1990). MI ranged from 1 to 5. 210 211 Nematode species with a cp = 1 were considered 212 r-strategists (colonizers) with short-generation times, 213 high fecundity and extreme population changes whereas those with a cp = 5 were defined as K-strat-214 215 egists (persisters) with lower breeding efficiency. The 216 MI is expected to decrease during disturbed periods, 217 when opportunistic nematodes are favoured (Bongers & Bongers, 1998). Over a 1-year period from 218 219 September 2008 to September 2009 (N = 37), at least 220 100 individual nematode body dimensions (length and 221 maximum width) were measured on each sampling 222 occasion from microscopic pictures taken while 223 counting. Mean individual wet weight (WW) was 224 then determined after Andrássy (1956).

Abiotic environmental factors

Mean Daily Discharge (MDD) was supplied by a 226 gauging station of the French water management 227 authority (DIREN Midi-Pyrénées, Marquefave sta-228 229 tion) located at 10 km upstream the study site-with no tributary and no dam between the gauging station 230 and the study site. The Mean Weekly Discharge 231 (MWD) before each sampling occasion was consid-232 ered in statistical analysis. To better reflect the effect 233 of flood disturbance, days after flood (DAF), which 234 were effective days after the last flood (MDD > 235 $300 \text{ m}^3 \text{ s}^{-1}$), were calculated for each sampling 236 occasion and considered in statistical analysis. Water 237 temperature, conductivity, pH and dissolved oxygen 238 concentration were measured every 30 min during the 239 whole study period with an automated multi-param-240 eter probe (YSI 6000, YSI inc., Yellow springs, OH, 241 USA) which was permanently settled at 5 cm above 242 the streambed at the study site. 243

Biofilm microalgal composition and biomass 244

Microalgal pigments extraction and HPLC-analysis 245

On each sampling occasion, four replicate biofilm 246 suspensions were centrifuged (3,220 g, 20 min). 247 Pellets were freeze-dried and thoroughly homoge-248 nized. Then, 250 mg aliquots were removed from 249 each pellet. Algal pigments from each pellet aliquot 250 were then extracted three times (15 min at -20° C) 251 with a total of 25 ml (10, 10 and 5 ml) 98% cold-252 buffered methanol (with 2% of 1 M ammonium 253 acetate) following Buffan-Dubau & Carman (2000b). 254 255 Algal pigment release was favoured at each extraction step by an ultrasonication probe (Sonifier 250A, 256 Branson Ultrasonics corp., Danbury, CT, USA). 257

One millilitre of the pigment solution so obtained 258 was then filtered on 0.2 µm PTFE syringe filter and 259 analyzed using a high-performance liquid chromato-260 graph (HPLC) consisting of a 100 µl loop auto-261 sampler and a quaternary solvent delivery system 262 coupled to a diode array spectrophotometer (LC1200 263 series, Agilent Technologies inc., Santa Clara, CA, 264 USA). The mobile phase was prepared and pro-265 grammed according to the analytical gradient protocol 266 described in Barlow et al. (1997). Pigment separation 267 was performed through a C8, 5 µm column (MOS-2 268



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Table 1 CHEMITAA pignent failo matrix										
Algal group	Species	Biomarker pigment ratios to Chl a								
		Fuco	Lut	Viola	Diad	Zea	β -car	Chl a	Chl b	Chl c
Green algae	P. borianum		0.143	0.049		0.014	0.043	1	0.088	
Diatoms	N. palea	0.477			0.102		0.002	1		0.121
Cyanobacteria	S. leopoliensis					0.411	0.011	1		

 Table 1
 CHEMTAX pigment ratio matrix

Ratios were calculated considering the relative concentrations of fucoxanthin (Fuco), lutein (Lut), violaxanthin (Viola), diadinoxanthin (Diad), zeaxanthin (Zea), β -carotene (β -car), chlorophyll *b* (Chl *b*) and chlorophyll *c* (Chl *c*) versus chlorophyll *a* (Chl *a*) concentrations from corresponding microalgal cultures. For green algae and diatoms these ratios were obtained from pure cultures of, respectively, *Pediastrum boryanum* and *Nitzschia palea*. For cyanobacteria, pigment ratios were obtained from *Synechococcus leopoliensis* (Schlüter et al., 2006)

