

THERMOPHILES FROM DEEP-SEA HYDROTHERMAL VENTS

J Quérellou, K Alain, M A Cambon-Bonavita

▶ To cite this version:

J Quérellou, K Alain, M A Cambon-Bonavita. THERMOPHILES FROM DEEP-SEA HYDROTHER-MAL VENTS. Vie et Milieu / Life & Environment, 2001, pp.161-172. hal-03192126

HAL Id: hal-03192126 https://hal.sorbonne-universite.fr/hal-03192126v1

Submitted on 7 Apr 2021

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

THERMOPHILES FROM DEEP-SEA HYDROTHERMAL VENTS

J.QUÉRELLOU*, K. ALAIN, M.A. CAMBON-BONAVITA

Laboratoire de Microbiologie et Biotechnologie des Extrêmophiles, Ifremer, Plouzané, France corresponding author: jquerell@ifremer.fr

THERMOPHILES HYDROTHERMAL VENT ARCHAEA THERMOSTABILITY ABSTRACT. – Deep-sea hydrothermal vents represent extreme environments where original microbial communities have evolved during geological times. Of particular interest are the thermophilic and hyperthermophilic microbes thriving in the various microhabitats determined by the geochemical processes at work in the hydrothermal vent fields. In the present review, the general characteristics of the hydrothermal vents as habitats for thermophiles and hyperthermophiles are described. The microbial diversity is discussed on the basis of molecular and cultivation approaches. The most original features of thermophily at molecular level are briefly reviewed. Data gathered during the last two decades have substantially modified our understanding of diversity and evolution of deep-sea hydrothermal vent communities and ecology.

THERMOPHILES SOURCES HYDROTHERMALES ARCHAEA THERMOSTABILITÉ RÉSUMÉ. – Les sources hydrothermales profondes constituent des environnements extrêmes où des communautés bactériennes originales se sont développées au cours des temps géologiques. Les thermophiles et hyperthermophiles colonisant les divers microhabitats déterminés par les processus géochimiques prévalant dans ces milieux présentent un intérêt particulier. Les caractéristiques principales de ces habitats sont passées en revue. Les connaissances sur la diversité microbienne, estimées à la fois par les approches moléculaires et de culture, sont discutées. Les aspects les plus originaux de l'adaptation à la thermophilie sont brièvement décrits. Les données accumulées au cours des deux dernières décades ont considérablement modifié notre compréhension de la diversité et de l'évolution des communautés microbiennes des sources hydrothermales profondes et de leur écologie.

INTRODUCTION

Since the discovery of microbial life in Yellowstone hot springs by Thomas Brock more than 30 years ago, it has become more and more evident that life is present in a great variety of extreme environments. In an anthropocentric view, most of these extreme environments were unsuitable for microbial life and this discovery opened the way to a new era for microbiologists. Inventorying a priori hostile environments demonstrated that most of them were not only able to tolerate some sporadic microbial forms of life, but constitute the selective habitats of a great variety of microorganisms belonging to the 2 domains defined by Woese (1977): Archaea and Bacteria. These microorganisms have been called 'extremophiles'. Considering the pa-rameters controlling their optimal growth, they are named thermophiles (>60°C), hyperthermophiles

(>80°C), psychrophiles, acidophiles, alcalophiles, barophiles or halophiles. Not only do they tolerate extreme environments but these extreme conditions are required for proper growth of microorganisms that have been selected over geologic time scale. Generally, they cannot grow in conditions suitable for eukaryotic forms of life. Following the work of Brock on terrestrial hot springs and thermophiles (Brock 1969), the discovery of deep-sea vents at the end of the 1970s boosted the research of the most extreme microorganisms. Among the different teams contributing to the research on extremophiles biodiversity, a remarkable contribution to the description of new genera and new species of hyperthermophiles is due to Stetter and his colleagues at University of Regensburg, Germany (Stetter 1998, Stetter 1999a,b). Besides these recent reviews, the present one focuses more specifically on hyperthermophiles from deep-sea hydrothermal vents.

Hydrothermal vents as habitats for thermophiles and hyperthermophiles

Thermophiles and hyperthermophiles have been isolated from various habitats: terrestrial hot springs, solfatara and volcano acidic hot springs, shallow marine hot springs, submarine vents, deepsea sediments and deep-sea hydrothermal vents. Moreover, evidence of the existence of microorganisms in the deep terrestrial and oceanic subsurface has been increasing during the last decade (Delaney et al. 1998, Summit & Baross 2001). Since the temperature and pressure of subsurface environments increase with depth, they are potentially among the most appropriate habitats for hyperthermophiles. An exhaustive list of thermophiles and hyperthermophiles isolated from these various environments as well as related references can be found in a recent review (Amend & Shock 2001) and on a dedicated web site at http: //levee.wustl.edu/~chan/Research/research.html.

Geochemical mechanisms at work on mid-oceanic ridges result from the interaction between seawater and volcanic rocks and from seawater and hydrothermal fluxes of heat and dissolved matter. Since their discovery in 1977 at the Galapagos Spreading Center (GSC) at 2500 m water depth (Hessler & Smithey 1983), it has been shown that seafloor hydrothermal vents constitute, in almost every mid-oceanic ridges investigated, ecosystems supporting important biomass and high productivity in contrast with other deep sea environments. Faunal assemblages, similar to those discovered at GSC, have been sampled at various sites on the East Pacific Rise (EPR) (Lutz & Kennish 1993), Juan de Fuca Ridge (Tunnicliffe 1988), Mid-Atlantic Ridge (MAR) (Van Dover 1995), and the spreading centres of Western and South-Western Pacific Back Arc Basin (Desbruyères et al. 1994, Hessler et al. 1987). Numerous active sites are still under investigation along the mid-ocean ridges in the Pacific Ocean, the Atlantic Ocean and more recently in the Indian Ocean (Fig. 1). Intensive research has been conducted on different vent fields at East Pacific Rise from 26°S to 21°N and at Mid-Atlantic Ridge between 15 and 40°N. Currently eight vent fields have been described from MAR, including Mount Saldanha (36°N) discovered in 1998 during the Portuguese-French SALDANHA cruise (Desbruyères et al. 2000). Most known hydrothermal vents along the mid-ocean ridges are located on young crust where the cooling of hot basaltic material drives hydrothermal flow (Fornari & Embley 1995). Recently an off-axis hydrothermal vent field named "Lost City" has been discovered near the Mid-Atlantic Ridge at 30°N at nearly 15 km from the spreading axis on a 1.5 Myr old crust, demonstrating that hydrothermal venting occurs not only along mid-ocean ridges, but also on old regions of the oceanic crust away from spread-

ing centres (Kelley et al. 2001). These last findings indicate that a much larger portion of the oceanic crust may support hydrothermal activity and microbial life than previously thought. On mid-ocean ridges, at depth ranging from 1 500 to 4 000 m, hydrothermal fluids can reach temperatures up to 400°C whereas ambient sea water is around 2°C, producing steep temperature gradients within diffusion structures. Hydrothermal solutions reaching the seafloor contain various products resulting from hot acidic seawater and volcanic rocks: metals (Fe, Mn, Zn, Cu, etc.) and reduced sulphur (Zierenberg et al. 2000). Most of the hydrothermal fluids display physico-chemical compositions that are unsuitable for eukaryotic organisms due to their high toxicity. However, active chimneys (white and black smokers) are densely populated, hosting not only microorganisms belonging to Bacteria and Archaea, but also a very rich macrofauna typical of these environments (Desbruyères & Segonzac 1997). Strikingly, fluids of "Lost City" off-axis vent field were found to be relatively cool (40-75°C) and alkaline (pH 9.0-9.8), supporting dense microbial communities including anaerobic thermophiles, but rare macrofaunal assemblages that typify most vent environments (Kelley et al. 2001).

