



Open Archive Toulouse Archive Ouverte (OATAO)

OATAO is an open access repository that collects the work of Toulouse researchers and makes it freely available over the web where possible.

This is an author-deposited version published in: <http://oatao.univ-toulouse.fr/>
Eprints ID: 5373

To link to this article: DOI: 10.1007/s10750-011-0781-6

URL: <http://dx.doi.org/10.1007/s10750-011-0781-6>

To cite this version: Nabil Majdi, Walter Traunspurger, Stéphanie Boyer, Benoît Mialet, Michèle Tackx, Robert Fernandez, Stefanie Gehner, Loïc Ten-Hage and Evelyne Buffan-Dubau (2011) Response of biofilm-dwelling nematodes to habitat changes in the Garonne River, France: influence of hydrodynamics and microalgal availability. *Hydrobiologia*, vol. 673 (n° 1). pp. 229-244. ISSN 0018-8158

Any correspondence concerning this service should be sent to the repository administrator: staff-oatao@listes.diff.inp-toulouse.fr

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

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

8 Received: 1 February 2011 / Revised: 14 May 2011 / Accepted: 22 May 2011
9 © Springer Science+Business Media B.V. 2011

10 **Abstract** Lotic epilithic biofilms are submitted to
11 seasonal disturbances (e.g. flood events, self-detach-
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

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 · 42
Feeding types · Algae · Environmental factors 43

A1 Handling editor: Stefano Amalfitano

A2 N. Majdi (✉) · S. Boyer · B. Mialet · M. Tackx ·
A3 R. Fernandez · L. Ten-Hage · E. Buffan-Dubau
A4 EcoLab, University Paul Sabatier, 118 route de Narbonne,
A5 31062 Toulouse, France
A6 e-mail: majdi@cict.fr

A7 N. Majdi · S. Boyer · B. Mialet · M. Tackx ·
A8 R. Fernandez · L. Ten-Hage · E. Buffan-Dubau
A9 EcoLab, CNRS, 31062 Toulouse, France

A10 W. Traunspurger · S. Gehner
A11 Animal Ecology, University of Bielefeld, Morgenbreede
A12 45, 33615 Bielefeld, Germany

Introduction 45

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
et al., 2008). This organic layer which is named either 51
epilithic biofilm, epilithon, 'Aufwuchs' or periphyton 52
can comprise more than 30% of microalgae in terms 53
of biomass (Peterson, 1996). Consequently, epilithic 54

55 biofilms can constitute the main site of primary
56 production in shallow water rivers harbouring hard
57 substrates such as the Garonne in its middle part
58 (Ameziane et al., 2003). These biofilms contribute
59 substantially to benthic food web functioning (Liess
60 & Hillebrand, 2004) and to biogeochemical processes
61 such as decomposition and nutrient retention (e.g.
62 Ford & Lock, 1987; Battin et al., 2003; Teissier et al.,
63 2007). However, epilithic biofilms are unstable
64 habitats, well-exposed to environmental perturba-
65 tions. Hence they are strongly influenced by seasonal
66 disturbances such as floods (Biggs & Close, 1989)
67 and self-detachment, a temperature-dependent bacte-
68 rial degradation of the mat (Biggs, 1996; Boulêtreau
69 et al., 2006). These disturbances are recognized to
70 shape the biomass, diversity and viability of the algal
71 and bacterial communities inhabiting the mat (e.g.
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
78 important food resource for them (e.g. Peters &
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
87 (mainly nematodes) can influence the release of
88 unpleasant odorous metabolites (e.g. geosmin) by
89 cyanobacterial biofilms, implying high economic
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 &
100 Traunspurger, 2005; Peters et al., 2005). So far, only
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.,

2006; Caramujo et al. 2008). But, long-term studies of 104
biofilm-dwelling nematodes are still lacking, which 105
hampers the assessment of how epilithic nematode 106
communities react and adapt to recurrent (seasonal) 107
abiotic disturbances and/or to fluctuations of food 108
resources 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

132 Methods

133 Study site and sample collection

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

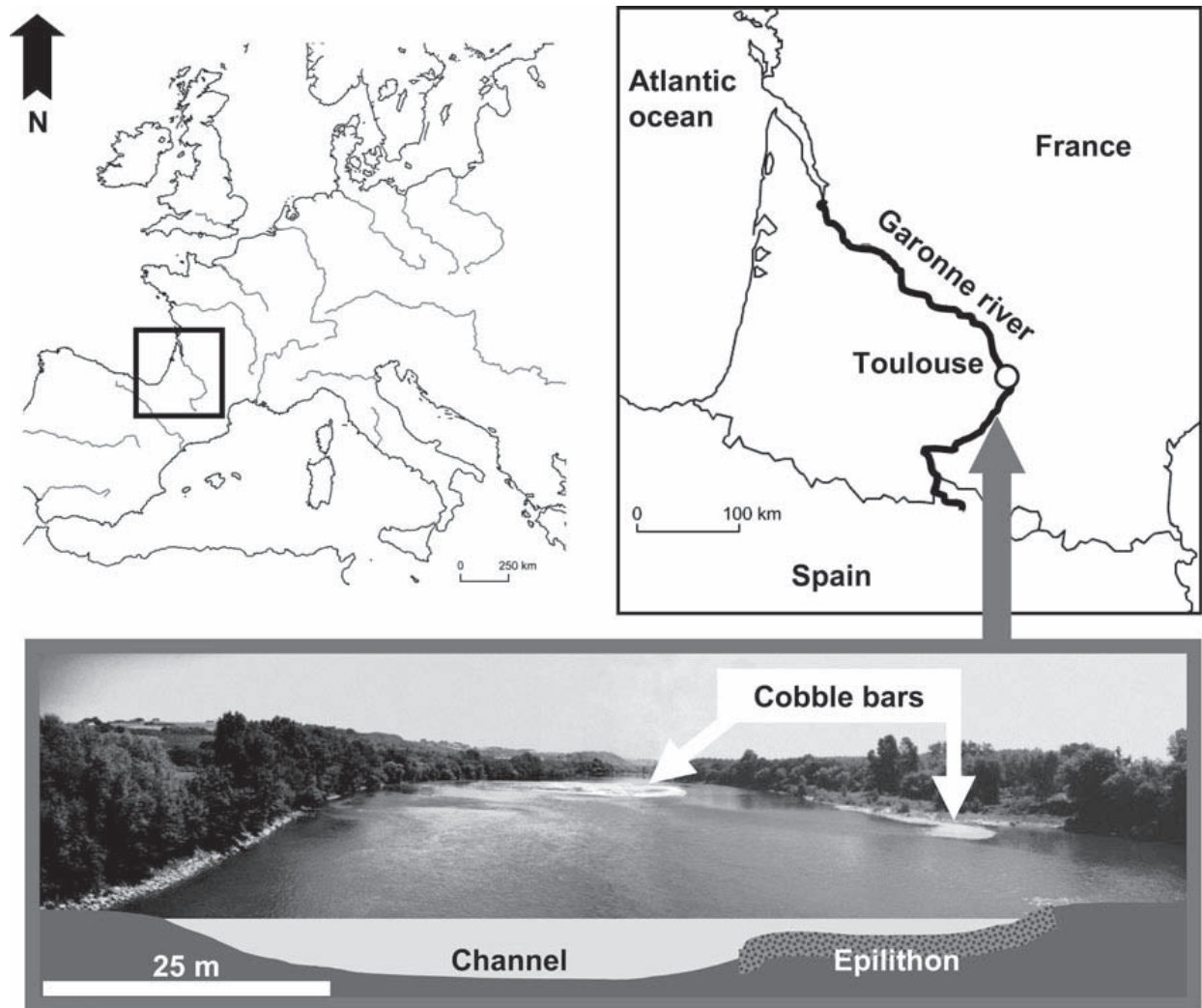


Fig. 1 Location of the sampling site and cross-section view of the Garonne River at the sampling site

148 the city of Toulouse (01°17'53"E, 43°23'45"N; eleva-
 149 tion 175 m a.s.l.), where the Garonne is of sixth-order
 150 (Fig. 1).

