### 1 Supplementary material

### 2 Hydrochemical analyses of water samples

Samples for Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup>, SO<sub>4</sub><sup>-2</sup> and HCO<sub>3</sub><sup>-2</sup> analysis were filtered through 0.2- $\mu$ m nylon filters, 3 4 stored at 4°C and analyzed using high performance liquid chromatography (HPLC) with a 5 WATERS 515 HPLC pump, IC-PAC anion columns, and a WATERS 432 detector. Samples for the 6 determination of cations were filtered through a 0.2-µm filter, acidified in the field with 1% 7 HNO3<sup>-</sup> and stored at 4°C. Cations were analyzed using inductively coupled plasma-optical 8 emission spectrometry (ICP-OES, Perkin-Elmer Optima 3200 RL). Samples for DOC analysis 9 were filtered through a 0.45-μm nylon filter and collected in muffled (450°C, 4.30 h) glass 10 bottles, acidified and stored at 4°C. In addition, water for TOC determination was sampled and stored at 4°C. TOC and DOC were analyzed with an infrared detector using the NPOC method 11 12 (Shimadzu TOC-Vcsh).

### 13 Molecular analyses for liquid and soil samples

Liquid samples were filtered through 0.22-µm GV Durapore® membrane filters (Merck 14 15 Millipore, USA) and stored at -80°C. Soil samples were centrifuged at 14.000 rpm and the 16 liquid fraction was discarded before pellets were cold-stored at -80°C. Total DNA extraction 17 was conducted using PowerWater® and PowerSoil® DNA Isolation Kits (MoBio Laboratories, 18 USA) for water (100 mL) and soil (200 mg) samples, respectively. For bacterial analyses, a 550bp DNA fragment in the 16S region of the small-subunit ribosomal RNA gene was amplified 19 20 using the primer set 341f/907r (Muyzer et al., 1993) a with a GC clamp added at the 5' end of 21 the forward primer. Final concentrations of the PCR reactions consisted of 1x PCR buffer, 2 22 mM of MgCl<sub>2</sub>, 200 µM of each deoxynucleoside triphosphate, 500 nM of each primer, 2.5 U of 23 Taq DNA polymerase (Invitrogen, ThermoFisher Scientific, USA) and 10 ng of template DNA. 24 The amplification protocol consisted of: 94ºC for 5 min; 20 cycles of 94ºC for 1 min, 65ºC for 1

25 min (-0.5°C/cycle), 72°C for 3 min; 15 cycles of 94°C for 1 min, 55°C for 1 min, 72°C for 3 min;
26 and a single final extension of 72°C for 7 min.

27 Denaturing gradient gel electrophoresis (DGGE) was performed using the Dcode Universal 28 Mutation Detection System (Bio-Rad, Spain). First, 900 ng of DNA from PCR products were 29 loaded onto 6% (w/v) polyacrylamide gels (acrylamide/bis solution 37.5:1) containing linear 30 chemical gradients of 30-70% denaturant. The 100% denaturing solution contained 7 M urea 31 and 40% (v/v) deionized formamide. Gels were run in 1X Tris acetate-EDTA (TAE) for 16 h at 75 32 V and 60°C, stained with 1 µg/mL ethidium bromide solution for 25 min, washed with 33 deionized water for 25 min and photographed with Universal Hood II (Bio-Rad, Spain). DGGE 34 images were analyzed using InfoQuest™ FP software. Dice's coefficient and the unweighted 35 pair group method with arithmetic averages (UPGMA) were employed for the clustering of 36 DGGE gel profiles. Non-metric Multidimensional Scaling (NMD) was performed using the 37 Vegan package (Oksanen et al., 2017). Prominent bands from the DGGE were excised, re-38 amplified and sequenced by Macrogen (South Korea). The obtained sequences were trimmed 39 with FinchTV software and checked for chimeras using the UCHIME algorithm (Edgar et al., 40 2011) integrated in Mothur version 1.38 (Schloss et al., 2009). Each 16S rRNA sequence was 41 assigned to its closest neighbor according to the Basic Local Alignment Search Tool (BLAST) 42 results (Altschul et al., 1997). Curated sequences were deposited in the National Center for 43 Biotechnology Information (NCBI) GenBank database under accession numbers MF471641-44 MF471667.