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269 HYPERSIL, Thermo Fisher Scientific inc., Waltham, 270 MA, USA). The diode array detector was set at 440 nm 271 to detect carotenoids, and at 665 nm to detect chloro-272 phylls and pheopigments (Wright et al., 1991). Data 273 analysis was performed using ChemStation soft-274 ware (version A.10.02, Agilent Technologies inc.). 275 Pigments were identified by comparing their retention 276 time and absorption spectra with those of pure 277 standards pigments (DHI LAB products, Hørsholm, 278 Denmark). Each pigment concentration was calculated 279 by relating its chromatogram's peak area with the 280 corresponding area of calibrated standard.

281 Microalgal cultures and chemotaxonomy

282 Algal pigment analysis by HPLC coupled with 283 chemotaxonomic analysis using CHEMTAX program 284 (Mackey et al., 1996) has proven to be a fast and 285 precise method to determine the biomass of phyto-286 planktonic and microphytobenthic groups in marine and freshwater environments (e.g. Schlüter et al., 287 288 2006; Caramujo et al., 2008; Lionard et al., 2008). As 289 reported by Leflaive et al. (2008), microalgal groups 290 inhabiting epilithic biofilms of the Garonne River are 291 diatoms, green algae and cyanobacteria. The bio-292 marker pigment composition found in the biofilm can 293 be used to estimate the biomass of each of these 294 microalgal groups by chemotaxonomy. Prior to the 295 chemotaxonomic analysis, biomarker pigment ratio 296 to chlorophyll a (Chl a) for each microalgal group 297 has to be obtained. Thus, a green algae species, 298 Pediastrum boryanum (Turpin) Meneghini (strain 299 Pedbo01) and a diatom species, Nitzschia palea 300 (Kützing) W. Smith (strain Nitpa01) were isolated 301 from the biofilm of the Garonne River and main-302 tained on Combo medium (Kilham et al., 1998) at

18°C (light:dark 16:8, 45 μ mol m⁻² s⁻¹). An aliquot 303 of each algal culture (10 mL) was filtered on 304 0.7 µm glass fibre filter (GF/F, Whatman, Clifton, 305 NJ, USA) and algal pigments were extracted and 306 analysed from the filters following the same procedure 307 than biofilm samples. Concerning cyanobacteria, pig-308 ment ratios calculated by Schlüter et al. (2006) for 309 Synechococcus leopoliensis (Raciborski) Komrek 310 (University of Toronto Culture Collection strain 102) 311 were considered. 312

The biomarker pigment ratio to Chl a so obtained 313 were used to supply the initial matrix needed for 314 CHEMTAX analysis (Table 1). Then, CHEMTAX 315 version 1.95 software (Mackey et al., 1996) was run 316 to estimate the biomass of diatoms, green algae and 317 cyanobacteria which were expressed as Chl a equiv-318 alents and considered as environmental biotic factors 319 320 in further statistical analysis.

Total epilithic biomass and autotrophic index 321

On each sampling occasion, four biofilm suspensions 322 were dried at 105°C for 18 h, weighted and then 323 combusted at 450°C for 8 h to weight the ash-free dry 324 mass (AFDM) of the biofilm. The Autotrophic Index 325 (AI) was determined as the ratio AFDM/Chl a. This 326 index is commonly used to describe the trophic status 327 of biofilm communities, e.g. higher AI values are 328 found in biofilms with higher proportions of hetero-329 trophs and/or organic detritus (Biggs & Close, 1989). 330

Statistical analysis

To investigate seasonal changes of the nematode 332 community structure, the differences in biomass, 333 diversity, age, sex, feeding types and MI were 334



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335 analysed between samples assigned to their corresponding sampling season (i.e. summer: 21 June-21 336 337 September, N = 15; autumn: 21 September-21 338 December, N = 18; winter: 21 December–21 March, 339 N = 15 and spring: 21 March-21 June, N = 3). The 340 homogeneity of variance was assessed with Levene's 341 test, and differences were examined either by one-342 way ANOVA followed by a post-hoc Tukey HSD test or by Kruskal-Wallis ANOVA. The same statistical 343 344 procedures were applied to investigate seasonal 345 changes of biofilm and microalgal biomass. The correlations between total nematode density and 346 347 biotic and abiotic factors were investigated by 348 Spearman's rank correlation test. These tests were 349 performed with STATISTICA software (version 8.0, 350 Statsoft inc., Tulsa, OK, USA).