Hydrothermal vent ecosystems stability is directly dependent upon the variations in fluid fluxes and chemical composition of the fluids. Microbial blooms are observed at the initiation of magmatic cycle, suggesting that the upper oceanic crust is inoculated with microbial communities (Zierenberg et al. 2000). Volcanic events initially are followed by relative increase in fluid temperature and magmatic gases, often accompanied by decrease salinity in hydrothermal fluids as a result of supercritical phase separation in seawater in the subsurface. The high-salinity components of the phase separated fluids reaches the seafloor a few years later (Von Damm 1995). Direct observation of discontinuity in vent emission has been done in different ridge dynamic contexts (Desbruyères et al. 2000). High variability in fluid composition at a same vent according to volcanic activity has been reported. In some cases, like at Lucky Strike (MAR 37°N), variability in chemical composition suggests the existence of two different sources (Von Damm et al. 1998). As an example, the concentration of most of the major chemical components in MAR vent fluid end-members is given in table I (Desbruyères et al. 2000). Despite the important differences between sites, it can be noticed that H_2S , CO_2 , NH_4 and CH_4 are present in vent fluids and are available as primary compounds for chemolithoautotrophic microbes essential to the good running of the ecosystems. Another source of variability between different vent fields, acting as a major ecological factor, is the particulate (mineral and organic) content of the fluids. Finally, these

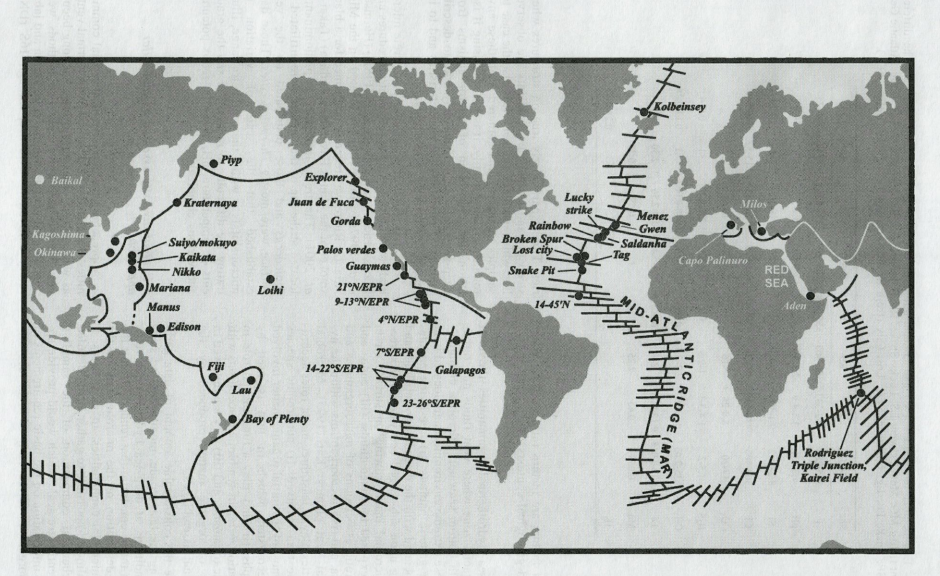


Fig. 1. – Distribution of deep-sea hydrothermal vents along oceanic ridges (from Desbruyères & Ségonzac 1997 modified and completed).

DEEP-SEA HYDROTHERMAL VENTS: THERMOPHILES

Site	MG	LS	Rb	BS	T.A.G.	SP	Lg	LC
Т	265-284	152-333	360-365	356-364	270-363	335-356	>353	40-75
pH	4.2-4.8	3.5-4.9	2.8-3.1		2.5-3.4	3.7-3.9	<3.3	9-9.8
Si	8.2-11.2	9.1-17.5	6.9-8.0		18-22	18-20	7-8.2	9-19
Cl	360-400	410-540	>750	469	633-675	550-563	515-522	546-549
CO ₂	17-20	8.9-28	<16		2.9-4.1	10.63		
H ₂ S	1.5-2	1.4-3.3	1-2.5	9.3	2.5-6.7	2.7-6.1	<1	0.064
CH4	1.35-2.63	0.5-0.97	2.2-2.5	0.065	0.14-0.62	0.046-0.062	2.1	0.13-0.28
Fe	0.002-0.018	0.13-0.86	24	1.68-2.16	1.64-5.45	1.8-2.56	2.50	
Mn	0.068	0.45	2.25	0.26	1	0.49	0.33	
H ₂								0.25-0.43

Table I. – End-member temperature (°C) and concentrations of chemical species (mmol kg⁻¹) for the different M.A.R.vent fields: MG: Menez Gwen; LS: Lucky Strike; Rb: Rainbow; BS: Broken Spur; T.A.G.: Trans Atlantic Geotraverse; SP: Snake Pit; Lg: Logatchev; LC: Lost City (from Desbruyères 2000 completed Kelley 2001).

groups of factors are in turn influenced by water depth and geological contexts.

Deep-sea hydrothermal communities thrive in the interfacial zone where vent fluids mix turbulently with bottom seawater. Microbial communities in these extreme environments are located in at least four generic habitats: (i) free-living populations associated with the discharged vent fluids and depending upon sub-seabed ecosystems, (ii) freeliving microbial mats growing on surface strata that are exposed to mixed fluids, (iii) endo- and exosymbiotic associations of microorganisms and vent fauna, and (iv) microorganisms within the hydrothermal vent plumes (Karl 1995). Microbial communities play at least three major roles according to the taxa involved: (i) autotrophic species produce most of the initial organic matter involved in the ecosystem which in turn is used by heterotrophs, (ii) they play a fundamental role in detoxification, often in symbiotic associations with eukaryotic organisms, and (iii) they are involved in recycling of organic matter produced by the macrofaunal assemblages.

