Extremophiles

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Continuous enrichment cultures: insights into prokaryotic diversity and metabolic interactions in deep-sea vent chimneys

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Abstract:

The prokarvotic diversity of culturable thermophilic communities of deep-sea hydrothermal chimneys was analysed using a continuous enrichment culture performed in a gas-lift bioreactor, and compared to classical batch enrichment cultures in vials. Cultures were conducted at 60°C and pH 6.5 using a complex medium containing carbohydrates, peptides and sulphur, and inoculated with a sample of a hydrothermal black chimney collected at the Rainbow field, Mid-Atlantic Ridge, at 2,275 m depth. To assess the relevance of both culture methods, bacterial and archaeal diversity was studied using cloning and sequencing, DGGE, and whole-cell hybridisation of 16S rRNA genes. Sequences of heterotrophic microorganisms belonging to the genera Marinitoga, Thermosipho, Caminicella (Bacteria) and Thermococcus (Archaea) were obtained from both batch and continuous enrichment cultures while sequences of the autotrophic bacterial genera Deferribacter and Thermodesulfatator were only detected in the continuous bioreactor culture. It is presumed that over time constant metabolite exchanges will have occurred in the continuous enrichment culture enabling the development of a more diverse prokaryotic community. In particular, CO2 and H2 produced by the heterotrophic population would support the growth of autotrophic populations. Therefore, continuous enrichment culture is a useful technique to grow over time environmentally representative microbial communities and obtain insights into prokaryotic species interactions that play a crucial role in deep hydrothermal environments.

Keywords: Microbial diversity - Deep-sea hydrothermal vent - Continuous enrichment cultures -Bioreactor - Thermophiles - 16S rRNA gene - Metabolic interactions

46 Introduction

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The widespread application of 16S rRNA gene based molecular methods to identify microorganisms in natural 48 49 samples has revealed an extensive and, in many cases, unexpected microbial diversity. Within deep-sea 50 hydrothermal environments, the diversity of microbial communities associated with in situ colonizers 51 (McCliment et al. 2006), mats (Moussard et al. 2006), animals (DeChaine et al. 2006), sediments (Inagaki et al. 52 2006) and chimneys (Kormas et al. 2006) have been reported in recent molecular surveys. The rise of molecular 53 microbial ecology has resulted in the detection of many microorganisms that have as yet not been cultivated, for 54 example the widespread Marine Crenarchaeota Group I (MGI) and the Korarchaeota. To understand the 55 physiology and ecological significance of these uncultivated microorganisms, an effort has to be made to improve and develop cultural approaches. Indeed, culture conditions routinely used reveal only a small fraction 56 57 of the global microbial community. As an alternative to batch cultures in vials, a gas-lift bioreactor was 58 developed to grow anaerobic and hyperthermophilic microorganisms in continuous culture (Raven et al. 1992). 59 Recently, it has been used to study the metabolism of members of the Thermococcales, including Pyrococcus 60 abyssi (Godfroy et al. 2000), to develop minimal media and to optimize the growth conditions of Pyrococcus 61 furiosus and Thermococcus hydrothermalis (Raven and Sharp 1997; Postec et al. 2005a). In addition to the study 62 of pure cultures, the gas-lift bioreactor can also be used to cultivate representative microorganisms from 63 environmental samples in continuous enrichment culture under controlled conditions. The bioreactor allows the 64 long-term cultivation of microbes by enabling a continuous substrate supply, the elimination of volatile 65 metabolic end-products (potentially toxic for microbial growth) by gas sparging and pH and temperature regulation. These features help to grow less dominant microorganisms having poor representation, long latency 66 phase and/or slow growth. In a previous study, the gas-lift bioreactor was used to enrich microorganisms from a 67 68 black smoker collected at 2275 m depth on the Rainbow hydrothermal field of the Mid-Atlantic Ridge (Postec et 69 al. 2005b). A fifty-days continuous culture at 90°C on a rich medium containing sulphur under anaerobic 70 conditions demonstrated a large diversity inside the cultivated community, including (in addition to archaeal 71 species belonging to the order *Thermococales*) moderately thermophilic members of the orders *Clostridiales* and 72 Thermotogales, and members of the Epsilon proteobacteria that were not detected in vial cultures. In the present 73 study, the same black smoker chimney was used as inoculum to perform similar experiments on both batch and 74 continuous enrichment cultures under the same conditions, except the temperature lowered to 60°C. The 75 microbial diversity of the communities enriched in vials and bioreactor was analysed using the following

76	techniques based on the 16S rRNA genes: cloning, sequencing, denaturing gradient gel electrophoresis (DGGE)
77	and whole-cell hybridisation. Molecular results were the guidelines for subsequent isolation of microorganisms
78	from the enrichment cultures.
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80	Materials and methods
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82	Samples
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84	During the ATOS cruise (European project VENTOX) on the Rainbow field (36°13'N 33°54W, 2275 m) located
85	on the Mid-Atlantic Ridge (MAR), an active black smoker was collected by the Remotely Operated Vehicle
86	(ROV) Victor, and brought to the surface in an insulated box under aseptic conditions. On board, eight fragments
87	of the chimney were subsampled according to mineral zonations from the inner part to the outer part. The
88	subsamples were crushed in an anaerobic chamber (La Calhene, France) and stored in sterile serum vials filled
89	with sterile seawater containing 0.5 mg Γ^1 of Na ₂ S. All subsamples were pooled to represent the whole chimney
90	and this suspension was used to inoculate the enrichment cultures.
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92	Continuous enrichment culture in bioreactor: conditions and monitoring
93	
94	Medium
95	The growth medium was the modified SME medium (Sharp and Raven 1997) in which yeast and peptone were
96	replaced by: 1 g yeast extract (Difco), 0.5 g casaminoacids (Difco), 0.4 g glucose, 0.4 g dextrin (from corn), 0.2
97	g galactose, 0.2 g dextran, 0.1 g glycogen, 0.2 g pyruvate and 0.1 g acetate (all purchased from Sigma). The
98	medium was supplemented with 3 g l^{-1} colloidal sulphur.
99	
100	Culture conditions
101	The medium was sterilized by filtration (Sartroban, $0.22 \ \mu m$) in a 20 litre Nalgene bottle containing the colloidal
102	sulphur, previously sterilized by heating twice at 100°C for 30 min on two successive days. The culture was
103	performed at 60°C and pH 6.5 in a 2 litre glass gas-lift bioreactor as previously described (Raven et al. 1992;