151 Samplings ($N = 51$) were regularly performed
 152 from September 2008 to March 2010 when hydro-
 153 logical conditions permitted it (sampling was only
 154 possible when discharge was lower than $175 \text{ m}^3 \text{ s}^{-1}$).
 155 On each sampling occasion, 12 immersed cobbles
 156 (mean diameter: 10 cm) were collected underwater
 157 using plastic bags to prevent any biofilm detach-
 158 ment during removal. To consider water level changes
 159 and 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
 disturbance. The biofilm was gathered by scraping the
 upper surface of each cobble with a scalpel and a
 toothbrush. Biofilm samples were finally suspended in
 MilliQ water to obtain 12 biofilm suspensions (25 ml
 each), in which algal aggregates were carefully crum-
 bled with scissors. These 12 biofilm suspensions were
 used for the three following treatments: (1) nematode
 species identification and density and biomass mea-
 surements, (2) HPLC analyses of microalgal pigments
 and (3) epilithic ash-free dry mass (AFDM) measure-
 ments. Four replicate suspensions were used for each
 treatment. Scraped cobbles were photographed, and
 the surface of biofilm which had been removed
 was clearly visible and measured using ImageJ soft-
 ware version 1.38 (Abramoff et al., 2004). Removed

180	biofilm surfaces were then reported to corresponding	Abiotic environmental factors	225
181	biofilm suspension volumes, so as densities, bio-		
182	mass and pigment concentrations were quantitatively	Mean Daily Discharge (MDD) was supplied by a	226
183	expressed per area unit.	gauging station of the French water management	227
		authority (DIREN Midi-Pyrénées, Marquefave sta-	228
184	Nematode processing	tion) located at 10 km upstream the study site—with	229
		no tributary and no dam between the gauging station	230
185	Nematodes were extracted from four replicate biofilm	and the study site. The Mean Weekly Discharge	231
186	suspensions using a modified gravity gradient centri-	(MWD) before each sampling occasion was consid-	232
187	fugation technique involving Ludox HS-40 after	ered in statistical analysis. To better reflect the effect	233
188	Pfannkuche & Thiel (1988). Nematodes so extracted	of flood disturbance, days after flood (DAF), which	234
189	were cleaned from Ludox by sieving through a 40 µm	were effective days after the last flood (MDD >	235
190	sieve, then preserved in formaldehyde (5% final	300 m ³ s ⁻¹), were calculated for each sampling	236
191	concentration) and stained with 1% Rose Bengal.	occasion and considered in statistical analysis. Water	237
192	Nematodes were counted in a Dolfuss cell (Elvetec	temperature, conductivity, pH and dissolved oxygen	238
193	services, Clermont-Ferrand, France) under a Leica	concentration were measured every 30 min during the	239
194	MZ 9.5 stereomicroscope (9×–90×) and their den-	whole study period with an automated multi-param-	240
195	sity was expressed per cm ² . According to nematode	eter probe (YSI 6000, YSI inc., Yellow springs, OH,	241
196	density, between 12 and 25 individuals were ran-	USA) which was permanently settled at 5 cm above	242
197	domly picked up from each replicate while counting,	the streambed at the study site.	243
198	transferred to glycerol solution (Seinhorst, 1959),		
199	mounted on slides and identified to the best species	Biofilm microalgal composition and biomass	244
200	level using a Leitz Dialux microscope at 1250×		
201	magnification.	<i>Microalgal pigments extraction and HPLC-analysis</i>	245
202	Nematodes were classified according to their age		
203	(juveniles, fourth stage juveniles and adults), their	On each sampling occasion, four replicate biofilm	246
204	sexual category (females, gravid females and males),	suspensions were centrifuged (3,220 g, 20 min).	247
205	and their feeding type (epistrate-feeders, deposit-	Pellets were freeze-dried and thoroughly homoge-	248
206	feeders, suction-feeders and chewers) after Traun-	nized. Then, 250 mg aliquots were removed from	249
207	spurger (1997). The Maturity Index (MI) was	each pellet. Algal pigments from each pellet aliquot	250
208	calculated on each sampling occasion as the weighted	were then extracted three times (15 min at –20°C)	251
209	mean frequency of individual colonizer–persister val-	with a total of 25 ml (10, 10 and 5 ml) 98% cold-	252
210	ues (cp) after Bongers (1990). MI ranged from 1 to 5.	buffered methanol (with 2% of 1 M ammonium	253
211	Nematode species with a cp = 1 were considered	acetate) following Buffan-Dubau & Carman (2000b).	254
212	r-strategists (colonizers) with short-generation times,	Algal pigment release was favoured at each extrac-	255
213	high fecundity and extreme population changes	tion step by an ultrasonication probe (Sonifier 250A,	256
214	whereas those with a cp = 5 were defined as K-strat-	Branson Ultrasonics corp., Danbury, CT, USA).	257
215	egists (persisters) with lower breeding efficiency. The	One millilitre of the pigment solution so obtained	258
216	MI is expected to decrease during disturbed periods,	was then filtered on 0.2 µm PTFE syringe filter and	259
217	when opportunistic nematodes are favoured (Bongers	analyzed using a high-performance liquid chromato-	260
218	& Bongers, 1998). Over a 1-year period from	graph (HPLC) consisting of a 100 µl loop auto-	261
219	September 2008 to September 2009 (N = 37), at least	sampler and a quaternary solvent delivery system	262
220	100 individual nematode body dimensions (length and	coupled to a diode array spectrophotometer (LC1200	263
221	maximum width) were measured on each sampling	series, Agilent Technologies inc., Santa Clara, CA,	264
222	occasion from microscopic pictures taken while	USA). The mobile phase was prepared and pro-	265
223	counting. Mean individual wet weight (WW) was	grammed according to the analytical gradient protocol	266
224	then determined after Andrassy (1956).	described in Barlow et al. (1997). Pigment separation	267
		was performed through a C8, 5 µm column (MOS-2	268

Table 1 CHEMTAX pigment ratio matrix

Algal group	Species	Biomarker pigment ratios to Chl <i>a</i>								
		Fuco	Lut	Viola	Diad	Zea	β -car	Chl <i>a</i>	Chl <i>b</i>	Chl <i>c</i>
Green algae	<i>P. boryanum</i>		0.143	0.049		0.014	0.043	1	0.088	
Diatoms	<i>N. palea</i>	0.477			0.102		0.002	1		0.121
Cyanobacteria	<i>S. leopoliensis</i>					0.411	0.011	1		