Despite the considerable amount of data gathered during the last decade, monitoring and sampling conditions at deep-sea hydrothermal vents are still a major constraint that render difficult the accurate description of the microhabitats of the various microbial communities. In most cases, it remains almost impossible to gather *in situ* detailed physico-chemical composition of the sampling sites at centimetre scale which would provide appropriate information on the basic requirements of the microorganisms further isolated in the laboratories. This is particularly true on structures where steep pH and temperature gradients are observed like white and black smokers. It is also the case for samples of macrofauna gathered from these environments. Consequently, it is not surprising if isolation and cultivation of microorganisms from deep-sea vents are still the only way to describe their basic physiological characteristics and to infer their role in the ecosystems.

The first report of occurrence of discrete microbial communities using enrichment procedures and fluorescence *in situ* hybridisation (FISH) (Harmsen *et al.* 1997a,b) for a chimney located on the Mid-Atlantic Ridge was recently confirmed for a black smoker of the Manus Basin (New Guinea) (Takai *et al.* 2001). The archaeal phylotypes predominantly located at the top of the smoker were affiliated to *Ignicoccus* and those distributed on the vent surface were mostly *Thermococcales*. These data established that the scale of variation of microhabitats is within the centimetre range, illustrating the difficulty to measure *in situ* the gradients of temperature, pH, oxidation-redox potential and various chemicals.

Microbial diversity assessed by molecular approach

Approaching the diversity of microbial communities associated with deep-sea hydrothermal vents, as well as from other ecosystems, has long been a very tedious task since only two methods were available: direct microscopic observation and laboratory cultivation. Evidence, through 16S rDNA

sequences, of numerous uncultured microorganisms in natural communities has gained strength during the last decade since pioneering work in that field (Ward et al. 1990, Ward et al. 1992). Despite specific bias introduced by molecular methods, demonstrated by the possible lack within the pool of amplified 16S rDNA of sequences corresponding to already isolated microorganisms, it is generally admitted that no more than 1 to 5% of the total microorganisms of a natural community are available through isolation procedures. Consequently, the molecular approach is complementary to the standard enrichment and cultivation procedures in assessing the diversity of microbial communities in extreme environment where our knowledge of appropriate isolation and culture techniques is still limited. This approach, based on extraction of total DNA from crude samples, amplification by PCR of 16S rDNA genes by use of universal primers for Bacteria and Archaea, cloning of these genes for isolation prior sequencing, has allowed, besides determination of microbial communities in situ, the discovery of entirely new phylogenetic lineages of prokaryotes (Moyer et al. 1995, Moyer et al. 1998, Takai & Horikoshii 1999). These novel taxa, based on FISH experiments, appear to be major components of the deep-sea vent ecosystems. Microbial mats are widely distributed at most deep-sea hydrothermal vent sites from Guaymas Basin to East Pacific Rise and Mid-Atlantic Ridge. Recently new phylotypes belonging to a group of uncultured E-Proteobacteria have been identified using an in situ growth chamber at a MAR hydrothermal vent (Reysenbach et al. 2000). The same research group established on a different hydrothermal vent site located on the Southern East Pacific Rise that more than 98% of the sequences of 16S rDNA obtained from a sample of microbial mat were closely related to the previously identified ε -Proteobacteria. They concluded from these data and from previous indications that this new clade of ε -Proteobacteria may be endemic and widely distributed among deep-sea hydrothermal vents (Longnecker & Reysenbach 2001). Two recent results reinforce this hypothesis. Firstly, DGGE profiles and phylogenetic analysis, conducted on enrichment cultures of samples of hydrothermal fluids, chimney-like structures and tubes and specimens of Alvinella spp. collected on a deep-sea vent at 13°N EPR, demonstrated that archaeal sequences were related to the genus Thermococcus and the bacterial sequences to uncultured *ɛ-Proteobacteria*, and Deferribacter thermophilus, Bacillus halodurans, Marinitoga Ralstonia pickettii and camini (Slobodkin et al. 2001). Secondly, we demonstrated (unpublished results) by ARDRA profiles and phylogenetic analysis, conducted on enrichment cultures, using complex proteinaceous substrates, from white tubes of Alvinella spp. collected on 13°N EPR, that most archaeal sequences were related to the Thermococcus genus as previously

mentioned and that the bacterial sequences belonged to *Clostridiales* (20%) and ε -*Proteobacteria* (30%).

The diversity of metabolic types within the ε -*Proteobacteria* already isolated and cultivated from other environments makes it difficult to hypothesise about the metabolic capabilities of the various strains inhabiting deep-sea vents. Considering the high prevalence of the uncultured ε -*Proteobacteria* in both 16S rDNA gene populations of crude samples and of enrichment cultures and their putative role in sulphur cycling in deep-sea vents environments hypothesised by Longnecker *et al.*, inferred from the white sulphur-like coloration of the mat filaments, it is of considerable interest to succeed in culturing them in order to assess their basic metabolic properties.

Cultivated thermophilic microorganisms from deep-sea hydrothermal vents

Research in this field had two main incentives. thermophilic and Firstly, more generally extremophilic microorganisms represent an unprecedented source of enzymes displaying original properties interesting many industrial activities, and known as extremozymes. Hyperthermostable polymerases from Thermococcus DNA and Pyrococcus genera such as Deep-Vent from Pyrococcus sp. GB-D (New England Biolabs, Boston, USA) and Tfu Pol and Pab Pol respectively from Thermococcus fumicolans (Cambon Bonavita et al. 2000) and Pyrococcus abyssi (QbioGene, Illkirch, F) are examples of extremozymes. Despite the fact that extremozymes properties do not generally fit exactly the industry requirements, extremozymes are a matchless material for directed evolution and protein engineering programs. The second major incentive is related to molecular evolution and the origin of life question. Phylogenetic analyses, based on comparisons of 16S rDNA sequences, place the hyperthermophiles as the most slowly evolving of all forms of life, the first to have diverged from the last common universal ancestor. This suggests that life first emerged on this planet in hyperthermophilic conditions. This statement represents a controversial dogma (Wiegel & Adams 1998) that strongly fuels research on hyperthermophiles and deep-sea hydrothermal vents.