105 temperature was controlled by a heated circulating bath filled with water and monitored with a standard PT100

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Godfroy et al. 2005). The bioreactor was inoculated at 2% (v/v) with the chimney sample suspension. The

106 probe covered with Teflon. The pH was monitored using a combination gel pH electrode (Mettler Toledo). Acid 107 (1N HCl) and base (1N NaOH) were added with peristaltic pumps (Masterflex). Temperature and pH were 108 controlled with a 4-20 mA controller and AFS Biocommand system from New Brunswick (Nijmegen, 109 Netherlands). Fresh medium addition and culture withdrawal were performed using peristaltic pumps (Masterflex). The culture was sparged with N₂ (0.1 v v^1 min⁻¹) to maintain anaerobic conditions, and to 110 111 eliminate volatile metabolic and products that might inhibit the growth of microorganisms (i.e. HS). The bioreactor was maintained as a batch culture for the first 34 h to prevent wash-out of the cells before they have 112 grown. After 34 h, fresh medium was provided by applying a dilution rate of 0.04 h⁻¹ (80 ml h⁻¹). To test the 113 114 effect of the temperature on the composition of the cultivated microbial community, the temperature was 115 increased from 60°C to 70°C at T31 (day 31) and to 80°C at T36 (day 36).

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117 Culture monitoring and sample preservation

118 The culture was maintained for 45 days. Culture samples from the bioreactor were collected every 24 hours, 119 from T0 (day 0) to T45 (day 45). Cell concentration was determined every day by direct cell counting, using a 120 Thoma chamber (0.02 mm depth) viewed with an Olympus BX60 phase contrast microscope (×400). For each 121 sampling, 10 ml of culture were preserved anaerobically at 4°C in a serum vial, and eight cryotubes containing each 1.8 ml of culture were frozen at -20°C with 5% DMSO (v/v). For DNA extraction, cells were recovered 122 123 from 15 ml of culture by centrifugation (20 min at 8,000 g). Cell pellets were washed with 23 g Γ^1 sterile NaCl, 124 then mixed to 5 ml of lysis buffer TE Na 1X (Tris-HCl pH 8, 100 mM; NaCl, 100 mM, EDTA pH 8, 50 mM), 125 and stored at -20°C until the DNA extraction procedure was undertaken. For whole-cell hybridisation, 12 ml of 126 culture sample were fixed for two hours with 3% (v/v) formaldehyde. Fixed cells recovered by centrifugation (10 127 min at 6,000 rpm) were washed with PBS buffer (phosphate-buffered: 8 g NaCl, 0.2 g KCl, 1.44 g Na₂HPO₄, 128 0.24 g KH₂PO₄, per litre of distilled water, pH 7.4) before storage in 50% (v/v) ethanol in PBS at -20°C. For 129 HPLC analyses, 1.5 ml of culture sample was centrifuged (10 min at 10,000 rpm) and the supernatant was stored 130 at 4°C until analysis.

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132 Batch enrichment cultures in vial: conditions and monitoring

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Cultures were performed anaerobically in 100 ml serum vials (Godfroy et al. 1996), using the medium described for the enrichment culture in the bioreactor to which 6.05 g I^1 PIPES was added and colloidal sulphur was replaced by 10 g Γ^1 sulphur powder. Inoculation was performed with 2% (v/v) of the chimney sample suspension. The same temperature and pH (60°C, pH 6.5) as in the bioreactor were used. Cell pellets dedicated to DNA extractions were recovered from 15 ml of culture after 24 h (sample A1) and 41 h (sample A2) of incubation. The 24 h culture (A1) was subcultured for 17 h in vial in the same conditions (sample B).

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141 Nucleic acid extraction

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143 DNA was extracted from frozen cell pellets in lysis buffer (cells recovered from 15 ml of culture, see procedure 144 above). A modified version of the protocol described by Alain et al. (2002) was followed combining chemical 145 and enzymatic lysis. For these culture samples, the applied lysis treatment was 1.5 h and 2.5 h. Afterwards, intact cells could not be observed by microscopy. Supernatants from the 1.5 h and 2.5 h lysis were pooled and 146 147 extracted twice with equal volumes of buffered (pH 8.0) PCI (phenol/chloroform/isoamylic acid : 25/24/1) and 148 once with an equal volume of chloroform. DNA was finally precipitated by addition of 70 % (v/v) isopropanol. 149 After centrifugation at 11,000×g for 30 min, DNA was air dried before being resuspended in 250 μ l TE 1× buffer (10 mM Tris-HCl, 2 mM EDTA, pH 7.5). The extracted DNA quality was routinely checked using 0.8% 150 151 agarose-TAE-1× gels.

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153 Amplification of the 16S rRNA gene and DGGE analysis

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The variable v3 region of 16S rRNA genes from extracted DNA were amplified using the primers 341F-GC and 907R specific to the bacterial domain (Muyzer et al. 1993; Muyzer and Smalla 1998). The PCR procedure and the analysis of the fragments by DGGE using the Bio-Rad Dcode apparatus are described in Muyzer et al. (1993). Electrophoresis conditions, gel staining, DGGE band extraction, DNA reamplification, and PCR product purification were performed using the conditions described by Postec et al. (2005b).

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161 Amplification of the 16S rRNA gene and cloning

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Archaeal DNA was amplified using the primer A24F (5'-TTC CGG TTG ATC CTG CCG GA-3') and the reverse primer 1407R (5'-GAC GGG CGG TGW GTR CAA-3') or alternatively A23SR (5'-CTT TCG GTC

165 GCC CCT ACT-3', position 257-234 on *Thermococcus celer* 23S rRNA gene sequence). Bacterial DNA was

amplified using primer E8F (5'-AGA GTT TGA TCA TGG CTC AG-3') and the reverse primer U1492R (5'-166 GTT ACC TTG TTA CGA CTT-3'). PCR reactions were performed on a Robocycler Gradient 96 (Stratagene) 167 (Wery et al. 2002; Nercessian et al. 2003). PCR products were then checked on a 0.8% (w/v) agarose gel and 168 directly cloned using the TOPO TA Cloning[®] kit (pCR2.1 vector), according to the manufacturer's instructions 169 170 (Invitrogen). Clone libraries were constructed by transforming E. coli TOP10F' cells. An archaeal and a bacterial library were constructed from two culture samples from the enrichment culture in bioreactor named T7 and T28, 171 172 collected respectively after 7 and 28 days of culture, and from each sample of enrichment cultures in vials (A1, 173 A2 and B).