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)

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
287 and freshwater environments (e.g. Schlüter et al.,
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 $\mu\text{mol m}^{-2} \text{s}^{-1}$). 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
in further statistical analysis. 320

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 331

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

335 analysed between samples assigned to their corre- 367
 336 sponding sampling season (i.e. summer: 21 June–21 368
 337 September, $N = 15$; autumn: 21 September–21 369
 338 December, $N = 18$; winter: 21 December–21 March, 370
 339 $N = 15$ and spring: 21 March–21 June, $N = 3$). The 371
 340 homogeneity of variance was assessed with Levene's 372
 341 test, and differences were examined either by one- 373
 342 way ANOVA followed by a post-hoc Tukey HSD test 374
 343 or by Kruskal–Wallis ANOVA. The same statistical 375
 344 procedures were applied to investigate seasonal 376
 345 changes of biofilm and microalgal biomass. The 377
 346 correlations between total nematode density and 378
 347 biotic and abiotic factors were investigated by 379
 348 Spearman's rank correlation test. These tests were 380
 349 performed with STATISTICA software (version 8.0, 381
 350 Statsoft inc., Tulsa, OK, USA).

351 The influence of biotic and abiotic environmental 377
 352 factors on the nematode species distribution was 378
 353 analyzed through canonical ordination analysis with 379
 354 CANOCO software (version 4.5, Biometris, Wagen- 380
 355 ingen, The Netherlands). Rare species (with relative 381
 356 occurrence $<0.1\%$) were not considered in this anal- 382
 357 ysis. Species densities were square-root transformed 383
 358 prior to the analysis. The distribution of nematodes 384
 359 was first analyzed by a detrended correspondence 385
 360 analysis (DCA). As the total inertia observed was less 386
 361 than 2.6, a predominance of linear species response 387
 362 curves could be expected (Ter Braak, 1987, 1994). 388
 363 Therefore, a redundancy analysis (RDA) in which the 389
 364 ordination axes were constrained to be linear combi- 390
 365 nations of provided environmental factors was used to 391
 366 investigate the relationships between these factors and 392
 393

the distribution of main nematode species. Environ- 367
 368 mental factors were also listed (conditional effects) 369
 370 according to the variance they explained singly (i.e. 371
 372 without eventual co-variability with other factors). 373
 374 The statistical significance was tested with Monte 375
 376 Carlo permutation test (499 unrestricted permutations) 377
 378 with applying Bonferroni's correction (significance 379
 380 level set at $P < 0.005$). 381
 382

383 Results 384

385 Dynamics of the epilithic biofilm 386

387 The range and annual mean values of each measured 388
 389 abiotic and biotic factor are listed in Table 2. AFDM 390
 391 and Chl *a* content of the epilithic biofilm were 392
 393 significantly positively correlated (Spearman rank: 394
 395 $R = 0.75$; $P < 0.001$) and showed considerable vari- 396
 397 ations throughout the sampling period, being partic- 398
 399 ularly dampened after floods (Fig. 2a). The AI was 399
 400 significantly higher during summer than during the 400
 401 other seasons (ANOVA: $F = 60.2$; $P < 0.001$), 401
 402 implying globally a lower availability of microalgae 402
 403 within summer biofilm communities. Diatoms dom- 403
 404 inated the epilithic microalgal assemblage over the 404
 405 whole sampling period (Fig. 2b, Table 2). The dia- 405
 406 tom biomass was significantly higher during winter 406
 407 than during the other seasons (ANOVA: $F = 16.1$; 407
 408 $P < 0.001$). Conversely, cyanobacterial biomass was 408
 409 significantly higher during summer (ANOVA: $F =$ 409
 410 4.6 ; $P < 0.01$), and green algal biomass was 410

Table 2 Measured abiotic and biofilm biotic factors

	Annual mean \pm SE	Min	Max
Temperature ($^{\circ}\text{C}$)	14.6 \pm 0.05	1.7	27.3
O ₂ (mg l ⁻¹)	11.5 \pm 0.02	7.4	22.1
pH (-)	7.6 \pm 0.004	6.7	9.1
Conductivity ($\mu\text{S cm}^{-1}$)	270.9 \pm 0.001	154	493
Mean daily discharge (m ³ s ⁻¹)	124.7 \pm 6.0	18	814
Days after flood (day)	89.4 \pm 11.1	7	233
AFDM (g m ⁻²)	27.4 \pm 2.7	4.4	79.7
Chlorophyll <i>a</i> (mg m ⁻²)	321.5 \pm 50	10.7	1012.8
Green algae (%)	17.1 \pm 2.3	0	36.3
Cyanobacteria (%)	2.2 \pm 0.6	0	14.6
Diatoms (%)	80.7 \pm 2.7	50.6	100

Annual means refer to 2009. For temperature, O₂, pH and conductivity ($N = 17507$). For days after flood and the biotic factors ($N = 31$). Minimum and maximum values refer to the whole sampling period (i.e. September 2008–March 2010)