Within the Archaea inhabiting deep-sea hydrothermal vents are found the most extreme of known hyperthermophiles with growth temperature optima above 100°C. *Pyrolobus fumarii*, a facultatively aerobic chemolithoautotroph isolated from chimney walls MAR (Blöchl *et al.* 1997) with an optimum growth temperature of 106°C, measurable growth up to 113°C and ability to remain viable after one hour treatment in the autoclave at 121°C, has the highest cardinal temperatures for any known microorganism. Next to *P. fumarii* are found the methanogen *Methanopyrus kandleri*, isolated from chimney walls of a black smoker where it reaches numbers of 10^8 per gram of chimney rock (Kurr *et al.* 1991), the obligate anaerobic heterotroph isolated from the Guaymas Basin *Pyrodictium abyssi* (Pley *et al.* 1991) and the barophilic chemoorganotrophic *Pyrococcus* sp. ES4 isolated from Juan de Fuca Ridge (Pledger & Baross 1991) which are able to grow up to 110 °C.

free living microorganisms, As the Thermococcaceae appear largely distributed on the various hydrothermal fields whether in shallow water or in the deep-sea. Strains belonging to this family are strictly anaerobic hyperthermophilic fermentative sulphur reducing heterotrophs. They use a variety of carbon substrates such as peptides and/or carbohydrates and are probably involved in recycling organic matter produced by autotrophs and higher organisms belonging to deep-sea vent communities. They are the predominant hyperthermophilic group of microorganisms isolated from deep-sea vents. Three genera are represented: Paleococcus with only one species, Pyrococcus (3 from the deep-sea among 5 valid species) and Thermococcus (10 species from the deep-sea among 21). Optimal temperatures for growth are at between 80-88 °C for Paleococcus and Thermococcus and between 90-95 °C for Pyrococcus. Differentiation of the species within the Thermococcus genus requires DNA-DNA hybridisation since they are morphologically and physiologically rather similar to each other. Some of them are barophilic with a direct effect of pressure on growth, like for Paleococcus ferrophilus (1-600 bar range, optimum at 300 bar) (Takai et al. 2000), Pyrococcus abyssi (optimum growth temperature 96 °C at 1 bar, 100 °C at 200 bars) (Erauso et al. 1993). Thermococcus barophilus, isolated from MAR at a depth of 3550 m on the external layer of a chimney wall and from enrichment cultures conducted at 400 bar, requires a minimum of 150-170 bar for growth in the range of 95-100°C (Marteinsson et al. 1999b).

Hyperthermophilic methanogens are also well distributed in deep-sea hydrothermal vent environments. Besides *Methanopyrus kandleri* already mentioned, the genus *Methanococcus* has been documented from MAR (*M. infernus* (Jeanthon *et al.* 1998)), from EPR (*M. vulcanius* (Jeanthon *et al.* 1999)) and from Guaymas Basin, Gulf of California (*M. jannaschii* (Jones *et al.* 1983), *M. fervens* (Jeanthon *et al.* 1999)). Obligate anaerobic, they are chemolithotrophs. They use H_2 and CO_2 as the only substrate for growth and methane production. In the presence of CO_2 and H_2 , they reduce elemental sulphur to hydrogen sulphide.

The other Archaea isolated from deep-sea hydrothermal vents are the sulphate reducer Archaeoglobus profondus (Burggraf et al. 1990), the sulphite reducer Archaeoglobus veneficus (Huber et al. 1997) and members of the kingdom Crenarchaeota including sulphur metabolisers from the genus Desulfurococcus, Staphylothermus (Fiala et al. 1986) and Ignicoccus (Huber et al. 2000).

Many of these genera and species are also found in other environments characterised by their extreme temperatures. Thermococcus litoralis (Neuner et al. 1990) initially isolated from a shallow marine hot spring was also isolated from solfatara and subterranean oil reservoirs. The genus Archaeoglobus is also found in Alaskan and North Sea oil reservoirs (Stetter et al. 1993) as well as from coastal submarine hot springs in Vulcano Island (Stetter 1988). Genera Thermococcus, Archaeoglobus and to a less extent Methanococcus appear to be ubiquitous in marine thermal habitats (Holden et al. 2001).

Comparatively, only a limited number of species belonging to the Bacteria domain have been isolated from deep-sea hydrothermal vents. Among described species of the Thermotogales Thermosipho melanesiensis (Antoine et al. 1997) was isolated from gills of a deep-sea hydrothermal mussel from the Lau Basin in South Western Pacific Ocean, Thermosipho japonicus isolated from the Iheya Basin of Okinawa (Takai & Horikoshi 2000). Marinitoga camini (optimum growth temperature of 55 °C) a chemoorganotrophic sulphur reducing strain (Wery et al. 2001a) and Caloranaerobacter azorensis, an anaerobic thermophilic bacterium belonging to the cluster XII of the Clostridiales (Wery et al. 2001b) were isolated from a chimney sample collected at MAR. A chemolithoautotrophic sulphur reducing Bacteria, Desulfurobacterium thermolithotrophum, with an optimum growth temperature of 70 °C, was isolated from a deep-sea hydrothermal chimney sample collected at the MAR (L'Haridon et al. 1998). More recently our laboratory has succeeded in culturing an *ɛ-Proteo*bacteria, Caminibacter hydrogenophilus (Alain et al. in press), proposed as the representative strain of a new family of *ε-Proteobacteria* well distributed in deep-sea hydrothermal vent environments as reported by several authors (Longnecker & Reysenbach 2001, Reysenbach et al. 2000), on the basis of molecular approaches of microorganisms diversity. This strain was isolated from fragments of chimney rocks and emptied white tubes of Alvinella pompejana samples, collected at EPR 13°N. It represents a novel thermophilic, anaerobic, obligate hydrogen-oxidising bacterium and is able to grow chemolithoautotrophically under an atmosphere of H2/CO2 with So or NO3- as electron acceptor and is probably involved in sulphur cycling in deep-sea hydrothermal vent ecosystems.

Aerobic thermophilic strains are less frequently isolated from deep-sea vent environments, probably due to anaerobic prevalent conditions above 35°C and to unbalanced efforts to uncover new species. However, aerobic Bacteria isolated from vents at 2000 m depth from the Guaymas Basin and MAR at 3500 m and belonging to the genera *Bacillus* and *Thermus* have been reported (Marteinsson *et al.* 1995, Marteinsson *et al.* 1999a).

Molecular adaptations to thermophily

Studies on life at high temperatures are comparatively recent and, despite the results gathered during the last two decades, remain a promising field of interest for topics as diverse as 'origin of life', biomolecules (in)stability, biotechnology and extremozymes, strategies for detection of novel hyperthermophilic organisms in extreme environments on Earth and preparing sampling for life discovery on other planets. All cell components of extremophiles microorganisms have to be heat adapted or at least heat resistant for periods of time compatible with corresponding metabolic constraints. This implies that not only proteins are stable at temperatures above 80, 100 or even 110°C according to the species considered, but that many low molecular weight metabolites and coenzymes, nucleic acids and lipids are either heat resistant or protected against denaturation and/or degradation by specific mechanisms. The upper limit of life at high temperatures evidenced from isolated Archaea might be slightly revised in the future. However, it depends on the stability of biomolecules, availability of energy and thermodynamics, and not the least on the ability of cells to maintain an appropriate proton permeability range of membranes in order to keep electrochemical proton gradients for energy gain (Albers et al. 2000).