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175 16S rRNA gene sequencing and phylogenetic analysis

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177 DNA fragments obtained by DGGE were sequenced by Genome Express S.A. (Grenoble, France). From clone 178 libraries, each clone was cultivated overnight at 37°C with shaking (320 rpm) on deepwell microplates, in 1 ml Luria Bertani broth 2X medium containing ampicillin (50 µg ml⁻¹). Plasmids were extracted and purified using 179 Montage Plasmid Miniprep₉₆ Kits (Millipore) and partially sequenced using the BigDye Terminator chemistry 180 181 with an automated capillary sequencer (Applied Biosystem). Sequences were compared to all GenBank, RefSeq 182 Nucleotides, EMBL, DDBJ and PDB sequences using the BLAST (http://www.ncbi.nlm.nih.gov/BLAST) 183 network service (Altschul et al. 1990) in order to determine phylogenetic affiliations and detect chimeric 184 sequences. Alignment of 16S rRNA gene sequences was performed using the CLUSTALW program (Thompson 185 et al. 1994), then refined manually using the SEAVIEW program (Galtier 1996). Sequences displaying more than 97% similarity were considered to be related and grouped in the same phylotype. Complete 16S rRNA gene 186 187 sequences were obtained for the representative clone of each unique phylotype: the related partial sequences 188 were first assembled using the SEQMAN module of the DNASTAR software (Madison, WI, USA), and the 189 complete sequences were analysed with the BLAST program.

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191 Whole-cell hybridisation

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Fixed cells in PBS/ethanol 50% (v/v) (fixation procedure described above) were diluted if necessary and filtered
on a 0.2 µm pore size white polycarbonate filter (Isopore Membrane Filters, Millipore) laying on a nitrocellulose
membrane. After drying at room temperature, cells were hybridised with the archaeal universal probe ARCH915

(5'-GTG CTC CCC CGC CAA TTC CT-3') labelled with indocarbocyanin (Cy3) (Eurogentec) as well as with 196 the universal bacterial probe EUB338 (5'-GCT GCC TCC CGT AGG AGT-3') labelled with fluorescein-197 198 isothiocyanate (FITC) (Eurogentec). Whole-cell hybridisations were carried out at 46°C after addition of 1.5 µl ARCH915 and 1.5 µl EUB338 each at 50 ng µl⁻¹ and 12 µl hybridisation solution [NaCl 5 M: 360 µl, Tris-HCl 1 199 200 M pH 7.4: 200 µl, formamide: 400 µl, sodium dodecyl sulphate (SDS) 10%: 1 µl, deionised water: 1039 µl] onto 201 each filter. After 2 h hybridisation, the filters were rinsed up 15 min at 48°C in a wash solution (NaCl 5M: 1.8 ml, Tris-HCl 1 M pH 7.4: 5 ml, SDS 10%: 25 µl, deionised water: 43.2 ml). After the hybridisation procedure, 202 each sample was stained with 10 µl of 4',6'-diamidino-2-phenylindole (DAPI 1 µg ml⁻¹). After addition of 203 204 Citifluor (Citifluor, UK) to the filters, the hybridised cells were viewed with an Olympus microscope (BX60) 205 equipped with a UV lamp and filters for DAPI (excitation 365 nm, emission 397 nm), FITC (excitation 492 nm, 206 emission 520 nm), or Cy3 (excitation 550 nm, emission 570 nm). 207 208 Analyses of amino acids, organic acids, and glucose

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210 The HPLC procedure used for analyses of amino acids, organic acids and glucose is in Wery et al. (2001).

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212 Gas analyses

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The gas outflow from the bioreactor was directly analysed using a MTI M200D micro gas chromatograph equipped with a thermal conductivity detector. A Molecular Sieve column with argon as the carrier gas was used at a temperature of 30° C to detect H₂. CO₂ and H₂S were detected with a Poraplot U column at 100° C, with helium as the carrier gas.

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219 Subcultures and isolations

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The media employed for the subcultures were designed to cultivate heterotrophic as well as autotrophic microorganisms that might be expected to grow with different electron acceptors (sulphur, nitrate, sulphate). Isolation of strains whose phylotypes were recovered in the clone libraries was attempted from the enrichment culture samples T7 and T28 from the bioreactor. Four culture media were used. The enrichment medium was used as described above and also according to the three following modifications. Vitamins and minerals were

226	preserved and organic substrates were replaced by: (1) $NH_4Cl \ 0.03 \ g \ \Gamma^1$, acetate 0.016 g Γ^1 , yeast extract 0.01 g Γ
227	¹ , and sulphur 5 g Γ^1 in the DS medium, or (2) NaNO ₃ 0.2 g Γ^1 in the DN medium, or (3) Na ₂ SO ₄ 0.3 g Γ^1 in the
228	T medium. The DS, DN and T media were adjusted at pH 6.5. The DS medium was sterilized by tyndallisation
229	(twice 30 min at 100°C) while the DN and T media were autoclaved (20 min at 121°C). In the anaerobic
230	chamber, the media were reduced with Na_2S (final concentration: 0.05 g Γ^1), then aliquoted into Hungate tubes
231	or penicillin vials under $N_2/H_2/CO_2$ (90:5:5). The gas phase was then replaced performing 10 cycles of vacuum
232	extraction / addition of H_2/CO_2 (80/20, v/v, 2 bar). All the incubations were performed at 60°C and pH6. Strains
233	were isolated by repeated dilutions-to-extinction cultures or streaking on solidified enrichment medium.
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235	Nucleotide sequence accession numbers
236	
237	The sequence data used in this study have been submitted to the EMBL databases under accession number
238	AJ874300 to AJ874328.
239	
240	Results
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242	Monitoring of the continuous enrichment culture in bioreactor: cell morphologies, whole-cell hybridisation,
243	DGGE, gas chromatography and HPLC
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245	Starting from 2.7 10 ⁶ cell ml ⁻¹ at T0, cell density reached 7.4 10 ⁸ cell ml ⁻¹ at T2 and its maximal value 2.2 10 ⁹
246	cell ml ⁻¹ at T31 (Fig. 1). Coccoid cells single or in pairs were dominant at T2 (Fig. 2a, 2d). Afterwards rods
247	displaying various morphologies became widely dominant (Fig. 2b, 2e). Short, rod-shaped cells appeared single
248	or in chains within an outer sheath-like structure, similarly to the specific toga of Thermotogales. Long rods
249	exhibiting a terminal endospore were observed. From T27, coccoid cells, single or in pairs, increased in density
250	compared to the rod morphologies. At the end of the culture, rod morphologies had nearly disappeared.
251	The relative proportions of archaeal and bacterial cells were determined by whole-cell hybridisation in eleven
252	culture samples of the bioreactor, from T2 to T41 (Fig. 1). Approximately 2,000 cells were counted on filters for
253	each sample. 99.0% of the cells detected at T2 belonged to the Archaea (Fig. 2a, 2d); Bacteria were dominant
254	from T4 to T31 (between 94.8 to 99.8% until T24, then 60.0% at T28 and 54.6% at T31). At T36 and T41,