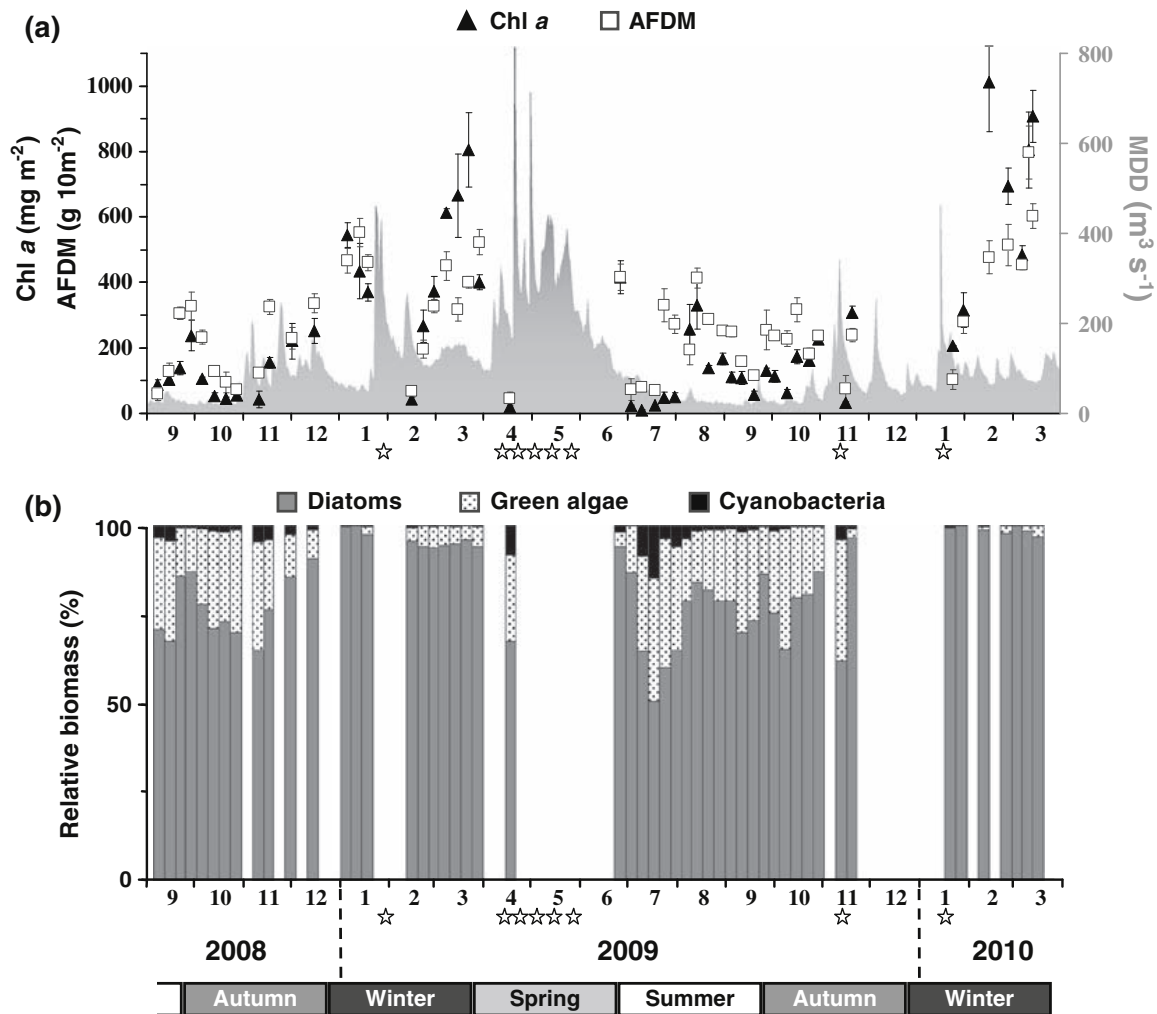


Fig. 2 Temporal dynamics of **a** epilithic chlorophyll *a* (Chl *a*) concentration (\pm SE, $N = 4$), ash-free dry mass (AFDM) of the biofilm (\pm SE, $N = 4$) and mean daily discharge (MDD), and **b** the relative proportion (%) of epilithic microalgal groups to

total Chl *a* biomass ($N = 4$). Months, years, seasons and floods during which MDD $> 300 \text{ m}^3 \text{ s}^{-1}$ (represented by stars) are indicated on the X axis

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

$R = 0.36$; $P < 0.01$), AFDM (Spearman rank: $R =$ 408
409 0.41 ; $P < 0.01$) and Chl *a* (Spearman rank: $R = 0.47$;
410 $P < 0.001$). From September 2008 to September 2009,
411 the nematode individual wet weight averaged $0.3 \mu\text{g}$.
412 The individual biomass was significantly lower during
413 summer (ANOVA: $F = 14.1$; $P < 0.001$) than during
414 the other seasons (Fig. 3a).

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

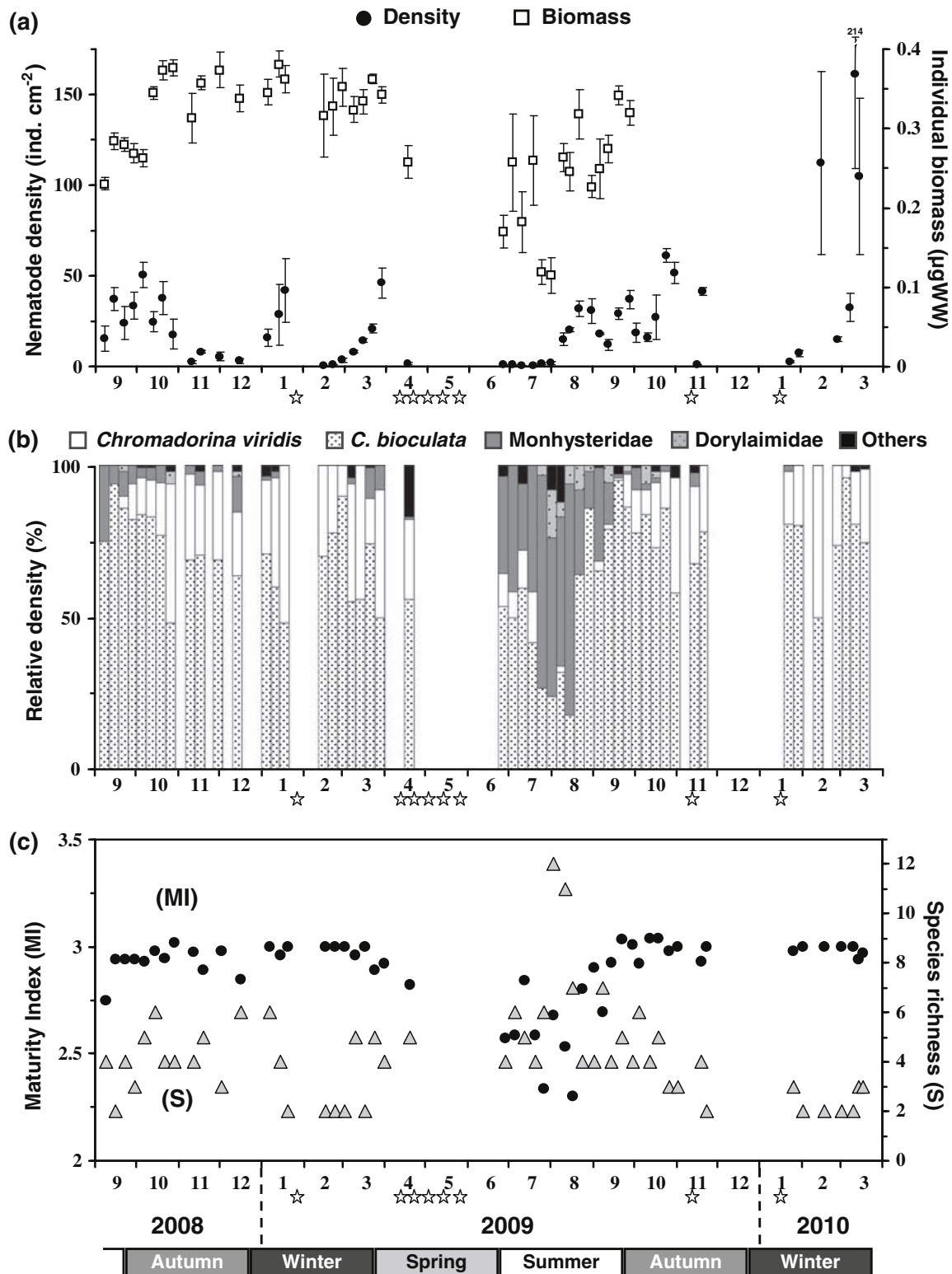


Fig. 3 Temporal dynamics of **a** nematode density (\pm SE, $N = 4$) and individual wet weight (WW) biomass (\pm SD, $N \geq 100$), **b** relative density of main nematode taxa, and **c** Maturity index (MI) and species richness (S) in the epilithic

biofilm. Months, years, seasons and floods during which $MDD > 300 \text{ m}^3 \text{ s}^{-1}$ (represented by stars) are indicated on the X axis