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	Table	251-3	Summ	ary of	nyaro	cnemi	cal pai	ramet	ers at		obrega		k site i	n Dotr	i scena	arios L	ry and	i wet.						
Sampling location (depth- <i>masl</i> )	P1-BG	G (-0.6)	P8	(1)	P8	(-3)	P2	(3)	P2	(-2)	P2	(-5)	P5 (	2.3)	P5 (	-2.2)	РЗ (	0.8)	P3 (-	4.2)	P10	(-1)	Infiltı ba	ration asin
Scenario	W	D	W	D	W	D	W	D	W	D	W	D	W	D	W	D	W	D	W	D	W	D	W	D
рН	7.05	7.36	7.45	7.13	7.39	7.27	7.30			7.34	7.28		7.39	6.98	7.42		7.15	7.15	7.15		7.28	7.35	8.47	
DOC (mg/l)	1.63	1.93	13.65	3.96	8.02	1.652	3.17			1.80	4.70		3.61	1.53	3.76		3.22	2.33	3.15		2.57	1.61	6.6	
TOC (mg/L)	2.48	1.97	7.95	4.05	13.05	1.80	1.99			2.00	3.64		2.96	1.50	2.70		2.13	1.88	2.94		3.58	1.70		
HCO3 <sup>-</sup> (mg/L)	354.3	382.6	241.3	379.3	253.8	408.1	261.2			314.2	272.6		256.1	408.1	255.2		399.0	294.3	385.8		327.1	396.4	144.5	
NO <sub>3</sub> <sup>-</sup> (mg/L)	5.02	17.79	2.75	2.82	2.88	15.49	3.13			12.9	3.64		3.41	17.7	4.08		6.06	18.2	8.73		5.88	16.28	6.14	
SO4 <sup>-</sup> <sub>2</sub> (mg/L)	173.0	181.5	120.0	141.23	152.1	199.54	138.4			140.0	138.4		133.8	200.3	147.8		168.6	177.0	168.7		180.0	174.8	123.4	
Cl <sup>-</sup> (mg/L)	224.8	212.4	267.5	186.1	262.8	193.5	267.0			211.8	212.0		274.6	191.7	269.7		241.0	193.2	241.7		265.5	189.1	245.9	
Ca (mg/L)	135.2	142.6	92.5	123.0	99.9	72.3	94.2			121.4	103.5		97.4	141.9	96.7		144.2	133.3	141.3		123.3	132.7	67.6	
Na (mg/L)	138.9	116.3	160.0	110.1	161.2	116.7	153.9			113.5	158.2		163.4	118.0	165.0		142.5	108.8	143.9		162.8	117.5	147.9	
K (mg/L)	19.6	23.7	30.0	23.0	29.4	20.7	30.2			22.9	31.0		38.6	24.4	31.9		24.2	23.9	24.4		26.3	27.5	28.0	
Mg (mg/L)	33.73	36.3	26.7	30.8	28.5	38.3	27.4			31.9	30.0		29.1	38.2	28.8		24.2	33.0	35.2		31.4	37.2	23.0	
S (mg/L)	52.2	63.5	40.5	51.2	47.2	64.3	41.7			58.1	42.6		42.9	63.4	42.7		47.7	58.7	47.9		51.4	59.9	42.2	
Fe (mg/L)	<0.2	<0.2.	<0.2	<0.2	<0.2	<0.2	<0.2			<0.2	<0.2		<0.2	<0.2	<0.2		<0.2	<0.2	<0.2		<0.2	<0.2	<0.2	
Mn (ppb)	28.9	0.87	2.46	145.30	2.77	<0.8	10.79			5.507	8.94		2.17	<0.8	1.46		3.67	0.81	1.74		1.58	<0.8	4.76	

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# Table S2. Sequence information corresponding to the DGGE bands obtained