Different recent reviews deal with the stability of proteins at high temperatures (Daniel & Cowan 2000, Jaenicke 2000, Ladenstein & Antranikian 1998, Scandurra et al. 2000). A surprising conclusion emerges from these studies: there are no general rules to achieve protein stabilisation. Each extremophilic protein adopts various strategies and the outstanding adaptation to extreme temperature and solvent conditions is realised through the same weak electrostatic and hydrophobic interactions among the ordinary amino acid residues which are also responsible for the proper balance between protein stability and flexibility in mesophilic proteins. Comparative studies between mesophilic enzymes and their hyperthermophilic counterparts indicate that hyperthermostability is gained through a variable combination of a small number of noncovalent features: minimisation of the surface energy and the hydration of apolar surface groups, increased compactness with a decrease of internal cavities and a high number of ionic interactions. The adaptation of proteins to extreme temperatures

appears then to be the result of a compromise between the increased rigidity responsible for thermal stability and the flexibility required for playing their physiological roles. Besides these intrinsic stability factors, protein stability can be enhanced by molecular chaperonins as demonstrated with the thermosome characterised from Methanopyrus kandleri (Andra et al. 1998). This property is probably critical at the upper temperature border of life where heat-shock proteins could play a key role: cultures of P. fumarii with the thermosome fully induced were able to survive 1 h autoclaving at 2 bar and 121°C (Blöchl et al. 1997). Recently, it has been shown that when overexpressed in E. coli the recombinant small heat shock protein (sHSP) from the hyperthermophile Pyrococcus furiosus (involved in adaptation to exposure to temperatures over growth temperature optimum), prevented the majority of E. coli proteins from aggregating in vitro for up to 40 min at 105 °C. Apparently, the sHSP confers a survival advantage on mesophilic Bacteria by preventing protein aggregation at supraoptimal temperatures (Laksanalamai et al. 2001). For more detail, the reader is directed to a recent review on the mechanisms for thermostability of hyperthermophilic enzymes (Vieille & Zeikus 2001).

High temperatures require also a tight control of exchanges at cytoplasmic membrane level. The membrane lipids, as basic components of the thermophilic microbial cell membrane, play a key role in thermophily. The ion permeability of the membrane increases with temperature and is a major factor determining the maximum growth temperature. As demonstrated by Könings, the most important finding is that the proton permeability of most bacterial and all archaeal membranes at their temperature of growth is maintained within a narrow window (H+ permeability coefficient near 10^{-9} cm.s⁻¹). In contrast, the permeability of the membranes for sodium ions at different growth temperatures was not constant, but was found to increase exponentially with temperature in a similar way for all organisms studied (Albers et al. 2000). Changes in lipid composition of membrane is a common strategy of Bacteria and Archaea in response to changes of ambient temperatures. The 'core' lipids of the Archaea are mainly based on saturated isoprenoid chains linked to a glycerol backbone by ether bonds. The increase in cyclization of transmembrane lipids contributes to a better adaptation to high temperatures by reducing membrane fluidity (Gliozzi et al. 1983). Increase in growth temperatures of M. jannaschii induces an increase in the ratio tetraether/diether lipids. The resulting cyclization of the chains tends to decrease the motion of the lipids and therefore contributes to an acceptable membrane fluidity at elevated growth temperature (Sprott et al. 1991).

The maintenance of the genomes and more generally of nucleic acids of hyperthermophiles inhabiting deep-sea hydrothermal vents is also a challenging question since to high temperatures is added a variety of chemical compounds with possible genotoxic effects. Considerable research has been done during the last two decades to analyse the strategies of hyperthermophilic Bacteria and Archaea in DNA and RNA maintenance and summarised in a recent review (Daniel & Cowan 2000). The main problem of DNA stability at high temperatures is thermal degradation due to depurination and subsequent breakage of the phosphodiester bonds (Marguet & Forterre 1994). The DNA of P. furiosus was found to be 20 times more resistant to thermal degradation than DNA of E. coli on the basis of the number of DNA backbone breaks after incubation of the cells at 105 °C (Peak et al. 1995). Less is known about strategies developed to cope with the other harsh conditions found in these environments where the relative instability of fluids composition and of the vents themselves with time imply appropriate molecular adaptations. The case of an other extremophile, Deinococcus radiodurans, isolated from canned meat exposed to X rays is of great interest. The complete sequencing of its genome (White et al. 1999) demonstrated that all systems for DNA repair, DNA damage export, desiccation and starvation recovery, and genetic redundancy are present in one cell. However, a subsequent analysis of this genome reached the conclusion that the fundamental questions underlying the extreme resistance phenotype of D. radiodurans remain unanswered (Makarova et al. 2001). Besides features revealed by genome analysis, resistance is probably the result of a very complex combination of modifications of proteins, nucleic acids and other cell components that are not readily inferred from the sequences. The availability of several complete genomes of hyperthermophiles, including 3 different species of Pyrococcus (Kawarabayasi et al. 1998, Lecompte et al. 2001, Robb et al. 2001), opens the way for more detailed research on specific metabolic pathways which are involved in genome integrity and cell detoxification. Like D. radiodurans, Pyrococcus furiosus has an extraordinarily high capacity for repair of radiation-induced double-strand breaks (DiRuggiero et al. 1997). Interestingly, the same authors demonstrated that at the sequence level, only a few genes share homology with known bacterial repair genes. Phylogenetic analysis indicates that archaeal recombinases occur in two paralogous gene families, one of which is very deeply branched, and both recombinases are more closely related to the eukaryotic Rad51 and Dmc1 gene families than to the E. coli recA gene (DiRuggiero et al. 1999). Comparisons of doublestrand breaks induced by gamma radiation in genomic DNA of *P. furiosus* and *abyssi*, *D.* radiodurans and E. coli have shown that the strong

radioresistance of *Pyrococcus* is not related to a specific protection of DNA. It might partly be linked to the smaller size of the *Pyrococcus* chromosome compared to *E. coli* (1.8 Mb vs 4.6 Mb) and to the homologous recombination processes, taking advantage of the existence of several copies of their chromosomes during the log and stationary phases of growth (Gérard *et al.* 2001).

Many low molecular weight metabolites and coenzymes like ATP and NAD(P) used by archaeal hyperthermophiles have short half-lives at their optimum growth temperatures and are unstable at 110°C (Daniel & Cowan 2000). The way these microorganisms have circumvent this instability is still open to research.

CONCLUSIONS

During the last two decades, more and more new thermophiles and hyperthermophiles from deep-sea hydrothermal vents have been discovered despite the high cost and relative difficulties in sampling. Assessing the global diversity of these extreme environments for microorganisms remains a challenging task as demonstrated by the incongruity of the cultivation and molecular retrieval approaches. A combination of both approaches is obviously needed to better characterise the microbial diversity and obtain novel species in culture. Allocating already isolated species and microorganisms evidenced only by their 16S rDNA sequences to a precise microhabitat on chimney structures has long been difficult for technical reasons. In the near future, the improvement of sampling procedures and of in situ monitoring of physico-chemical parameters is expected to produce data which are deeply needed to understand the ecology of hypermicrobiology thermophiles. Coupling with geochemical and physical dynamics of deep-sea hydrothermal vent structure is also a promising approach. The main interest of the molecular approach was the construction of 16S rDNA libraries from which completely new taxa emerged. From sequences, it is impossible to infer the thermophily of the corresponding microorganisms and much broader temperature ranges have consequently to be tested for enrichment cultures and subsequent isolation procedures. Isolation of novel E-Proteobacteria like Caminibacter hydrogenophilus illustrates this approach.