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

Nematode taxa	%	cp	FT
CHROMADORIDA Filipjev, 1929			
Chromadoridae Filipjev, 1917			
<i>Chromadorina bioculata</i> (Schultze in Carus, 1857)	68.87	3	E
<i>Chromadorina viridis</i> (Linstow, 1876)	17.15	3	E
Plectidae Örley, 1880			
<i>Plectus opisthocirculus</i> Andrassy, 1952	0.59	2	D
<i>Plectus aquatilis</i> Andrassy, 1985	0.14	2	D
<i>Plectus rhizophilus</i> de Man, 1880	<0.1	2	D
<i>Plectus cirratus</i> Bastian, 1865	<0.1	2	D
Prismatolaimidae Micoletzky, 1922			
<i>Prismatolaimus</i> cf. <i>intermedius</i> (Bütschli, 1873)	<0.1	3	E
Rhabdolaimidae Chitwood, 1951			
<i>Rhabdolaimus aquaticus</i> de Man, 1880	<0.1	3	D
MONHYSTERIDA Filipjev, 1929			
Monhysteridae de Man, 1876			
<i>Eumonhystera dispar</i> (Bastian, 1865)	6.92	2	D
<i>Eumonhystera vulgaris</i> (de Man, 1880)	1.84	2	D
<i>Eumonhystera simplex</i> (de Man, 1880)	0.35	2	D
<i>Eumonhystera barbata</i> Andrassy, 1981	0.31	2	D
<i>Eumonhystera</i> cf. <i>filiformis</i> (Bastian, 1865)	<0.1	2	D
<i>Eumonhystera longicaudatula</i> (Gerlach & Riemann, 1973)	<0.1	2	D
<i>Eumonhystera</i> sp.	<0.1	2	D
<i>Monhystrella paramacrura</i> (Meyl 1954)	1.04	2	D
DORYLAIMIDA Pearse, 1942			
Dorylaimidae de Man, 1876			
<i>Mesodorylaimus</i> cf. <i>subtiliformis</i> (Andrassy, 1959)	1.04	4	S
<i>Mesodorylaimus</i> sp.	<0.1	4	S
<i>Eudorylaimus</i> sp.	<0.1	4	S
<i>Dorylaimus stagnalis</i> Dujardin, 1845	<0.1	4	S
Mermithidae Braun, 1883			
Mermithidae	<0.1	1	P
ENOPLIDA Filipjev, 1929			
Tobrilidae Filipjev, 1918			
<i>Brevitobrilus stefanskii</i> (Micoletzky, 1925)	0.56	3	C
<i>Tobrilus gracilis</i> (Bastian, 1865)	<0.1	3	C
Tripylidae de Man, 1876			
<i>Tripyla</i> cf. <i>filicaudata</i> de Man, 1880	<0.1	3	C
<i>Tripyla glomerans</i> Bastian, 1865	<0.1	3	C
Alaimidae Micoletzky, 1922			
<i>Paramphidelus</i> sp.	<0.1	2	D
TYLENCHIDA Thorne, 1949			
Aphelenchoididae Skarbilovich, 1947			
<i>Aphelenchoides</i> sp.	0.24	2	S
Tylenchidae Örley, 1880			

Table 3 continued

Nematode taxa	%	cp	FT
<i>Coslenchus</i> sp.	<0.1	3	S

The proportion (%) of each species to the total number of identified nematodes ($N = 2875$) is provided. Each species is assigned to its corresponding colonizer–persistor value (cp) after Bongers & Bongers (1998) and to its corresponding feeding type (FT) after Traunspurger (1997): epistrate-feeders (E), deposit-feeders (D), suction-feeders (S) chewers (C) and insect parasites (P)

423 *paramacrura*—represented only 10% of all identified
 424 nematodes over the whole period, they clearly
 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
 430 was significantly higher during summer (ANOVA:
 431 $F = 6.5$; $P < 0.001$) than during the other seasons.
 432 Conversely, the Maturity Index (MI) was signifi-
 433 cantly lower (MI = 2.67) during summer (Kruskal–
 434 Wallis ANOVA: $H = 31.5$; $P < 0.001$) than during
 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
 442 chewers were less common representing, respec-
 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-feed-
 447 ers were significantly more represented (Kruskal–
 448 Wallis ANOVA: $H = 38.7$; $P < 0.001$) than during
 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

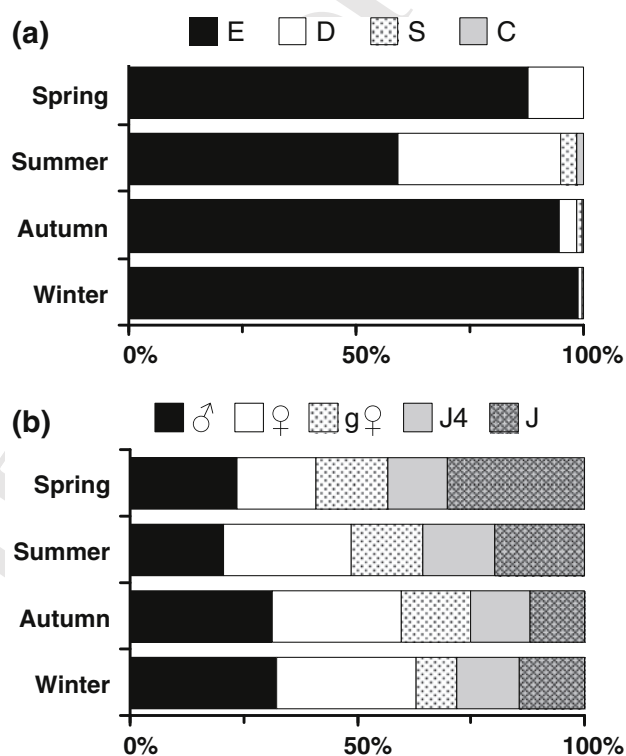


Fig. 4 Seasonal variations of the nematode community structure in the biofilm: **a** seasonal proportion of epistrate-feeders (E), deposit-feeders (D), suction-feeders (S) and chewers (C), and **b** seasonal proportion of males (♂), females (♀), gravid females (g♀), fourth stage juveniles (J4) and juveniles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

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

Influence of environmental factors on nematode 464
 species distribution 465

The results of the redundancy analysis (RDA) testing 466
 the influence of biotic and abiotic factors on nematode 467
 species and feeding-types distribution are presented in 468
 Fig. 5 and Table 4. The temporal distribution of 469
 nematode species was significantly influenced by 470