## from bacterial populations' fingerprints in the Llobregat MAR system studied.

Phylot	type	Phylogenetic affiliation				
Band	Accesion	Taxonomic linage (Phylum, Class, Order, Family,	Closest match	Similarity		
code <sup>a</sup>	number	Genus) <sup>b</sup>	(accesion no.) <sup>c</sup>	(%) <sup>d</sup>		
<b>P</b> 1	ME471641	Firmicutes(100); Bacilli(100); Bacillales(100);	Chryseomicrobium sp.	08		
DI	WIF471041	Planococcaceae(100)	(KX889925)	90		
B2	ME471642	Proteobacteria(100); Betaproteobacteria(100);	Vogesella indigofera	100		
52	WII 47 1042	Neisseriales(94); Neisseriaceae(94)	(KF951043)	100		
B3	MF471643	Firmicutes(100); Bacilli(100); Bacillales(100);	Chryseomicrobium sp.	99		
55	WII 47 1045	Planococcaceae(100)	(KX889925)	55		
			Uncultured beta			
B4	ME471644	Proteobacteria(94); Betaproteobacteria(87); TRA3-20(54);	proteobacterium	94		
D4	WII 47 1044	TRA3-20_fa(54)	(KF182906)	54		
		Chloroflexi(100); Dehalococcoidia(100); vadinBA26(61); vadinBA26 fa(61)	Dehalogenimonas			
B5	MF471645		alkenigignens	100		
			(JQ994267)			
		Proteobacteria(100); Gammaproteobacteria(100);	Stenotrophomonas sp.			
B6	MF471646	Xanthomonadales(100); Xanthomonadaceae(100);	(LC136883)	97		
		Stenotrophomonas(98)	( ,			
		Nitrospirae(100): Nitrospira(100): Nitrospirales(100):	Uncultured bacterium			
B7	MF471647	Nitrospiraceae(100); Nitrospira(100)	(HM445209)	97		
			Dehalogenimonas			
B8	MF471648	Chloroflexi(100); Dehalococcoidia(100); vadinBA26(52); 1648	alkenigignens	100		
		vadinBA26_fa(52)	(JQ994267)			
		Firmicutes(100); Bacilli(100); Bacillales(100);	Chryseomicrobium sp.			
B9	MF471649	Planococcaceae(100)	(KX889925)	99		
B10	MF471650	Cyanobacteria(100); Cyanobacteria(100); SubsectionI(100);	Uncultured	99		

1		FamilyI(100)	cyanobacterium	
			(FJ916292)	
B11			Uncultured	
	MF471651	Cyanobacteria(100); Cyanobacteria(100); SubsectionI(100);	cyanobacterium	97
		Family(100)	(FJ916292)	
			Uncultured	
B12	MF471652	Proteobacteria(97); Betaproteobacteria(86);	Burkholderiaceae	94
		Nitrosoffonadales(38); Nitrosoffonadaceae(34)	(AM935619)	
		Proteobacteria(100); Betaproteobacteria(100);	Uncultured	
B13	MF471653	Nitrosomonadales(89); Nitrosomonadaceae(89);	Burkholderiaceae	97
		uncultured(88)	(AM935619)	
			Uncultured beta	
B14	MF471654	Proteobacteria(100); Betaproteobacteria(100); TRA3- 20(100); TRA3-20_fa(100); TRA3-20_ge(100)	proteobacterium	99
			(EU979071)	
		Proteobacteria(99); Betaproteobacteria(94); Hydrogenophilales(47); Hydrogenophilaceae(47)	Uncultured beta	
B15	MF471655		proteobacterium	92
			(JN868168)	
		Proteobacteria(82): Retanroteobacteria(56):	Uncultured beta	
B16	MF471656	Nitrosomonadales(20); Nitrosomonadaceae(20)	proteobacterium	88
			(AM935274)	
		Nitrospirae(100); Nitrospira(100); Nitrospirales(100); Nitrospiraceae(100); <i>Nitrospira</i> (100)	Uncultured	
B17	MF471657		Nitrospiraceae	97
			(EU298577)	
		Bacteroidetes(100); Cytophagia(100); Cytophagales(100);	Uncultured	
B18	MF471658		Bacteroidetes	99
		-,,,,,,,,,	(HF564274)	
B19	MF471659	Chlorobi(80); Chlorobia(80); Chlorobiales(80); OPB56(80);	Uncultured bacterium	99
_		OPB56_ge(80)	(KC666711)	
		Actinohacteria(99). Actinohacteria(99). Frankiales(99).	Uncultured	
B20	MF471660	Sporichthyaceae(99); Sporichthyaceae ge(99)	actinobacterium	99
			(LC018957)	
		Actinobacteria(93); Actinobacteria(93): Micrococcales(86):	Uncultured	
B21	MF471661	Microbacteriaceae(77)	actinobacterium	96
		.,	(LC018957)	
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		Acidahactoria(01): Solihactoroc(01): Solihactoraloc(01):	Uncultured			
B22	MF471662	Solibactoraciana (Subaroun 2)(01)	Acidobacteria	97		
		2010acteraceae_/2008r00h_2/21	(KM016273)			
	NAE 474 CC2	Proteobacteria(100); Betaproteobacteria(100);	Uncultured bacterium	erium 95		
BZ3	MF471663	Burkholderiales(99); Comamonadaceae(99)	(EU465081)			
			Uncultured	94		
B24	MF471664	Proteobacteria(100); Betaproteobacteria(100); Burkholderiales(94); Comamonadaceae(79)	Comamonadaceae			
			(LT679549)			
		Proteobacteria(96); Alphaproteobacteria(90); Rhizobiales(75); Rhizobiales_Incertae_Sedis(25)	Uncultured alpha			
B25	MF471665		proteobacterium	98		
			(HF584680)			
		Proteobacteria(100); Betaproteobacteria(100);				
B26	MF471666	Burkholderiales(100); Comamonadaceae(100);	Uncultured bacterium	99		
		Aquabacterium(100)	(KF065163)			
		Bacteria(100); Proteobacteria(100);				
B27		Betaproteobacteria(100); Methylophilales(100);	Methylotenera mobilis	00		
	WIF4/166/	Methylophilaceae(100); Methylophilus(100)	(AB698738)	99		