255 Archaea became predominant again, representing 61.6% of the cells at T36 and 98.8% of the cells at T41.

Temperature was increased from 60°C to 70°C at T31, and from 70°C to 80°C at T36, which was associated with a significant decrease in the cell density (2.2 10^9 cell ml⁻¹ at T31 and 1.8 10^7 cell ml⁻¹ at T41).

The forty-five days enrichment culture was further investigated by DGGE analysis. The v3 hypervariable region 258 259 of the bacterial 16S rRNA gene was amplified from T1, T4, T9, T13, T16, T24, T28, T31 and T36 culture 260 samples. No amplification was obtained from samples collected after T36, probably due to the low cell density 261 corresponding to the temperature increase at T36. Representative DGGE bands migrating at different distances and originating from various samples were extracted from gels (not shown) and re-amplified directly. The 262 263 DGGE sequence types (approximately 500 pb) were affiliated to Thermosipho MV1063 (99% identity), Marinitoga camini (96%), Caminicella sporogenes (99-98%) and Deferribacter abyssi (87%). Thermosipho spp. 264 265 was detected from T1 to T36, Marinitoga spp. at T31, Caminicella spp. from T4 to T31 and Deferribacter spp. from T1 to T31. 266

267 H₂, CO₂ and H₂S were detected in T8, T21, T30 and T38 samples by gas chromatography. H₂S production was also detected using Zn acetate strips (Lead Acetate, Whatman) from T2 until the end of the culture. From T3 268 269 until T41, HPLC analysis of the free amino acids in the culture medium showed that they were all completely 270 consumed, and might be therefore a limiting factor for microbial growth. Analysis of glucose by HPLC indicated an initial concentration of 0.440 g l⁻¹, between T4 and T32 glucose was not detected at all showing a complete 271 consumption, and endly detected again and reaching 0.389 g Γ^1 . Also organic acids as pyruvate, succinate, 272 273 lactate, formate, acetate, propionate, butyrate, isobutyrate and isovalerate were detected and assumed to be 274 metabolic end products.

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276 16S rRNA gene libraries from the continuous enrichment culture in bioreactor

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278 Archaeal and bacterial 16S rRNA genes were amplified from T7 and T28 culture samples from the continuous 279 enrichment culture in bioreactor. All the archaeal sequences from T7 and T28 were related to the genus 280 Thermococcus (Table 1). The sequence types A704 and A800 were affiliated to T. siculi, and shared more than 281 97% of identity with a large number of 16S rRNA gene sequences related to members of the group T. siculi -282 T. celer, according to the BLAST analysis. Bacterial sequences affiliated with the orders Clostridiales, 283 Thermotogales and Deferribacterales were retrieved in both T7 and T28 libraries (Table 1, Fig. 3) and were closely related to the hydrothermal species Caminicella sporogenes (97% 16S rRNA gene identity with clones 284 775 and 813), Marinitoga camini (94% identity with the clone 716, 95% with clone 805) and Deferribacter 285

abyssi (98% identity with clones 737 and 820), respectively. The proportion of clones related to *Deferribacter* spp. and *Marinitoga* spp. increased slightly at T28, while the number of clones related to *Caminicella* spp. decreased from 50% to 7% (Fig. 3). Two phylotypes were recovered only in the T28 bacterial library: (i) 12 clones were affiliated to *Thermosipho* spp. and the sequence type 840 shared 99% identity with *Thermosipho* MV1063, the closest species being *T. melanesiensis* (96% identity) and (ii) 3 clones were affiliated to *Thermodesulfatator* spp. and the sequence type 850 shared 96% identity with *T. indicus* (Table 1).

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293 16S rRNA gene libraries from the batch enrichment cultures in vials

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295 Total DNA was extracted from A1, A2 and B culture samples. Archaeal 16S rRNA gene was amplified by PCR 296 only from sample A1, corresponding to the shortest incubated enrichment culture (24 h incubation). No archaeal 297 16S rRNA gene sequences was amplified after longer incubation (A2: 41 h incubation), and after subculturing 298 from A1 (B). The 55 archaeal clones from the A1 library were all related to the genus Thermococcus (Table 1). 299 The sequence type A254 displayed 97 % identity with the closest described strain T. barophilus (AY099172), 300 also originating from the Mid-Atlantic Ridge (Snake Pit) (Marteinsson et al. 1999) and growing in the range of 301 48-95°C under atmospheric pressure. Bacterial 16S rRNA gene was amplified by PCR from all three samples. 302 Sequences related to Marinitoga spp. and Caminicella spp. were retrieved in each library. Sequences related to 303 the Thermosipho spp. were not recovered in the A1 culture, but only after 41 hours of incubation (A2) and after 304 17 h subculturing (B). Moreover, a shift was observed in the library compositions; sequences related to 305 Marinitoga spp. were widely dominant in the A1 library whereas the A2 and B libraries were largely dominated by sequences related to Caminicella spp. (Fig. 3). Extending the incubation time or subculturing from the 306 307 primary A1 culture resulted in similar changes in the composition of the bacterial libraries.