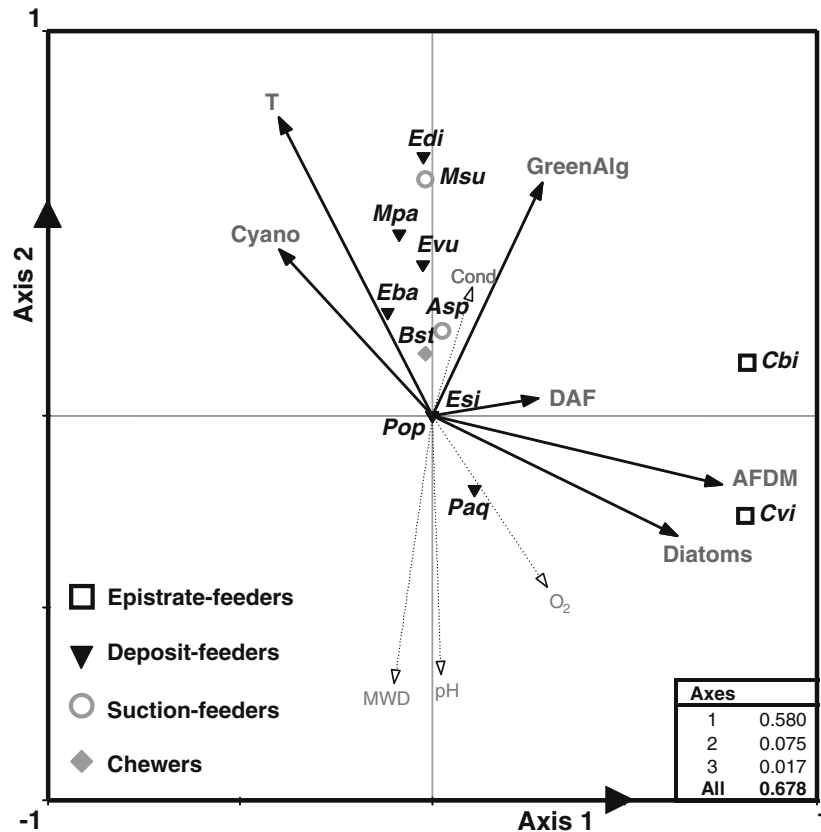


Fig. 5 Biplot from the redundancy analysis (RDA) explaining the distribution of nematode species densities according to environmental factors. Ordination axes were rescaled to range from -1 to 1 . *Slim dotted arrows* are non-significant factors. *Bold arrows* are significant factors (Monte Carlo permutation 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

dry mass (AFDM), water temperature (T), pH, dissolved O_2 (O_2), 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*)

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
 477 *C. bioculata* and *C. viridis*. These two species are
 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.

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

Discussion 498

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

Table 4 Conditional effects from the redundancy analysis (RDA)

Factors	λ	<i>P</i>
Diatoms	0.149	0.002**
<i>T</i>	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
pH	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-variability, 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)

520 biofilm-dwelling nematode community was not
521 diversified, two groups of species showing different
522 dynamics were clearly distinguished and seemed to
523 adapt to biofilm composition and seasonality: the first
524 group, consisting of the strongly dominating *Chrom-*
525 *adorina bioculata* and *C. viridis*, was mainly related
526 to biofilm composition (i.e. age, thickness and diatom
527 content) whereas the second group of species mainly
528 grew under summer conditions.

529 The nematode density averaged 25.4 ind cm⁻² and
530 ranged from 0.4 to 161.4 ind cm⁻² in the epilithic
531 biofilm over the whole study period. This result lies
532 within the range of values reported for lake epilithic
533 biofilms, i.e. 2.8–161.5 ind cm⁻² (Peters & Traun-
534 spurger, 2005) and for river epilithic biofilms, i.e.
535 10–100 ind cm⁻² (Gaudes et al., 2006). In our study,
536 the nematode community constituted a permanent
537 component of river epilithic biofilms. Mathieu et al.
538 (2007) suggested that nematode activity could affect
539 the oxygen turnover of diatom biofilms at density
540 values ≥ 50 ind cm⁻². This threshold value of density
541 was reached on several occasions during the study
542 period suggesting that this influence was substantial in
543 the epilithic biofilms of the Garonne River.

544 Nematode density positively correlated with
545 AFDM and Chl *a*. This strengthens the hypothesis

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

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

571 While diatoms dominated biofilm algal assem-
572 blages in terms of biomass, two epistrate-feeder
573 species *Chromadorina bioculata* and *Chromadorina*
574 *viridis* dominated strongly the nematode assemblage.
575 This observation supports the trend previously hypoth-
576 esized that, in freshwater benthic environments,

577 nematode communities are generally dominated by
 578 few species (e.g. Zullini & Ricci, 1980; Michiels &
 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.*
 585 *bioculata* and *C. viridis* were clearly segregated from
 586 the other nematode species (Fig. 5) and primarily
 587 positively related to diatom biomass. Due to their high
 588 content of polyunsaturated fatty acids (Phillips, 1984),
 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
 604 previously reported for sediment-dwelling nematode
 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
 616 diversified group of nematode species (mainly
 617 deposit-feeding species) grew under summer condi-
 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 ètreau 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

626 suggested that the higher nematode diversity
 627 observed during summer could result from a decrease
 628 of interspecific competition while the microbial food
 629 resources are more diversified (e.g. cyanobacteria,
 630 green microalgae and potentially bacteria), confirm-
 631 ing that resource availability can structure nematode
 632 species composition and diversity (Michiels &
 633 Traunspurger, 2005b; Ristau & Traunspurger, 2011).

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

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

675 lifestyle fits well with corresponding biofilm biotic
676 conditions, suggesting that a close coupling occurs
677 between nematode assemblage functional structure
678 and biofilm characteristics.

679 Conclusion

680 Biomass of epilithic microalgae constituting potential
681 food sources for nematodes was plainly identified as
682 an important predictor of nematode community
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 nem-
688 atodes are involved in a strong trophic coupling with
689 their microbial habitat and should be taken into
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.

695 **Acknowledgments** We thank J. Ferriol for helping with algal
696 cultures. We are grateful to K. Martens and to two anonymous
697 reviewers for their helpful comments on the manuscript. This
698 study was funded by a national CNRS EC2CO-CYTRIX
699 program. Nabil Majdi received a doctoral grant no. 31381-2008
700 by the French department of higher education and research
701 (MESR).

702 References

703 Abramoff, M. D., P. J. Magelhaes & S. J. Ram, 2004. Image
704 processing with ImageJ. *Biophotonics International* 11:
705 36–42.
706 Ameziane, T., F. Garabetian, D. Dalger, S. Sauvage, A. Dauta
707 & J. Capblancq, 2002. Epilithic biomass in a large gravel
708 bed river (the Garonne, France): a manifestation of
709 eutrophication? *River Research and Applications* 18:
710 343–354.
711 Ameziane, T., A. Dauta & R. Le Cohu, 2003. Origin and
712 transport of phytoplankton in a large river: the Garonne,
713 France. *Archiv für Hydrobiologie* 156: 385–404.
714 Andrassy, I., 1956. Die Rauminhalts- und Gewichtsbestim-
715 mung der Fadenwürmer (Nematoden). *Acta Zoologica*
716 *Hungarica* 2: 1–15.
717 Barlow, R. G., D. G. Cummings & S. W. Gibb, 1997. Improved
718 resolution of mono- and divinyl chlorophylls *a* and *b* and
719 zeaxanthin and lutein in phytoplankton extracts using
720 reverse phase C-8 HPLC. *Marine Ecology Progress Series*
721 161: 303–307.