Considerable progress have been made in recent years in the study of molecular mechanisms that enable the proteins of hyperthermophiles to operate under extreme conditions. The major determinants of protein thermostability have been characterised for many different proteins and revealed the lack of universal law to solve the transition from mesophilic to hyperthermophilic status. Modelling the corresponding processes is consequently almost impossible and directed evolution through DNA shuffling is still the basic method to engineer proteins in order to adjust their properties to industrial requirements. The upper limit of life appears to be less constrained by protein stability, which in many cases could tolerate life up to 150°C, than by limited stability of nucleic acids and low molecular weight compounds. Moreover, most of available data were gathered from in vitro experiments and the shortage of data is acute for in vivo conditions. The complex interactions between the various cell components and the resulting possible stabilising effects make it difficult to evaluate the true stabilities of these molecules in vivo.

The increasing availability of complete genomes, and more specifically genomes from extremophiles isolated from deep-sea hydrothermal vents is an invaluable source of data for the near future. Despite the high percentage of ORFs with undetermined functions, averaging 50% in some cases, the evidence of interkingdom lateral gene transfer (Koonin et al. 1997, Nelson et al. 1999) and the possible resulting chimeric origin of Archaea are strong incentives for genomic research. Genome comparisons between phylogenetically close species like the 3 Pyrococcus have evidenced the high plasticity of these genomes (Lecompte et al. 2001, Myllykallio et al. 2000). It might also be of considerable interest in reconstructing metabolic pathways and help to design experiments for their validation.

Finally, it might be stressed that the way we see the extreme conditions prevailing at deep-sea vent fields is probably inappropriate. Microorganisms have been thriving for millions or even billions of years in these "extreme" conditions and not only survived, but developed strategies for colonising almost every type of habitats including subterranean reservoirs under the sea flour. These harsh conditions are probably the driving force to molecular adaptations which are still to be uncovered.

REFERENCES

- Albers SV, van dVJL, Driessen AJ, Konings WN 2000. Adaptations of the archaeal cell membrane to heat stress. *Front Biosci* 5: D813-20.
- Amend JP, Shock EL 2001. Energetics of overall metabolic reactions of thermophilic and hyperthermophilic Archaea and Bacteria. *Fems Microbiol Rev* 25(2): 175-243.

- Andra S, Frey G, Jaenicke R, Stetter KO 1998. The thermosome from *Methanopyrus kandleri* possesses an NH4[±]dependent ATPase activity. *Eur J Biochem* 255(1): 93-9.
- Antoine E, Cilia V, Meunier JR, Guézennec J, Lesongeur F, Barbier G 1997 Thermosipho melanesiensis sp. nov., a new thermophilic anaerobic bacterium belonging to the order Thermotogales, isolated from deep-sea hydrothermal vents in the Southern Pacific Ocean. Int J Syst Bacteriol 47(4): 1118-1123.
- Blöchl E, Rachel R, Burggraf S, Hafenbradl D, Jannasch HW, Stetter KO 1997. *Pyrolobus fumarii*, gen. and sp. nov., represent a novel group of archaea, extending the upper temperature limit for life to 113°C. *Extremophiles* 1: 14-21.
- Brock TD 1969. *Thermus aquaticus* gen. n. and sp. n., a non sporulating extreme thermophile. *J Bacteriol* 98: 289-297.
- Burggraf S, Jannasch HW, Nicolaus B, Stetter KO 1990. Archaeoglobus profundus sp. nov., represents a new species within the sulfate-reducing Archaebacteria. System Appl Microbiol 13: 24-28.
- Cambon Bonavita MA, Schmitt P, Zieger M, Flaman JM, Lesongeur F, Raguenes G, Bindel D, Frisch N, Lakkis Z, Dupret D et al. 2000. Cloning, expression, and characterization of DNA polymerase I from the hyperthermophilic archaea Thermococcus fumicolans. Extremophiles 4(4): 215-225.
- Daniel RM, Cowan DA 2000. Biomolecular stability and life at high temperatures. *Cell Mol Life Sci* 57(2): 250-64.
- Delaney JR, Kelley DS, Lilley MD, Butterfield DA, Baross JA, Wilcock W S, Embley RW, Summit M. 1998. The quantum event of oceanic crustal accretion: impacts of diking at mid-ocean ridges. *Science* 281: 222-30.
- Desbruyères D, Alayse-Danet A-M, Ohta S 1994. Deapsea hydrothermal communities in Southwestern Pacific back-arc basins (the North Fiji and Lau Basins): composition, microdistribution and food-web. *Marine Geol* 116: 227-242.
- Desbruyères D, Almeida A, Biscoito M, Comtet T, Khripounoff A, Le Bris N, Sarradin PM, Segonzac M 2000. A review of the distribution of hydrothermal vent communities along the northern Mid-Atlantic Ridge: dispersal vs. environmental controls. *Hydrobiologia* 440(1-3): 201-216.
- Desbruyères D, Segonzac M 1997. Handbook of deepsea hydrothermal vent fauna. Brest: IFREMER, 279 p.
- DiRuggiero J, Brown JR, Bogert AP, Robb FT 1999. DNA repair systems in archaea: mementos from the last universal common ancestor? J Mol Evol 49(4): 474-84.
- DiRuggiero J, Santangelo N, Nackerdien Z, Ravel J, Robb FT 1997. Repair of extensive ionizing-radiation DNA damage at 95 degrees C in the hyperthermophilic archaeon *Pyrococcus furiosus*. J Bacteriol 179(14): 4643-5.
- Erauso G, Reysenbach AL, Godfroy A, Meunier JR, Crump B, Partensky F, Baross JA, Marteinsson V, Barbier G, Pace NR *et al.* 1993. *Pyrococcus abyssi* sp. nov., a new hyperthermophilic archaeon isolated from a deep-sea hydrothermal vent. *Arch Microbiol* 160: 338-349.