The influence of biotic and abiotic environmental 351 factors on the nematode species distribution was 352 353 analyzed through canonical ordination analysis with 354 CANOCO software (version 4.5, Biometris, Wagen-355 ingen, The Netherlands). Rare species (with relative 356 occurrence <0.1%) were not considered in this anal-357 ysis. Species densities were square-root transformed 358 prior to the analysis. The distribution of nematodes 359 was first analyzed by a detrended correspondence 360 analysis (DCA). As the total inertia observed was less 361 than 2.6, a predominance of linear species response 362 curves could be expected (Ter Braak, 1987, 1994). 363 Therefore, a redundancy analysis (RDA) in which the 364 ordination axes were constrained to be linear combi-365 nations of provided environmental factors was used to 366 investigate the relationships between these factors and

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the distribution of main nematode species. Environ-367 mental factors were also listed (conditional effects) 368 according to the variance they explained singly (i.e. 369 without eventual co-variability with other factors). 370 The statistical significance was tested with Monte 371 Carlo permutation test (499 unrestricted permutations) 372 with applying Bonferroni's correction (significance 373 level set at P < 0.005). 374

Results

Dynamics of the epilithic biofilm

The range and annual mean values of each measured 377 abiotic and biotic factor are listed in Table 2. AFDM 378 and Chl a content of the epilithic biofilm were 379 significantly positively correlated (Spearman rank: 380 R = 0.75; P < 0.001) and showed considerable vari-381 ations throughout the sampling period, being partic-382 ularly dampened after floods (Fig. 2a). The AI was 383 significantly higher during summer than during the 384 other seasons (ANOVA: F = 60.2; P < 0.001), 385 implying globally a lower availability of microalgae 386 within summer biofilm communities. Diatoms dom-387 inated the epilithic microalgal assemblage over the 388 whole sampling period (Fig. 2b, Table 2). The dia-389 tom biomass was significantly higher during winter 390 than during the other seasons (ANOVA: F = 16.1; 391 P < 0.001). Conversely, cyanobacterial biomass was 392 significantly higher during summer (ANOVA: F =393 4.6; P < 0.01), and green algal biomass was 394

Table 2 Measured abioticand biofilm biotic factors		Annual mean \pm SE	Min	Max
	Temperature (°C)	14.6 ± 0.05	1.7	27.3
	$O_2 (mg l^{-1})$	11.5 ± 0.02	7.4	22.1
	pH (-)	7.6 ± 0.004	6.7	9.1
Annual means refer to 2009.	Conductivity (μ S cm ⁻¹)	270.9 ± 0.001	154	493
For temperature, O ₂ , pH	Mean daily discharge $(m^3 s^{-1})$	124.7 ± 6.0	18	814
and conductivity	Days after flood (day)	89.4 ± 11.1	7	233
(N = 1/50'). For days after flood and the biotic factors	AFDM (g m^{-2})	27.4 ± 2.7	4.4	79.7
(N = 31). Minimum and	Chlorophyll $a \ (mg \ m^{-2})$	321.5 ± 50	10.7	1012.8
maximum values refer to	Green algae (%)	17.1 ± 2.3	0	36.3
the whole sampling period	Cyanobacteria (%)	2.2 ± 0.6	0	14.6
(i.e. September 2008– March 2010)	Diatoms (%)	80.7 ± 2.7	50.6	100





Fig. 2 Temporal dynamics of **a** epilithic chlorophyll *a* (Chl *a*) concentration (\pm SE, *N* = 4), ash-free dry mass (AFDM) of the biofilm (\pm SE, *N* = 4) and mean daily discharge (MDD), and **b** the relative proportion (%) of epilithic microalgal groups to

395 significantly higher during summer and autumn 396 (ANOVA: F = 2.8; P < 0.05) than during the 397 remainder of the year.