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309 Subcultures and isolations

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Several strains were isolated from culture samples (T3, T7 and T28) from the bioreactor: (i) an archaeal strain *Thermococcus* spp. designated as AT1273 (99% 16S rRNA gene similarity with *Thermococcus siculi*) (ii) a strain *Thermosipho* spp. (order *Thermotogales*) designated as AT1272 (98% similarity with *Thermosipho* MV1063, 95% with *T. melanesiensis*) (iii) a new bacterial species of the *Marinitoga* genus (order *Thermotogales*) named *M. hydrogenitolerans* (Postec et al. 2005c); (iv) using the T medium, a new bacterial 316 species among the *Thermodesulfatator* genus capable of sulphate-reduction (96% similarity with the 317 hydrothermal species *T. indicus*), (v) using the DS and DN media, we isolated a strain closely affiliated to the 318 hydrothermal species *Deferribacter abyssi* (99% identity).

319 320

321 Discussion

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In this study, we used an original culture method to enrich thermophilic microorganisms from a hydrothermal black smoker: a continuous culture was performed in a gas-lift bioreactor during 45 days at 60°C and pH 6.5 under anaerobic conditions. The microbial diversity in continuous culture and classical batch cultures in vials was compared.

327

328 Methodological considerations

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The molecular inventories gave snapshots of the microbial diversity on a restricted number of samples while DGGE and whole-cell hybridisation revealed a temporal dynamics in the continuous culture. The use of different molecular techniques based on 16S rRNA gene analysis gave complementary data. For example, sequences of *Thermosipho* spp. were detected by DGGE at T7 in the bioreactor but were not evidenced by cloning. Inversely, *Thermodesulfatator* spp. was detected by cloning but not by DGGE. The possible limitations of primer selectivity and cloning biases (Theron and Cloete 2000) or PCR biases (Suzuki and Giovannoni 1996) are well established and can explain the variability in results from different methods.

337 Considering the gas-lift bioreactor as a system for continuous cultivation, the potential adhesion of bacteria on the inner wall and the formation of a biofilm have to be examined, since the bioreactor includes no device to 338 339 clean surfaces. From a precipitate sampled on the inner wall, no cells were detected by microscopy observation. 340 Culture attempt in vial and DNA extraction failed as well. X-RD analysis indicated that the precipitate was 341 mainly composed of sulphurs (data not shown). The microorganisms detected in this study represent therefore cells in suspension. The dilution rate applied in the bioreactor after 34 h of batch culture was 0.04 h^1 and 342 343 corresponded to a generation time of 17.25 h. Although the continuous culture involves a progressive dilution of the medium inside the bioreactor, four volume changes, corresponding to 100 h (about 4 days) at a dilution rate 344 of 0.04 h⁻¹, have been considered sufficient to completely renew the culture medium inside the bioreactor (Raven 345

et al. 1992). Thus, microorganisms thriving in the continuous culture from T6 were not washed-out and should have grown with a minimal growth rate of 0.04 h^{-1} .

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349 *Thermococcales* as early heterotrophic colonizers

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351 Thermococcales at deep-sea hydrothermal vents are widespread and members of the genus Thermococcus are 352 some of the most numerous hyperthermophiles described from deep-sea vents. Investigation of their natural 353 distribution showed that a viable *Thermococcus* population was present in the surface layers of mature 354 hydrothermal chimneys (Harmsen et al. 1997; Takai et al. 2001; Schrenk et al. 2003). The early growth of 355 Thermococcales was observed in enrichment cultures from hydrothermal chimney performed in vials and bioreactor at 90°C (Postec 2005b) and at 60°C (this study). Although all the members of the Thermococcales are 356 357 hyperthermophilic, their early growth at 60°C both in the bioreactor and in vials is possible since (i) some Thermococcus sp. are able to grow at 60°C (Godfroy et al. 1997), (ii) an absence of latency phase could explain 358 359 the early growth (T. hydrothermalis; Godfroy, pers. com.), (iii) the number of Thermococcus-related cells might 360 be abundant in the chimney sample used as inoculum. This is also suggested by the detection of Thermococcus 361 spp. in the molecular inventory performed directly on the studied chimney sample, while the bacteria grown in 362 this study were not detected (Postec 2005d). Their growth at 60°C at the beginning of the enrichment culture and 363 then when temperature was risen from 60 to 80°C after T31 may confer on Thermococcales a great ecological 364 advantage to colonize new hydrothermal environments and they may be the first heterotrophs colonizing this 365 ecosystem. This idea is supported by a recent study of nascent vent colonization in which protochimneys were deployed for short time on hydrothermal vents and heterotrophic groups including Thermococcales dominated 366 the colonization of mineral surfaces after 72h (McCliment et al. 2006). The temporal sequence of colonization 367 was analysed in the study of *in situ* samplers deployed on hydrothermal vents: the widespread occurrence of 368 369 Thermococcales was demonstrated in short deployments (4-7 days) and decreased with time, suggesting that this 370 group is an early surface colonizer (Nercessian et al. 2003).

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372 Bacterial diversity and insight into microbial interactions

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All the bacteria cultivated in this study were related to microorganisms from deep-sea hydrothermal vents. *Caminicella* spp. and *Marinitoga* spp. were detected in both vials and bioreactor, while the autotrophs *Deferribacter* spp. and *Thermodesulfatator* spp. were only detected in the bioreactor. Compared to batch cultures
in vials, a larger diversity was described in the enriched community in bioreactor continuous supply of nutrients,
gaseous inhibitory by-products removal and and pH regulation over time, what confirms previous results (Postec
et al. 2005b).

380 Guided by the results of the molecular analysis, we succeeded in isolating microorganisms (heterotrophs and 381 autotrophs) enriched in continuous in the gas-lift bioreactor. Three new species belonging to the genera 382 Thermosipho, Marinitoga and Thermodesulfatator were obtained in pure culture in vials. They were related to T. 383 melanesiensis and M. camini, both originating from Atlantic deep sea vents, and T. indicus, isolated from the Central Indian Ridge, respectively. Marinitoga sp. nov. was recently characterised and named M. 384 385 hydrogenitolerans because its growth is not inhibited by high hydrogen concentrations (Postec et al. 2005c). The 386 genus Thermodesulfatator is only represented so far by the species T. indicus, a thermophilic, anaerobic and 387 strictly chemolithoautotrophic bacterium growing exclusively with CO₂ as sole carbon source, H₂ as sole 388 electron donor and sulphate as sole electron acceptor (Moussard et al. 2004). A second chemolithoautotrophic to 389 mixotrophic strain isolated from the bioreactor shared 99% 16S rRNA gene similarity with Deferribacter abyssi 390 (Miroshnichenko et al. 2003). D. abyssi is thermophilic, anaerobic and facultative chemolithoautotrophic using 391 elemental sulphur or nitrate as electron acceptors, similarly to the strain isolated in this study.