Battin, T. J., L. A. Kaplan, J. D. Newbold & M. E. Hansen, 722
2003. Contributions of microbial biofilms to ecosystem 723
processes in stream mesocosms. *Nature* 426: 439–441. 724
Beier, S. & W. Traunspurger, 2003. Seasonal distribution of 725
free-living nematodes in the Körsch, a coarse-grained 726
submountain carbonate stream in southwest Germany. 727
Nematology 5: 481–504. 728
Biggs, B. J. F., 1996. Patterns in benthic algae of streams. In 729
Stevenson, R. J., M. L. Bothwell & R. L. Lowe (eds), 730
Algal Ecology: Freshwater Benthic Ecosystems. Aca- 731
demic Press, San Diego, CA, USA: 31–56. 732
Biggs, B. J. F. & M. E. Close, 1989. Periphyton biomass 733
dynamics in gravel bed rivers: the relative effects of flows 734
and nutrients. *Freshwater Biology* 22: 209–231. 735
Bongers, T., 1990. The maturity index: an ecological measure 736
of environmental disturbance based on nematode species 737
composition. *Oecologia* 83: 14–19. 738
Bongers, T. & M. Bongers, 1998. Functional diversity of 739
nematodes. *Applied Soil Ecology* 10: 239–251. 740
Boulêtreau, S., F. Garabetian, S. Sauvage & J. M. Sanchez- 741
Perez, 2006. Assessing the importance of a self-generated 742
detachment process in river biofilm models. *Freshwater* 743
Biology 51: 901–912. 744
Buffan-Dubau, E. & K. R. Carman, 2000a. Diel feeding behavior 745
of meiofauna and their relationships with microalgal 746
resources. *Limnology and Oceanography* 45: 381–395. 747
Buffan-Dubau, E. & K. R. Carman, 2000b. Extraction of 748
benthic microalgal pigments for HPLC analyses. *Marine* 749
Ecology Progress Series 204: 293–297. 750
Caramujo, M. J., C. R. B. Mendes, P. Cartaxana, V. Brotas & 751
M. J. Boavida, 2008. Influence of drought on algal bio- 752
films and meiofaunal assemblages of temperate reservoirs 753
and rivers. *Hydrobiologia* 598: 77–94. 754
Cardinale, B. J., 2011. Biodiversity improves water quality 755
through niche partitioning. *Nature* 472: 86–89. 756
Costerton, J. W., 2000. Biofilms in the new millennium: 757
musings from a peak in Xanadu. In Allison, D. G., P. 758
Gilbert, H. M. Lappin-Scott & M. Wilson (eds), *Com- 759*
munity Structure and Co-Operation in Biofilms. Cam- 760
bridge University Press, Cambridge, UK: 329–344. 761
Death, R. G. & M. J. Winterbourn, 1995. Diversity patterns in 762
stream benthic invertebrate communities: the influence of 763
habitat stability. *Ecology* 76: 1446–1460. 764
Deutsch, A., 1978. Gut ultrastructure and digestive physiology 765
of two marine nematodes, *Chromadorina germanica* 766
(Bütschli, 1874) and *Diplolaimella* sp. *Biological Bulletin* 767
155: 317–335. 768
Dole-Olivier, M. J., P. Marmonier & J. L. Befly, 1997. 769
Response of invertebrates to lotic disturbance: is the 770
hyporheic zone a patchy refugium? *Freshwater Biology* 771
37: 257–276. 772
Ford, T. E. & M. A. Lock, 1987. Epilithic metabolism of 773
dissolved organic carbon in boreal forest rivers. *FEMS* 774
Microbiology Letters 45: 89–97. 775
Gaudes, A., S. Sabater, E. Vilalta & I. Muñoz, 2006. The 776
nematode community in cyanobacterial biofilms in the 777
river Llobregat, Spain. *Nematology* 8: 909–919. 778
Goedkoop, W. & R. K. Johnson, 1996. Pelagic-benthic cou- 779
pling: profundal benthic community response to spring 780
diatom deposition in mesotrophic Lake Erken. *Limnology* 781
and *Oceanography* 41: 636–647. 782

- 783 Hawkins, C. P., J. N. Hogue, L. M. Decker & J. W. Feminella, 1997. Channel morphology, water temperature, and
784 assemblage structure of stream insects. *Journal of the*
785 *North American Benthological Society* 16: 728–749.
786 Hillebrand, H., 2009. Meta-analysis of grazer control of
787 periphyton biomass across aquatic ecosystems. *Journal of*
788 *Phycology* 45: 798–806.
789 Hillebrand, H., M. Kahlert, A. L. Haglund, U. G. Berninger, S.
790 Nagel & S. Wickham, 2002. Control of microbenthic
791 communities by grazing and nutrient supply. *Ecology* 83:
792 2205–2219.
793 Höckelmann, C., T. Moens & F. Jüttner, 2004. Odor com-
794 pounds from cyanobacterial biofilms acting as attractants
795 and repellents for free-living nematodes. *Limnology and*
796 *Oceanography* 49: 1809–1819.
797 Hodda, M., L. Peters & W. Traunspurger, 2009. Nematode
798 diversity in terrestrial, freshwater aquatic and marine
799 systems. In Wilson, M. J. & T. Kakouli-Duarte (eds),
800 *Nematodes as Environmental Indicators*. CABI Publish-
801 ing, Wallingford, UK: 45–94.
802 Kilham, S. S., D. A. Kreeger, S. G. Lynn, C. E. Goulden &
803 L. Herrera, 1998. COMBO: a defined freshwater culture
804 medium for algae and zooplankton. *Hydrobiologia* 377:
805 147–159.
806 Lawrence, J. R., B. Scharf, G. Packroff & T. R. Neu, 2002.
807 Microscale evaluation of the effects of grazing by inver-
808 tebrates with contrasting feeding modes on river biofilm
809 architecture and composition. *Microbial Ecology* 44:
810 199–207.
811 Leflaive, J., S. Boulétreau, E. Buffan-Dubau & L. Ten-Hage,
812 2008. Temporal patterns in epilithic biofilm-relation with
813 a putative allelopathic activity. *Fundamental and Applied*
814 *Limnology* 173: 121–134.
815 Liess, A. & H. Hillebrand, 2004. Invited review: direct and
816 indirect effects in herbivore-periphyton interactions.
817 *Archiv für Hydrobiologie* 159: 433–453.
818 Lionard, M., K. Muylaert, M. Tackx & W. Vymerman, 2008.
819 Evaluation of the performance of HPLC – CHEMTAX
820 analysis for determining phytoplankton biomass and
821 composition in a turbid estuary (Schelde, Belgium).
822 *Estuarine, Coastal and Shelf Science* 76: 809–817.
823 Lyautey, E., S. Boulétreau, E. Y. Madigou & F. Garabetian,
824 2010. Viability of differentiated epilithic bacterial com-
825 munities in the River Garonne (SW France). *Hydrobio-*
826 *logia* 637: 207–218.
827 Mackey, M. D., D. J. Mackey, H. W. Higgins & S. W. Wright,
828 1996. CHEMTAX – a program for estimating class
829 abundances from chemical markers: application to HPLC
830 measurements of phytoplankton. *Marine Ecology Pro-*
831 *gress Series* 144: 265–283.
832 Mathieu, M., J. Leflaive, L. Ten-Hage, R. de Wit & E. Buffan-
833 Dubau, 2007. Free-living nematodes affect oxygen turn-
834 over of artificial diatom biofilms. *Aquatic Microbial*
835 *Ecology* 49: 281–291.
836 Michiels, I. C. & W. Traunspurger, 2003. Maintenance of
837 biodiversity through predation in freshwater nematodes?
838 *Nematology Monographs and Perspectives* 2: 1–15.
839 Michiels, I. C. & W. Traunspurger, 2004. A three year study of
840 seasonal dynamics of a zoobenthos community in a
841 eutrophic lake. *Nematology* 6: 655–669.
842 Michiels, I. C. & W. Traunspurger, 2005a. Benthic community
843 patterns and the composition of feeding types and repro-
844 ductive modes in freshwater nematodes. *Nematology* 7:
845 21–36.
846 Michiels, I. C. & W. Traunspurger, 2005b. Impact of resource
847 availability on species composition and diversity in
848 freshwater nematodes. *Oecologia* 142: 98–103.
849 Michiels, I. C. & W. Traunspurger, 2005c. Seasonal variation
850 of biodiversity and assemblage structure in freshwater
851 nematodes. *Archiv für Hydrobiologie* 163: 183–194.
852 Moens, T., L. Verbeeck, A. de Maeyer, J. Swings & M. Vincx,
853 1999. Selective attraction of marine bacterivorous nema-
854 todes to their bacterial food. *Marine Ecology Progress*
855 *Series* 176: 165–178.
856 Palmer, M. A., P. Arensburger, A. P. Martin & D. W. Denman,
857 1996. Disturbance and patch-specific responses: the
858 interactive effects of woody debris and floods on lotic
859 invertebrates. *Oecologia* 105: 247–257.
860 Peters, L. & W. Traunspurger, 2005. Species distribution of
861 free-living nematodes and other meiofauna in littoral
862 periphyton communities of lakes. *Nematology* 7:
863 267–280.
864 Peters, L., W. Traunspurger, M. A. Wetzel & K. O. Rothhaupt,
865 2005. Community development of free-living aquatic
866 nematodes in littoral periphyton communities. *Nematol-*
867 *ogy* 7: 901–916.
868 Peterson, C. G., 1996. Response of benthic algal communities
869 to natural physical disturbance. In Stevenson, R. J., M.
870 L. Bothwell & R. L. Lowe (eds), *Algal Ecology: Fresh-*
871 *water Benthic Ecosystems*. Academic Press, San Diego,
872 CA, USA: 375–403.
873 Peterson, C. G. & R. J. Stevenson, 1992. Resistance and
874 resilience of lotic algal communities: importance of dis-
875 turbance timing and current. *Ecology* 73: 1445–1461.
876 Pfanckuche, O. & H. Thiel, 1988. Sample processing. In
877 Higgins, R. P. & H. Thiel (eds), *Introduction to the Study*
878 *of Meiofauna*. Smithsonian Institution Press, Washington,
879 DC, USA: 134–145.
880 Phillips, N. W., 1984. Role of different microbes and substrates
881 as potential suppliers of specific, essential nutrients to
882 marine detritivores. *Bulletin of Marine Science* 35:
883 283–298.
884 Ristau, K. & W. Traunspurger, 2011. Relation between nem-
885 atode communities and trophic state in southern Swedish
886 lakes. *Hydrobiologia* 663: 121–133.
887 Robertson, A. L., J. Lancaster, L. R. Belyea & A. G. Hildrew,
888 1997. Hydraulic habitat and the assemblage structure of
889 stream benthic microcrustacea. *Journal of the North*
890 *American Benthological Society* 16: 562–575.
891 Sabater, S., E. Vilalta, A. Gaudes, H. Guasch, I. Munoz & A.
892 Romani, 2003. Ecological implications of mass growth of
893 benthic cyanobacteria in rivers. *Aquatic Microbial Ecol-*
894 *ogy* 32: 175–184.
895 Schlüter, L., T. L. Lauridsen, G. Krogh & T. Jørgensen, 2006.
896 Identification and quantification of phytoplankton groups
897 in lakes using new pigment ratios – a comparison between
898 pigment analysis by HPLC and microscopy. *Freshwater*
899 *Biology* 51: 1474–1485.
900 Schroeder, F., D. Muschiol & W. Traunspurger, 2010. Fluc-
901 tuating food availability may permit coexistence in
902