398 Dynamics of biofilm-dwelling nematodes

399 Over the whole study period, the nematode density averaged 25.4 ± 4.3 ind cm⁻² and varied greatly 400 throughout the year: the lowest density (0.36 \pm 401 $0.14 \text{ ind } \text{cm}^{-2}$) occurred in early summer 2009 402 whereas the highest density $(161.36 \pm 52.5 \text{ ind } \text{cm}^{-2})$ 403 404 was attained during late winter 2010. As AFDM and Chl a, the nematode density was clearly dampened 405 406 after flood events (Fig. 3a). Nematode density was positively correlated with DAF (Spearman rank: 407

total Chl *a* biomass (N = 4). Months, years, seasons and floods during which MDD > 300 m³ s⁻¹ (represented by *stars*) are indicated on the *X* axis

R = 0.36; P < 0.01), AFDM (Spearman rank: R = 4080.41; P < 0.01) and Chl a (Spearman rank: R = 0.47;P < 0.001). From September 2008 to September 2009,410the nematode individual wet weight averaged 0.3 µg.411The individual biomass was significantly lower during412summer (ANOVA: F = 14.1; P < 0.001) than during413the other seasons (Fig. 3a).

From the 2,875 nematodes identified, 28 species 415 belonging to 11 families were found (see species list 416 in Table 3). Two species: Chromadorina bioculata 417 and Chromadorina viridis (family Chromadoridae) 418 strongly dominated the assemblage accounting for 419 86% of all identified nematodes. Although the family 420 Monhysteridae-particularly with species Eumonhys-421 tera dispar, Eumonhystera vulgaris and Monhystrella 422

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Fig. 3 Temporal dynamics of **a** nematode density (\pm SE, N = 4) and individual wet weight (WW) biomass (\pm SD, $N \ge 100$), **b** relative density of main nematode taxa, and **c** Maturity index (MI) and species richness (S) in the epilithic

biofilm. Months, years, seasons and floods during which MDD > 300 m³ s⁻¹ (represented by *stars*) are indicated on the X axis

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Table 3 Biofilm-dwelling nematode species in the study site between September 2008 and March 2010

CHROMADORIDA Filipjev, 1929 Chromadoridae Filipjev, 1917 Chromadorina bioculata (Schultze in Carus, 1857) 68. Chromadorina viridis (Linstow, 1876) Plectidae Örley, 1880 Plectus opisthocirculus Andrássy, 1952 0. Plectus aquatilis Andrássy, 1985 0. Plectus rhizophilus de Man, 1880	87 3 .15 3 .59 2 .14 2 1 2	E
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Plectus aquatilis Andrássy, 19850.Plectus rhizophilus de Man, 1880<0.	.14 2	D
Plectus rhizophilus de Man, 1880	1	D
	- 2	D
Plectus cirratus Bastian, 1865 <0.	.1 2	D
Prismatolaimidae Micoletzky, 1922		
Prismatolaimus cf. intermedius (Bütschli, 1873) <0	.1 3	E
Rhabdolaimidae Chitwood, 1951		
Rhabdolaimus aquaticus de Man, 1880 <0.	.1 3	D
MONHYSTERIDA Filipjev, 1929		
Monhysteridae de Man, 1876		
Eumonhystera dispar (Bastian, 1865)	.92 2	D
Eumonhystera vulgaris (de Man, 1880)	.84 2	D
Eumonhystera simplex (de Man, 1880)	.35 2	D
Eumonhystera barbata Andrássy, 1981	.31 2	D
Eumonhystera cf. filiformis (Bastian, 1865) <<0	.1 2	D
Eumonhystera longicaudatula (Gerlach & Riemann, 1973) </td <td>.1 2</td> <td>D</td>	.1 2	D
<i>Eumonhystera</i> sp. <0	.1 2	D
Monhystrella paramacrura (Meyl 1954)	.04 2	D
DORYLAIMIDA Pearse, 1942		
Dorylaimidae de Man, 1876		
Mesodorylaimus cf. subtiliformis (Andrássy, 1959)	.04 4	S
Mesodorylaimus sp. <0	.1 4	S
Eudorylaimus sp. <0	.1 4	S
Dorylaimus stagnalis Dujardin, 1845	.1 4	S
Mermithidae Braun, 1883		
Mermithidae <0	.1 1	Р
ENOPLIDA Filipjev, 1929		
Tobrilidae Filipjev, 1918		
Brevitobrilus stefanskii (Micoletzky, 1925) 0	.56 3	С
Tobrilus gracilis (Bastian, 1865) <0	.1 3	С
Tripylidae de Man, 1876		
Tripyla cf. filicaudata de Man, 1880 <0	.1 3	С
Tripyla glomerans Bastian, 1865 <0	.1 3	С
Alaimidae Micoletzky, 1922		
Paramphidelus sp. <0	.1 2	D
TYLENCHIDA Thorne, 1949		
Aphelenchoididae Skarbilovich, 1947		
Aphelenchoides sp. 0	.24 2	S
Tylenchidae Örley, 1880		