In the enrichment culture, growth of *Thermococcales* clearly occurred first. Their fermentative metabolism on proteinaceous substrates and in smaller extent carbohydrates through sulphur reduction might have generated a propitious environment for the growth of bacterial heterotrophs *Caminicella* spp., *Thermosipho* spp. and *Marinitoga* spp) and then autotrophs (*Deferribacter* spp. and *Thermodesulfatator* spp.). The growth of *Thermosipho* sp. and *Marinitoga* sp. is correlated with the diminution of the glucose concentration in the medium and species of these genera are known to be able to use glucose as carbon substrate and to produce acetate as end metabolic product (Antoine et al. 1997; Alain et al. 2002b; Postec et al. 2005c).

Indeed 16S rRNA gene sequences related to the autotrophic microorganisms were detected by molecular analysis late at T28 but not at T7. Similarly, a temporal study of *in situ* collectors deployed on deep-sea hydrothermal vents showed that the microbial diversity of the colonizing community increased with time and that chemolithoautotrophs emerged during late stages (Nercessian et al. 2003).

The late growth of chemolithotrophs in co-culture with heterotrophs may be explained by interactions between species by means of metabolites exchanges. Organic carbon provided by the medium supported the growth of heterotrophs, afterwards the chemolithotrophs utilized carbon dioxide or acetate as carbon source and hydrogen 406 and acetate as possible electron donor, both compounds being end-products of fermentation. The study of a 407 natural community showed that acetate and a range of other organic electron donors can be oxidised under sulphate-reducing conditions in hydrothermal vents at high temperature (90°C) (Tor et al. 2003). It has been 408 409 suggested that acetate and hydrogen are the most prevalent organic fermentation products and important 410 extracellular intermediates in the degradation of organic matter in hot microbial ecosystems, and that cooperative 411 activity between fermentative microorganisms and sulphate reducers is important for the metabolism of 412 fermentable compounds. Syntrophic interactions can also have an impact in hyperthermophilic co-cultures, for 413 example on the metabolism of heterotrophic microorganisms co-cultivated with methanogens (Johnson et al. 414 2006).

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416 Conclusions

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418 A gas-lift bioreactor was used to cultivate in continuous a thermophilic microbial community from a deep sea 419 hydrothermal chimney, on an organic-rich medium with sulphur under anaerobic conditions. The enrichment 420 culture was monitored with molecular and chemical analyses. In the long-term running culture, the cultivated 421 populations were evidenced to continuously evolve with time, instead of reaching a stationary state. 422 Thermococcales dominated in the first hours of the enrichment cultures suggesting that this group early 423 colonizes hydrothermal edifices, and may represent the first heterotrophic colonizers. A larger diversity was 424 detected in the enrichment culture in bioreactor compared to culture in vials and most of the microorganisms 425 enriched in bioreactor, including three new bacterial species, were successfully isolated by subculturing in vials. Results indicate that the continuous culture in a gas-lift bioreactor, combined with the use of molecular tools, 426 427 could be of further use to access the "uncultivated" microbial community. The microorganisms isolated in this study displayed a phylogenetic and metabolic diversity. They are involved in the sulphur cycle (suphur- and 428 429 sulphate-reduction) and in the carbon cycle (autotrophy and heterotrophy). Autotrophic microorganisms were 430 enriched in co-culture with heterotrophs, suggesting that prokaryotic species interact by means of metabolite 431 exchange to support the growth of autotrophs. This emphasizes the importance of microbial interactions with 432 surrounding microorganisms, animals or minerals within ecological niches. Inter-species interactions should be 433 further taken into account to attempt the growth of as-yet uncultivated microorganisms and microbial 434 metabolism need to be examined inside communities rather than extrapolated from pure cultures (For et al. 435 2003). The bioreactor can be considered as a window to investigate *in vitro* interactions between population

436 interactions that may occur *in situ*. Considering that deep-sea hydrothermal vents are extreme environments 437 encompassing intense thermal and chemical gradient (Karl 1995), the microbial communities inhabiting these 438 disturbed systems must be strongly affected by environmental changes. The gas-lift bioreactor represents a 439 promising tool to investigate *in vitro* the effect of physico-chemical perturbations on the microbial community 440 structure.