- 903 bacterivorous nematodes. *Fundamental and Applied*
 904 *Limnology* 178: 59–66.
- 905 Seinhorst, J. W., 1959. A rapid method for the transfer of
 906 nematodes from fixative to anhydrous glycerin. *Nemato-*
 907 *logica* 4: 67–69.
- 908 Stead, T. K., J. M. Schmid-Araya & A. G. Hildrew, 2003. All
 909 creatures great and small: patterns in the stream benthos
 910 across a wide range of metazoan body size. *Freshwater*
 911 *biology* 48: 532–547.
- 912 Teissier, S., M. Torre, F. Delmas & F. Garabetian, 2007.
 913 Detailing biogeochemical N budgets in riverine epilithic
 914 biofilms. *Journal of the North American Benthological*
 915 *Society* 26: 178–190.
- 916 Ter Braak, C. J. F., 1987. Ordination. In Jongman, R. H. G., C.
 917 J. F. Ter Braak & O. F. R. Van Tongeren (eds), *Data*
 918 *Analysis in Community and Landscape Ecology*. Cam-
 919 *bridge University Press, Wageningen, The Netherlands:*
 920 91–173.
- 921 Ter Braak, C. J. F., 1994. Canonical community ordination.
 922 Part I: Basic theory and linear methods. *Ecoscience* 1:
 923 127–140.
- 924 Tietjen, J. H. & J. J. Lee, 1977. Feeding behaviour of marine
 925 nematodes. In Coull, B. C. (ed.), *Ecology of Marine*
 926 *Benthos*. University of South Carolina Press, Columbia,
 927 SC, USA: 21–36.
- 928 Traunspurger, W., 1991. *Fischbiologie des Königssees.*
 929 *Nahrungsangebot und Nahrungswahl. Band I: Das*
Meiobenthos des Königssees. Forschungsbericht 22. Na-
tionalpark Berchtesgaden, Germany: 152 pp.
- Traunspurger, W., 1992. A study of the free-living freshwater
 nematodes of hard substrates in the littoral of the oligo-
 trophic Koenigssee (National Park Berchtesgaden,
 F.R.G.). *Spixiana* 15: 233–238.
- Traunspurger, W., 1997. Bathymetric, seasonal and vertical
 distribution of the feeding-types of nematodes in an
 oligotrophic lake. *Vie et milieu* 47: 1–7.
- Traunspurger, W., 2002. Nematoda. In Rundle, S. D., A.
 L. Robertson & J. M. Schmid-Araya (eds), *Freshwater*
Meiofauna: Biology and Ecology. Backhuys Publishers,
 Leiden, The Netherlands: 63–104.
- Traunspurger, W., I. C. Michiels & E. Abebe, 2006. Compo-
 sition and distribution of free-living aquatic nematodes:
 global and local perspectives. In Abebe, E., W. Traun-
 spurger & I. Andrassy (eds), *Freshwater Nematodes:*
Ecology and Taxonomy. CABI Publishing, Wallingford,
 UK: 46–77.
- Wright, S. W., S. W. Jeffrey, R. F. C. Mantoura, C. A.
 Llewellyn, T. Bjornland, D. Repeta & N. Welschmeyer,
 1991. Improved HPLC method for the analysis of chlo-
 rophylls and carotenoids from marine phytoplankton.
Marine Ecology Progress Series 77: 183–196.
- Zullini, A. & C. Ricci, 1980. Bdelloid rotifers and nematodes
 in a small Italian stream. *Freshwater Biology* 10: 67–72.