<sup>a</sup> band numbers correspond to those presented in figure 5

52 <sup>b</sup> taxonomic string with bootstrap values (in parentheses), generated in mothur using SILVA

53 database reference file release 119

54 closest relative according to INSA (International Nucleotide Sequence Database)

55 <sup>d</sup> percentage sequence similarity with closest INSA using BLAST tool

Table S3 – Granulometric composition of sediments in Llobregat MAR site									
Sampling	ocation	>12.7 mm	2-12.7 mm	0.3-2 mm	0.056-0.3 mm	<0.056 mm			
Pre-sedimentation basin	Decantation	39.48	6.68	20.66	28.10	5.07			
Le Ciline i tra a la sette	Entrance	54.05	25.43	12.93	6.91	0.68			
Infiltration basin	Midfield	40.35	30.36	20.90	7.62	0.76			
	End	15.36	40.26	31.54	11.47	1.37			

### 59 References

- 60 Altschul, S., Madden, T., Schaffer; AA, Zhang, J., Zhang, Z., Miller, W.
- and Lipman, D.: Gapped BLAST and PSI-BLAST: a new generation of
- 62 protein database search programs, Nucleic Acids Res, 25(17), 3389-
- 63 3402, doi:10.1093/nar/25.17.3389, 1997.
- 64 Edgar, R. C., Haas, B. J., Clemente, J. C., Quince, C. and Knight, R.:
- 65 UCHIME improves sensitivity and speed of chimera detection,
- 66 Bioinformatics, 27(16), 2194–2200 [online] Available from:
- 67 http://dx.doi.org/10.1093/bioinformatics/btr381, 2011.
- 68 Muyzer, G., De Waal, E. C. and Uitterlinden, A. G.: Profiling of
- 69 complex microbial populations by denaturing gradient gel
- 70 electrophoresis analysis of polymerase chain reaction-amplified
- 71 genes coding for 16S rRNA, Appl Environ Microbiol, 59(3), 695–700,
- 72 doi:0099-2240/93/030695-06\$02.00/0, 1993.
- 73 Oksanen, J., Blanchet, F., Kindt, R., Legendre, P., Minchin, P., O'Hara,
- 74 R., Simpson, G., Solymos, P., Stevens, M. and Wagner, H.: Vegan:
- 75 Community Ecology Package. R package version 2.4–3. The R Project
- 76 for Statistical Computing, Vienna, Austria., [online] Available from:
- 77 https://cran.r-project.org/web/packages/vegan/index.html
- 78 (Accessed 21 July 2017), 2017.
- 79 Schloss, P. D., Westcott, S. L., Ryabin, T., Hall, J. R., Hartmann, M.,
- 80 Hollister, E. B., Lesniewski, R. A., Oakley, B. B., Parks, D. H.,
- 81 Robinson, C. J., Sahl, J. W., Stres, B., Thallinger, G. G., Van Horn, D. J.
- 82 and Weber, C. F.: Introducing mothur: open-source, platform-

- 83 independent, community-supported software for describing and
- 84 comparing microbial communities., Appl Environ Microbiol, 75(23),
- 85 7537–41, doi:10.1128/AEM.01541-09, 2009.

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