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- 448 **References**
- 449
- Alain K, Olagnon M, Desbruyeres D, Page A, Barbier G, Juniper SK, Querellou J, Cambon-Bonavita MA
 (2002) Phylogenetic characterization of the bacterial assemblage associated with mucous secretions of the
 hydrothermal vent polychaete *Paralvinella palmiformis*. FEMS Microbiol Ecol 42:463-476
- Altschul SF, Gish W, Miller W, Meyers EW, Lipman DJ (1990) Basic Local Alignment Search Tool. J Mol Biol
 215:403-410
- DeChaine EG, Bates AE, Shank TM, Cavanaugh CM (2006) Off-axis symbiosis found: characterization and
 biogeography of bacterial symbionts of *Bathymodiolus* mussels from Lost City hydrothermal vents. Environ
 Microbiol 8:1902-1912
- Galtier N, Gouy M, Gautier C (1996) SEAVIEW and PHYLO_WIN: two graphic tools for sequence alignment
 and molecular phylogeny. CABIOS 12:543-548
- Godfroy A, Meunier JR, Guezennec J, Lesongeur F, Raguénès G, Rimbault A, Barbier G (1996) *Thermococcus fumicolans* sp. nov. a new hyperthermophilic archaeum isolated from deep-sea hydrothermal vent in North
 Fiji bassin. Int J Syst Bacteriol 46:1113-1119
- Godfroy A, Lesongeur F, Raguénès G, Quérellou J, Antoine E, Meunier JR, Guezennec J, Barbier G (1997)
 Thermococcus hydrothermalis sp. nov., a new hyperthermophilic archaeon isolated from deep-sea
 hydrothermal vent. Int J Syst Bacteriol 47:622-626
- Godfroy A, Raven NDH, Sharp RJ (2000) Physiology and continuous culture of the hyperthermophilic deep-sea
 vent archaeon *Pyrococcus abyssi* ST549. FEMS Microbiol Lett 186:127-132
- Godfroy A, Postec A, Raven NDH (2005) Growth of hyperthermophilic microorganisms for physiological and
 nutritional studies. In: Rainey FA, Oren A (eds) Methods in Microbiology, Extremophiles. Academic Press,
 Oxford, pp 91-106
- Harmsen HJM, Prieur D, Jeanthon C (1997) Distribution of microorganisms in deep-sea hydrothermal vent
 chimneys investigated by whole-cell hybridization and enrichment culture of thermophilic subpopulations.
 Appl Environ Microbiol 63:2876-2883
- Inagaki F, Kuypers MMM, Tsunogai U, Ishibashi JI, Nakamura KI, Treude T, Ohkubo S, Nakaseama M, Gena
 K, Chiba H, Hirayama H, Nunoura T, Takai K, Jorgensen BB, Horikoshi K, Boetius A (2006) From the
 Cover: Microbial community in a sediment-hosted CO₂ lake of the southern Okinawa Trough hydrothermal
 system. PNAS 103:14164-14169
- Johnson MR, Conners SB, Montero CI, Chou CJ, Shockley KR, Kelly RM (2006) The *Thermotoga maritima* phenotype is impacted by syntrophic interaction with *Methanococcus jannaschii* in hyperthermophilic
 coculture. Appl Environ Microbiol 72:811-818
- Karl DM (1995) Ecology of free-living hydrothermal vent microbial communities. In: Karl DM (eds) The
 microbiology of deep-sea hydrothermal vents. CRC Press, pp 35-125
- 483 Kormas KA, Tivey MK, Von Damm K, Teske A (2006) Bacterial and archaeal phylotypes associated with
 484 distinct mineralogical layers of a white smoker spire from a deep-sea hydrothermal vent site (9°N, East
 485 Pacific Rise). Environ Microbiol 8:909-920
- López-García P, Duperron S, Philippot P, Foriel J, Susini S, Moreira D (2003) Bacterial diversity in
 hydrothermal sediment and epsilon-proteobacterial dominance in experimental microcolonizers at the Mid Atlantic Ridge. Environ Microbiol 5:961-976
- Marteinsson V, Birrien J, Reysenbach A, Vernet M, Marie D, Gambacorta A, Messner P, Sleytr U, Prieur D
 (1999) *Thermococcus barophilus* sp. nov., a new barophilic and hyperthermophilic archaeon isolated under
 high hydrostatic pressure from a deep-sea hydrothermal vent. Int J Syst Bacteriol 49:351-359
- McCliment EA, Voglesonger KM, O'Day PA, Dunn EE, Holloway JR, Cary SC (2006) Colonization of nascent,
 deep-sea hydrothermal vents by a novel archaeal and nanoarchaeal assemblage. Environ Microbiol 8:114-125
- Miroshnichenko ML, Slobodkin AI, Kostrikina NA, L'Haridon S, Nercessian O, Spring S, Stackebrandt E,
 Bonch-Osmolovskaya EA, Jeanthon C (2003) *Deferribacter abyssi* sp. nov., an anaerobic thermophile from
 deep-sea hydrothermal vents of the Mid-Atlantic Ridge. Int J Syst Evol Microbiol 53:1637-1641
- Moussard H, L'Haridon S, Tindall BJ, Banta A, Schumann P, Stackebrandt E, Reysenbach AL, Jeanthon C (2004) *Thermodesulfatator indicus* gen. nov., sp. nov., a novel thermophilic chemolithoautotrophic sulfate-reducing bacterium isolated from the Central Indian Ridge. Int J Syst Evol Microbiol 54:227-233
- Moussard H, Corre E, Cambon-Bonavita MA, Fouquet Y, Jeanthon C (2006) Novel uncultured
 Epsilonproteobacteria dominate a filamentous sulphur mat from the 13°N hydrothermal vent field, East
 Pacific Rise. FEMS Microbiol Ecol 58:449-463
- Muyzer G, Smalla K (1998) Application of denaturing gradient gel electrophoresis (DGGE) and temperature
 gradient gel electrophoresis (TGGE) in microbial ecology. Ant van Leeuw 73:127-141