(H)

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Table 3 continued			
Nematode taxa	%	ср	FT
Coslenchus sp.	<0.1	3	S



paramacrura-represented only 10% of all identified 423 nematodes over the whole period, they clearly 424 425 dominated the assemblage from mid-July to mid-426 August (Fig. 3b). Sixteen species were rare, account-427 ing for <0.1% of all identified nematodes (Table 3). 428 The species richness (S) varied from 2 to 12 species 429 averaging S = 4.23 over the whole study period. S was significantly higher during summer (ANOVA: 430 F = 6.5; P < 0.001) than during the other seasons. 431 Conversely, the Maturity Index (MI) was signifi-432 433 cantly lower (MI = 2.67) during summer (Kruskal-Wallis ANOVA: H = 31.5; P < 0.001) than during 434 435 the other seasons. This summer shift in S and MI is 436 illustrated in Fig. 3c.

437 Epistrate-feeders-mainly represented by C. bioc-438 ulata and C. viridis-dominated representing 86% of 439 nematodes identified over the whole sampling period. 440 Deposit-feeders were the second most observed 441 group representing 12% while suction-feeders and chewers were less common representing, respec-442 443 tively, 1.5 and 0.5%. Insect parasites (i.e. Mermi-444 thidae) represented <0.1%. During summer, the 445 epistrate-feeders were significantly less represented 446 (ANOVA: F = 28.5; P < 0.001) while deposit-feeders were significantly more represented (Kruskal-447 Wallis ANOVA: H = 38.7; P < 0.001) than during 448 449 the other seasons (Fig. 4a).

450 The seasonal proportion of juveniles, fourth stage 451 juveniles, females, gravid females and males is 452 presented in Fig. 4b. Concerning the age structure 453 of the community, adult nematodes averaged 70% of 454 all identified nematodes, while fourth stage juveniles 455 and early instar juveniles contributed, respectively, to 456 14 and 16%. Early instar juveniles were significantly 457 more represented during spring (ANOVA: F = 2.8; 458 P < 0.05) than during the other seasons. Concerning 459 the sex structure of the community, females repre-460 sented 28% (non-gravid females) and 14% (gravid 461 females) against 28% for males. Males contributed

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Fig. 4 Seasonal variations of the nematode community structure in the biofilm: **a** seasonal proportion of epistrate-feeders (E), deposit-feeders (D), suction-feeders (S) and chewers (C), and **b** seasonal proportion of males (\mathcal{J}), females (\mathcal{G}), gravid females ($g\mathcal{G}$), fourth stage juveniles (J4) and juveniles (J)

significantly less during summer (ANOVA: $F = 3.2$; 462
P < 0.05) than during winter.	463

Influence of environmental factors on nematode464species distribution465

The results of the redundancy analysis (RDA) testing466the influence of biotic and abiotic factors on nematode467species and feeding-types distribution are presented in468Fig. 5 and Table 4. The temporal distribution of469nematode species was significantly influenced by470

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Fig. 5 Biplot from the redundancy analysis (RDA) explaining the distribution of nematode species densities according to environmental factors. Ordination axes were rescaled to range from -1 to 1. Slim dotted arrows are non-significant factors. Bold arrows are significant factors (Monte Carlo permutation test with Bonferroni's correction, P < 0.005). The eigenvalues (λ) are indicated for main ordination axes. Environmental factor abbreviations: biomass of diatoms (Diatoms), green algae (GreenAlg) and cyanobacteria (Cyano), epilithic ash-free