- Fiala G, Stetter KO, Jannasch HW, Langworthy TA, Madon J 1986. *Staphylothermus marinus* sp. nov. represents a novel genus of extremely thermophilic submarine heterotrophic archaebacteria growing up to 98. *Syst Appl Microbiol* 8: 106-113.
- Fornari DJ, Embley RW 1995. Tectonic and volcanic controls on hydrothermal processes at the Mid-Ocean ridge: an overview based on near-bottom and submersible studies. *In* SE Humphris RAZ, LS Mullineaux & RE Thomson, editors. Seafloor Hydrothermal Systems: American Geophysical Union: 1-46.
- Gérard E, Jolivet E, Prieur D, Forterre P 2001. DNA protection mechanisms are not involved in the radioresistance of the hyperthermophilic archaea *Pyrococcus abyssi* and *P. furiosus*. *Mol Gen Gen* 261(1): 72-78.
- Gliozzi A, Rolandi R, De RM, Gambacorta A 1983. Monolayer black membranes from bipolar lipids of archaebacteria and their temperature-induced structural changes. J Membr Biol 75(1): 45-56.
- Harmsen HJM, Prieur D, Jeanthon C 1997a. Distribution of microorganisms in deep-sea hydrothermal vent chimneys investigated by whole-cell hybridization and enrichment culture of thermophilic subpopulations. Appl Environ Microbiol 63(7): 2876-2883.
- Harmsen HJM, Prieur D, Jeanthon C 1997b. Group-specific 16S rRNA-targeted oligonucleotide probes to identify thermophilic bacteria in marine hydrothermal vents. *Appl Environ Microbiol* 63(10): 4061-4068.
- Hessler RR, France SC, Boudrias MA 1987. Hydrothermal vent communities of the Mariana back-arc basin. *Eos* 68(44): 1531.
- Hessler RR, Smithey WM Jr 1983. The distribution and community structure of megafauna at the Galapagos Rift hydrothermal vents. *In* Rona PA, Boström K, Laubier L, Smith KL Jr editors. Hydrothermal processes at seafloor spreading centers. New York: Plenum Press: 735-770.
- Holden JF, Takai K, Summit M, Bolton S, Zyskowski J, Baross JA 2001. Diversity among three novel groups of hyperthermophilic deep-sea *Thermococcus* species from three sites in the northeastern Pacific Ocean. *FEMS Microbiol Ecol* 36(1): 51-60.
- Huber H, Burggraf S, Mayer T, Wyschkony I, Rachel R, Stetter KO 2000. *Ignicoccus* gen. nov., a novel genus of hyperthermophilic, chemolithoautotrophic Archaea, represented by two new species, *Ignicoccus islandicus* sp. nov. and *Ignicoccus pacificus* sp. nov. Int J Syst Evol Microbiol 50: 2093-1000.
- Huber H, Jannasch H, Rachel R, Fuchs T, Stetter KO 1997. Archaeoglobus veneficus sp. nov., a novel facultative chemolithoautotrohic hyperthermophilic sulfite reducer, isolated from abyssal black smokers. Syst Appl Microbiol 20: 374-380.
- Jaenicke R 2000. Stability and stabilization of globular proteins in solution. J Biotechnol 79(3): 193-203.
- Jeanthon C, L'Haridon S, Reysenbach AL, Corre E, Vernet M, Messner P, Sleytr UB, Prieur D 1999. Methanococcus vulcanius sp. nov., a novel hyperthermophilic methanogen isolated from East Pacific Rise, and identification of Methanococcus sp. DSM 4213T as Methanococcus fervens sp. nov. Int J Syst Bacteriol 49 Pt 2: 583-9.
- Jeanthon C, L'Haridon S, Reysenbach AL, Vernet M, Messner P, Sleytr UB, Prieur D 1998. *Methanococcus infernus* sp. nov., a novel hyperthermophilic lithotro-

phic methanogen isolated from a deep-sea hydrothermal vent. Int J Syst Bacteriol 48 Pt 3: 913-9.

- Jones WJ, Leigh JA, Mayer F, Woese CR, Wolfe RS 1983. Methanococcus jannashii sp. nov., an extremely thermophilic methanogen from a submarine hydrothermal vent. Arch Microbiol 136(1100a): 254-261.
- Karl DM, editor. 1995. The microbiology of deep-sea hydrothermal vents. 124 p.
- Kawarabayasi Y, Sawada M, Horikawa H, Haikawa Y, Hino Y, Yamamoto S, Sekine M, Baba S, Kosugi H, Hosoyama A *et al.* 1998. Complete sequence and gene organization of the genome of a hyper- thermophilic archaebacterium, *Pyrococcus horikoshii* OT3. *DNA Res* 5(2): 55-76.
- Kelley DS, Karson JA, Blackman DK, Früh-Green GL, Butterfield DA, Lilley MD, Olson EJ, Schrenk MO, Roe KK, Lebon GT *et al.* 2001. An off-axis hydrothermal vent field near the Mid-Atlantic Ridge at 30°N. *Nature* 412: 145-149.
- Koonin EV, Mushegian AR, Galperin MY, Walker DR 1997. Comparison of archaeal and bacterial genomes: computer analysis of protein sequences predicts novel functions and suggests a chimeric origin for archaea. *Mol Microbiol* 25(4): 619-637.
- Kurr M, Huber R, König H, Jannasch HW, Fricke H, Trincone A, Kristjansson JK, Stetter KO 1991. *Methanopyrus kandleri*, gen. and sp. nov. represents a novel group of hyperthermophilic methanogens, growing at 110°C. *Arch Microbiol* 156: 239-247.
- Ladenstein R, Antranikian G 1998. Proteins from hyperthermophiles: stability and enzymatic catalysis close to the boiling point of water. *Adv Biochem Eng Biotechnol* 61: 37-85.
- Laksanalamai P, Maeder DL, Robb FT 2001. Regulation and Mechanism of Action of the Small Heat Shock Protein from the Hyperthermophilic Archaeon *Pyrococcus furiosus*. J Bacteriol 183(17): 5198-202.
- Lecompte O, Ripp R, Puzos-Barbe V, Duprat S, Heilig R, Dietrich J, Thierry JC, Poch O 2001. Genome evolution at the genus level: comparison of three complete genomes of hyperthermophilic archaea. *Genome Res* 11(6): 981-93.
- L'Haridon S, Cilia V, Messner P, Raguenes G, Gambacorta A, Sleytr UB, Prieur D, Jeanthon C 1998. *Desulfurobacterium thermolithotrophum* gen. nov., sp. nov., a novel autotrophic, sulphur-reducing bacterium isolated from a deep-sea hydrothermal vent. *Int J Syst Bacteriol* 48: 701-11.
- Longnecker K, Reysenbach A 2001. Expansion of the geographic distribution of a novel lineage of epsilon-Proteobacteria to a hydrothermal vent site on the Southern East Pacific Rise. *FEMS Microbiol Ecol* 35(3): 287-293.
- Lutz R, Kennish M 1993. Ecology of Deep-sea Hydrothermal Vent Communities: A Review. *Rev Geophysics* 31(3)(4053): 211-242.
- Makarova KS, Aravind L, Wolf YI, Tatusov RL, Minton KW, Koonin EV, Daly MJ. 2001 Genome of the extremely radiation-resistant bacterium *Deinococcus radiodurans* viewed from the perspective of comparative genomics. *Microbiol Mol Biol Rev* 65(1): 44-79.
- Marguet E, Forterre P 1994. DNA stability at temperatures typical for hyperthermophiles. *Nucleic Acids Res* 22(9): 1681-1686.