- Muyzer G, De Waal EC, Uitterlinden AG (1993) Profiling of complex microbial populations by denaturing
 gradient gel electrophoresis analysis of polymerase chain reaction- amplified genes coding for 16S rRNA.
 Appl Environ Microbiol 59:695-700
- Nercessian O, Reysenbach AL, Prieur D, Jeanthon C (2003) Archaeal diversity associated with *in situ* samplers
 deployed on hydrothermal vents on the East Pacific Rise (13°N). Environ Microbiol 5:492-502
- Pace NR, Stahl DA, Lane DJ, Olsen GJ (1986) The analysis of natural microbial populations by ribosomal RNA
 sequences. Adv Microbial Ecol 9:1-55
- Postec A, Pignet P, Cueff-Gauchard V, Schmitt A, Querellou J, Godfroy A (2005a) Optimisation of growth
 conditions for continuous culture of the hyperthermophilic archaeon *Thermococcus hydrothermalis* and
 development of sulphur-free defined and minimal media. Res Microbiol 156:82-87
- Postec A, Urios L, Lesongeur L, Ollivier B, Querellou J, Godfroy A (2005b) Continuous enrichment culture and
 molecular monitoring to investigate the microbial diversity of thermophiles inhabiting the deep-sea
 hydrothermal ecosystems. Curr microbiol 50:138-144
- Postec A, Le Breton C, Fardeau ML, Lesongeur F, Pignet P, Querellou J, Ollivier B, Godfroy A (2005c)
 Marinitoga hydrogenitolerans sp. nov., a novel member of the order *Thermotogales* isolated from a black
 smoker chimney on the Mid-Atlantic Ridge. Int J Syst Evol Microbiol 55:1217-1221
- Postec A (2005d) Diversité de populations microbiennes thermophiles d'une cheminée hydrothermale océanique:
 cultures d'enrichissement en bioréacteur et isolement d'espèces nouvelles. In: Thesis, Université de Provence,
 France
- Raven N, Ladwa N, Sharp R (1992) Continuous culture of the hyperthermophilic archaeum*Pyrococcus furiosus*.
 Appl Microbiol Biotechnol 38:263-267
- Raven NDH, Sharp RJ (1997) Development of defined and minimal media for the growth of the
 hyperthermophilic archaeon *Pyrococcus furiosus* Vc1. FEMS Microbiol Lett 146:135-141
- Schrenk MO, Kelley DS, Delaney JR, Baross JA (2003) Incidence and diversity of microorganisms within the
 walls of an active deep-sea sulfide chimney. Appl Environ Microbiol 69:3580-3592
- Sharp RJ, Raven NDH (1997) Isolation and growth of hyperthermophiles. In: Rhodes PM, Stanbury PF (eds)
 Applied microbial physiology: a practical approach. IRL Press, Oxford, pp 23-51
- Suzuki M, Giovannoni S (1996) Bias caused by template annealing in the amplification of mixtures of
 16S rRNA genes by PCR. Appl Environ Microbiol 62:625-630
- Takai K, Komatsu T, Inagaki F, Horikoshi K (2001) Distribution of *Archaea* in a black smoker chimney
 structure. Appl Environ Microbiol 67:3618-3629
- Theron J, Cloete TE (2000) Molecular techniques for determining microbial diversity and community structure
 in natural environments. Crit Rev Microbiol 26:37-57
- Thompson J, Higgins D, Gibson T (1994) CLUSTAL W: improving the sensitivity of progressive multiple
 sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice.
 Nuc Acids Res 22:4673-4680
- Tor JM, Amend JP, Lovley DR (2003) Metabolism of organic compounds in anaerobic, hydrothermal sulphate reducing marine sediments. Environ Microbiol 5:583-591
- Wery N, Cambon-Bonavita MA, Lesongeur F, Barbier G (2002) Diversity of anaerobic heterotrophic
 thermophiles isolated from deep-sea hydrothermal vents of the Mid-Atlantic Ridge. FEMS Microbiol Ecol
 41:105-114
- Wery N, Lesongeur F, Pignet P, Derennes V, Cambon-Bonavita MA, Godfroy A, Barbier G (2001) Marinitoga
 camini gen. nov., sp. nov., a rod-shaped bacterium belonging to the order *Thermotogales*, isolated from a
- 548 deep-sea hydrothermal vent. Int J Syst Evol Microbiol 51:495-504
- 549



Fig. 1 Total cell densities of the bioreactor culture from T2 to T41 expressed in cell ml⁻¹ and Archaea and Bacteria ratio expressed in percentages of total cells determined by whole-cell hybridisation using the universal probes ARC915 and EUB338 respectively. The temperature was increased from 60°C to 70°C at T31 and from 70°C to 80°C at T36.





Fig. 2 Whole-cell hybridisation of fixed cells from 3 samples of the enrichment culture in bioreactor: T2 (a, d), T7 (b, e) and T28 (c, f). Cells were stained with DAPI (a, b and c), and hybridised with the FITC-labelled Eub338 probe (e), or the Cy-3-labeled Arch 915 (d and f). Cells were viewed by epifluorescence microscopy in which DAPI-, FITC- and Cy-3-specific filters were used. The relative proportions of archaeal and bacterial cells were determined by counting approximately 2,000 cells on filter and resulted in: 99.0% of the cells detected at T2 belonged to the *Archaea* (d), 99.0% of the cells detected at T7 belonged to the *Bacteria* (e), 40% of the cells detected at T28 belonged to the *Archaea* (f). Scale bars = $10 \mu m$.



Fig. 3 Composition of the bacterial clone libraries from enrichment cultures in vials (A1: 24 hours culture, A2:
41 hours culture and B: 17 hours subculture from A1) and from the enrichment culture in bioreactor (T7: 7 days
culture, T28: 28 days culture). The percentages of clones of each phylogenetic group are indicated on the piecharts.

Table 1. Distribution and phylogenetic affiliations of archaeal and bacterial 16S rDNA sequences ≥ 97%
similarity in each phylotype) from the enrichment culture in bioreactor and from enrichment cultures in flask
both performed at 60°C. Representative clones were completely sequenced. The sequence types deposited in
GenBank appear in bold.

	Culture sample	Phylogenetic affiliation	Representative clones	Number of clones	Closest match organism ^a	Identity ^a (%)
	Τ7	Thermococcales	A704- A710-A712- A715-	76	Thermococcus siculi (AY099185)	98
			A730-A732-A737-A739			
	T28	Thermococcales	A800 -A811- A816-A817	71	Thermococcus siculi (AY099185)	98
	Τ7	Clostridiales	775- 700-725-728-750-770-	38	Caminicella sporogenes (AJ320233)	97
2		Thermotogales	716 -705-706-709-724-768	29	Marinitoga camini (AJ250439)	94
aci		Deferribacterales	737 -711-740-754-769	9	Deferribacter abyssi (AJ515882)	98
Ire	T28	Clostridiales	813 -874	5	Caminicella sporogenes (AJ320233)	97
i		Thermotogales	805-802-822-823-832	35	Marinitoga camini (AJ250439)	95
р		0	840-812-821-825-868-884-	12	Thermosipho MV1063 (AJ419874)	99
		Deferribacterales	893	17	Deferribacter abyssi (AJ515882)	98
		Thermodesulfo-	820 -829-856	3	Thermodesulfatator indicus	96
		bacteriales	850 -816-858		(AF393376)	
	A1	Thermococcales	A254	55	Thermococcus barophilus (AY099172)	97
	A1	Clostridiales	238	1	Caminicella sporogenes (AI320233)	97
		Thermotogales	207-245-219-240-255	58	Marinitoga camini (AJ250439)	94
Sk	A2	Clostridiales	413 -404-462	47	Caminicella sporogenes (AJ320233)	97
la		Thermotogales	436	8	Marinitoga camini (AJ250439)	95
			440^b	1	Thermosipho MV1063 (AJ419874)	98
	В	Clostridiales	608 -626-669-635	68	Caminicella sporogenes (AJ320233)	97
		Thermotogales	660 ^b	1	Marinitoga camini (AJ250439)	93
		0	609	1	Thermosipho MV1063 (AJ419874)	97

^a based on BLAST search. GenBank accession numbers are in brackets.

^b partial sequence (600pb)