471 temperature, AFDM, DAF and biomass of cyanobac-472 teria, green algae and diatoms. The sum of all 473 significant factor eigenvalues explained 64.1% of the 474 variance. This analysis allowed to clearly distinguish 475 two groups of nematode species: The first group 476 comprised the two dominant epistrate-feeder species C. bioculata and C. viridis. These two species are 477 478 situated along axis 1, scoring towards the middle right 479 side of the biplot. Since axis 1 involved mainly factors 480 AFDM, DAF and diatom biomass, this indicated that 481 both species were more abundant during prolonged 482 undisturbed periods with a high biofilm and diatom 483 biomass. The second group comprised deposit-feeders 484 (i.e. Eumonhystera dispar, E. vulgaris, E. barbata, 485 Plectus aquatilis and Monhystrella paramacrura), 486 suction-feeders (i.e. Mesodorylaimus cf. subtilifor-487 mis and Aphelenchoides sp.) and chewers (i.e.

dry mass (AFDM), water temperature (T), pH, dissolved O₂ (O₂), conductivity (Cond), mean weekly discharge (MWD) and days after flood (DAF). Nematode species abbreviations: Aphelenchoides sp. (Asp), Chromadorina bioculata (Cbi), C. viridis (Cvi), Eumonhystera barbata (Eba), E. dispar (Edi), E. simplex (Esi), E. vulgaris (Evu), Brevitobrilus stefanskii (Bst), Monhystrella paramacrura (Mpar), Mesodorylaimus cf. subtiliformis (Msub), Plectus aquatilis (Paq) and P. opisthocirculus (Pop)

Brevitobrilus stefanskii). These species are distributed 488 along axis 2, scoring towards the upper part of the 489 biplot (except for *P. aquatilis*). Since axis 2 involved 490 491 mainly factors temperature and biomass of cyanobacteria and green microalgae, and since these both 492 microalgal groups were significantly more represented 493 during summer, this indicated that these nematode 494 species were more abundant under summer conditions. 495 No clear trend was observed for the distribution of 496 Plectus opisthocirculus and Eumonhystera simplex. 497

Discussion

To the best of our knowledge, the present study is the 499 first long-term monitoring of nematode assemblages 500 inhabiting lotic epilithic biofilms. Although the 501



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Factors	λ	Р
Diatoms	0.149	0.002**
Т	0.138	0.002**
DAF	0.104	0.002**
AFDM	0.102	0.002**
Cyano	0.084	0.004**
GreenAlg	0.064	0.004**
Cond	0.015	0.122
pН	0.013	0.154
MWD	0.006	0.502
O ₂	0.003	0.786

Each environmental factor is listed by its eigenvalue (λ) indicating the importance of its own contribution (i.e. without co-variabiliy, see "Methods") to explain the distribution variance of nematodes species. Significant factors (**) at P < 0.005 (see "Methods"). Biomass of diatoms (Diatoms), green algae (GreenAlg) and cyanobacteria (Cyano), epilithic ash-free dry mass (AFDM), water temperature (T), pH, dissolved O₂ (O₂), conductivity (Cond), mean weekly discharge (MWD) and days after flood (DAF)

502 biofilm-dwelling nematode community was not 503 diversified, two groups of species showing different 504 dynamics were clearly distinguished and seemed to 505 adapt to biofilm composition and seasonality: the first 506 group, consisting of the strongly dominating Chrom-507 adorina bioculata and C. viridis, was mainly related 508 to biofilm composition (i.e. age, thickness and diatom 509 content) whereas the second group of species mainly 510 grew under summer conditions.

The nematode density averaged 25.4 ind cm^{-2} and 511 ranged from 0.4 to 161.4 ind cm^{-2} in the epilithic 512 biofilm over the whole study period. This result lies 513 within the range of values reported for lake epilithic 514 biofilms, i.e. 2.8-161.5 ind cm⁻² (Peters & Traun-515 spurger, 2005) and for river epilithic biofilms, i.e. 516 10–100 ind cm^{-2} (Gaudes et al., 2006). In our study, 517 the nematode community constituted a permanent 518 519 component of river epilithic biofilms. Mathieu et al. 520 (2007) suggested that nematode activity could affect the oxygen turnover of diatom biofilms at density 521 values >50 ind cm⁻². This threshold value of density 522 523 was reached on several occasions during the study 524 period suggesting that this influence was substantial in 525 the epilithic biofilms of the Garonne River.

526 Nematode density positively correlated with 527 AFDM and Chl *a*. This strengthens the hypothesis Hydrobiologia

that the amount of microalgae and organic matter 528 favour meiobenthic organisms-such as nema-529 todes-in epilithic biofilms (Hillebrand et al., 2002; 530 Peters & Traunspurger, 2005). However, nematode 531 density and biofilm biomass were both clearly 532 dampened after floods (Figs. 2a, 3a). Moreover, the 533 positive relation found between nematode density and 534 DAF pointed out the negative impact of floods on 535 nematode populations. It is well-known that epilithic 536 biofilms are detached by shear stress, substratum 537 instability and abrasive effects of suspended solids 538 during flood events (Biggs & Close, 1989; Boulêtreau 539 et al., 2006). It is thus obvious that nematodes were 540 swept away with the biofilm when flood occurred. 541 542 This corroborates the studies of Robertson et al. (1997) and Palmer et al. (1996) showing that floods 543 are important factors shaping meiobenthic commu-544 nities in rivers. 545

The species richness observed in the present study 546 (i.e. 28 species over the whole study period) agreed 547 with those observed for several lake epilithic bio-548 films, i.e. 29 and 8-34 species (in, respectively, 549 Traunspurger, 1992; Peters & Traunspurger, 2005). 550 However, higher species richness values were often 551 reported for sediment-dwelling nematodes (see 552 review of Traunspurger, 2002). As previously shown 553 in lakes (Peters & Traunspurger, 2005), our results 554 suggest that, also in rivers, nematode diversity is 555 lower in biofilms than in sediments. Reasons for this 556 diversity difference remain complex and unclear 557 (Hodda et al., 2009). A possible explanation might 558 be that, in the Garonne river, nematodes had to totally 559 re-colonize the biofilm after critical floods several 560 times a year (e.g. in January, April-May and 561 November 2009, Fig. 3a). Conversely, in sediments, 562 meiobenthic organisms can migrate deeper towards 563 less disturbed sediment layers to shelter against 564 increasing discharge conditions (Dole-Olivier et al., 565 1997). Thus, biofilm-dwelling nematodes could be 566 more exposed than sediment-dwelling nematodes to 567 flood disturbances, which are known to decrease 568 benthic invertebrate diversity (Death & Winterbourn, 569 1995). 570

While diatoms dominated biofilm algal assem-
blages in terms of biomass, two epistrate-feeder571species Chromadorina bioculata and Chromadorina573viridis dominated strongly the nematode assemblage.574This observation supports the trend previously hypoth-
esized that, in freshwater benthic environments,576

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577 nematode communities are generally dominated by few species (e.g. Zullini & Ricci, 1980; Michiels & 578 579 Traunspurger, 2005a; Peters & Traunspurger, 2005). 580 Furthermore, this corroborates a previous study indi-581 cating that the epistrate-feeder Chromadorita leuck-582 arti (de Man, 1876) dominates the nematode 583 assemblages in diatom-dominated biofilms of the 584 Llobregat River, Spain (Gaudes et al., 2006). C. bioculata and C. viridis were clearly segregated from 585 586 the other nematode species (Fig. 5) and primarily positively related to diatom biomass. Due to their high 587 content of polyunsaturated fatty acids (Phillips, 1984), 588 589 diatoms are known to represent a high-quality food 590 resource often selected by benthic primary consumers 591 (e.g. Goedkoop & Johnson, 1996; Buffan-Dubau & 592 Carman, 2000a). Furthermore, it has been evidenced 593 that a marine nematode belonging to the Chromado-594 rina genus: Chromadorina germanica (Bütschli, 595 1874) feeds on benthic diatoms (e.g. Tietjen & Lee, 596 1977; Deutsch, 1978). Therefore, it is likely that the 597 presence of large amounts of a potential food resource 598 may favour C. bioculata and C. viridis. This finding 599 strengthens that nematode feeding strategies match 600 with the availability of their preys within the biofilm.

601 Our results indicate that a clear shift of the nem-602 atode community occurred during summer (Fig. 3b). 603 Such seasonal variations of species composition were previously reported for sediment-dwelling nematode 604 605 communities in lakes (Traunspurger, 1991; Michiels 606 & Traunspurger, 2005c) and in rivers (Beier & 607 Traunspurger, 2003). In our study, the summer nem-608 atode community is more diversified with a higher 609 proportion of deposit-feeders: e.g. Monhysteridae 610 (Figs. 3c, 4a). Concomitantly, the proportion of mic-611 roalgae in the biofilm (AI) was reduced, but the 612 microalgal community became more diversified. Sev-613 eral hypotheses can be advanced to account for this 614 summer shift:

615 Firstly, the RDA analysis (Fig. 5) evidenced that a diversified group of nematode species (mainly 616 deposit-feeding species) grew under summer condi-617 618 tions. It is known that summer temperatures enhance 619 the proportion of diversified bacterial assemblages 620 inside epilithic biofilms of the Garonne River (Boul-621 êtreau et al., 2006; Lyautey et al., 2010). Deposit-622 feeding nematodes can show species-specific feeding 623 response to bacterial and cyanobacterial diversity and 624 availability (Moens et al., 1999; Höckelmann et al., 625 2004; Schroeder et al., 2010). Therefore, it can be suggested that the higher nematode diversity 626 observed during summer could result from a decrease 627 of interspecific competition while the microbial food 628 resources are more diversified (e.g. cyanobacteria, 629 green microalgae and potentially bacteria), confirm-630 ing that resource availability can structure nematode 631 species composition and diversity (Michiels & 632 Traunspurger, 2005b; Ristau & Traunspurger, 2011). 633

Secondly, Michiels & Traunspurger (2003, 2004) 634 observed that the density of predators can increase the 635 number of co-existing nematode species by preventing 636 competitive exclusion due to dominant species. In the 637 present study, the density of the predatory nematode 638 Brevitobrilus stefanskii was positively linked to sum-639 mer conditions (Fig. 5). However, preventing com-640 petitive exclusion could also have resulted from 641 macrobenthic predators and grazers (e.g. insect larval 642 stages of Plecoptera, Trichoptera and Ephemeroptera), 643 which are particularly abundant during summer 644 (peaking in early July) in the Garonne River (Leflaive 645 et al., 2008, Majdi et al., unpubl. data). 646

Thirdly, temperature is known to strongly influence 647 benthic communities in running waters (Hawkins 648 et al., 1997; Stead et al., 2003). When temperature is 649 high, the biomass of the epilithic biofilm remains 650 severely controlled by self-generated detachment 651 processes and grazers (Boulêtreau et al., 2006; Hille-652 brand, 2009). Moreover, Lawrence et al. (2002) 653 experimentally showed that grazing of phototrophic 654 biofilm by macrobenthic invertebrates resulted in a 655 significant reduction of autotrophic biomass with an 656 increase of bacterial biomass within grazed regions, 657 corroborating the first hypothesis described above. 658 Thus, these disturbances can lead to a thin summer 659 biofilm layer with a high proportion of heterotrophic 660 organisms where intensive competition for space and 661 resources may create harsh life conditions for epiben-662 thic invertebrates. This suggestion is supported by the 663 decrease of the algal proportion in the biofilm 664 observed during this period. Therefore, it makes sense 665 that typical opportunistic and bacterial-feeding nem-666 atodes with a small body size and a low MI (e.g. 667 Monhysteridae) could benefit from these harsh condi-668 tions. Moreover, Monhysteridae species-especially 669 genus Eumonhystera-are known to reproduce 670 parthenogenetically (Traunspurger, 1991). This repro-671 ductive strategy probably accounted for the significant 672 reduction of the male proportion observed during 673 summer (Fig. 4b). Overall, summer nematode species 674



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675 lifestyle fits well with corresponding biofilm biotic676 conditions, suggesting that a close coupling occurs677 between nematode assemblage functional structure

678 and biofilm characteristics.

679 Conclusion

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680 Biomass of epilithic microalgae constituting potential 681 food sources for nematodes was plainly identified as an important predictor of nematode community 682 683 dynamics. Overall, our results strongly suggest that 684 variations in microalgal composition and proportion 685 in the biofilm might drive the observed changes in 686 nematode diversity and functional feeding group 687 composition. This supports the hypothesis that nematodes are involved in a strong trophic coupling with 688 their microbial habitat and should be taken into 689 690 consideration in further studies on biofilm dynamics 691 and functioning. Notably, studies of nematode feed-692 ing behaviour could disentangle trophic interactions 693 in epilithic biofilms and their potential feedback on 694 biofilm's structure and composition.

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