- Marteinsson VT, Birrien JL, Kristjansson JK, Prieur D 1995. First isolation of thermophilic aerobic non-sporulating heterotrophic bacteria from deep-sea hydrothermal vents. *FEMS Microbiol Ecol* 18: 163-174.
- Marteinsson VT, Birrien JL, Raguenes G, da CMS, Prieur D 1999a. Isolation and characterization of *Thermus thermophilus* Gy1211 from a deep-sea hydrothermal vent. *Extremophiles* 3(4): 247-51.
- Marteinsson VT, Birrien JL, Reysenbach AL, Vernet M, Marie D, Gambacorta A, Messner P, Sleytr UB, Prieur D 1999b. *Thermococcus barophilus* sp. nov., a new barophilic and hyperthermophilic archaeon isolated under high hydrostatic pressure from a deepsea hydrothermal vent. *Int J Syst Bacteriol* 49 Pt 2: 351-9.
- Moyer CL, Dobbs FC, Karl DM 1995. Phylogenetic diversity of the bacterial community from a microbial mat at an active, hydrothermal vent system, Loihi Seamount, Hawaii. *Appl Environ Microbiol* 61(4): 1555-62.
- Moyer C, Tiedje JM, Dobbs FC, Karl DM 1998. Diversity of deep-sea hydrothermal vent *Archaea* from Loihi Seamount, Hawaii. *Deep-Sea Research* II 45: 303-17
- Myllykallio H, Lopez P, Lopez Garcia P, Heilig R, Saurin W, Zivanovic Y, Philippe H, Forterre P 2000. Bacterial mode of replication with eukaryotic-like machinery in a hyperthermophilic archaeon. *Science* 288: 2212-2215.
- Nelson KE, Clayton RA, Gill SR, Gwinn ML, Dodson RJ, Haft DH, Hickey EK, Peterson JD, Nelson WC, Ketchum KA *et al.* 1999. Evidence for lateral gene transfer between Archaea and bacteria from genome sequence of *Thermotoga maritima*. *Nature* 399: 323-9.
- Neuner A, Jannasch HW, Belkin S, Stetter KO 1990. *Thermococcus litoralis* sp. nov.: a new species of extremely thermophilic marine archaebacteria. Arch Microbiol 153: 205-7.
- Peak MJ, Robb FT, Peak JG 1995. Extreme resistance to thermally induced DNA backbone breaks in the hyperthermophilic archaeon *Pyrococcus furiosus*. J *Bacteriol* 177(21): 6316-6318.
- Pledger RJ, Baross JA 1991. Preliminary description and nutritional characterization of a chemoorganotrophic archaeobacterium growing at temperatures of up to 110°C isolated from a submarine hydrothermal vent environment. J Gen Microbiol 137(1): 203-211.
- Pley U, Schipka J, Gambacorta A, Jannasch HW, Fricke H, Reinhard R, Stetter KO 1991. Pyrodictium abyssi, sp. nov., represents a novel heterotrophic marine archaeal hyperthermophilic growing at 110°C. Syst Appl Microbiol 14: 245-253.
- Reysenbach AL, Longnecker K, Kirshtein J 2000. Novel bacterial and archaeal lineages from an *in situ* growth chamber deployed at a Mid-Atlantic Ridge hydrothermal vent. *Appl Environ Microbiol* 66(9): 3798-806.
- Robb FT, Maeder DL, Brown JR, DiRuggiero J, Stump MD, Yeh RK, Weiss RB, Dunn DM 2001. Genomic sequence of hyperthermophile, *Pyrococcus furiosus*: implications for physiology and enzymology. *Methods Enzymol* 330: 134-57.
- Scandurra R, Consalvi V, Chiaraluce R, Politi L, Engel PC 2000. Protein stability in extremophilic archaea. *Front Biosci* 5: D787-95.

- Slobodkin A, Campbell B, Cary SC, Bonch-Osmolovskaya E, Jeanthon C 2001. Evidence for the presence of thermophilic Fe(III)-reducing microorganisms in deep-sea hydrothermal vents at 13 degrees N (East Pacific Rise). FEMS Microbiol Ecol 36(2-3): 235-243.
- Sprott GD, Meloche M, Richards JC 1991. Proportions of diether, macrocyclic diether, and tetraether lipids in *Methanococcus jannaschii* grown at different temperatures. J Bacteriol 173(12): 3907-10.
- Stetter KO 1988. Archaeoglobus fulgidus gen. nov., sp. nov.: a new taxon of extremely thermophilic archaebacteria. Syst Appl Microbiol 10: 172-173.
- Stetter KO 1998. Hyperthermophiles: isolation, classification, and properties. *In* Horikoshi K, Grant WD, editors. Extremophiles – Microbial life in Extreme Environments. New York: Wiley: 1-24.
- Stetter KO 1999a. Extremophiles and their adaptation to hot environments. *FEBS Lett* 452(1-2): 22-5.
- Stetter KO 1999b. Volcanoes, hydrothermal venting, and the origin of life. *In* Marti J, Ernst GJ, editors. Volcanoes and the Environment. Cambridge: Cambridge University Press.
- Stetter KO, Huber R, Blöchl E, Kurr M, Eden RD, Fielder M, Cash H, Vance I 1993. Hyperthermophilic archaea are thriving in deep North Sea and Alaskan oil reservoirs. *Nature* 365: 743-745.
- Summit M, Baross JA 2001. A novel microbial habitat in the mid-ocean ridge subseafloor. *Proc Natl Acad Sci.* USA 98(5): 2158-63.
- Takai K, Horikoshi K 1999. Genetic diversity of archaea in deep-sea hydrothermal vent environments. *Genetics* 152: 1285-97.
- Takai K, Horikoshi K 2000. Thermosipho japonicus sp. nov., an extremely thermophilic bacterium isolated from a deep-sea hydrothermal vent in Japan. Extremophiles 4(1): 9-17.
- Takai K, Komatsu T, Inagaki F, Horikoshi K 2001. Distribution of archaea in a black smoker chimney structure. Appl Environ Microbiol 67(8): 3618-29.
- Takai K, Sugai A, Itoh T, Horikoshi K 2000. Palaeococcus ferrophilus gen. nov., sp. nov., a barophilic, hyperthermophilic archaeon from a deep-sea hydrothermal vent chimney. Int J Syst Evol Microbiol 50: 499-500.
- Tunnicliffe V 1988. Biogeography and evolution of hydrothermal-vent fauna in the eastern Pacific Ocean. *Proc Royal Soc London* B 233: 347-366.
- Van Dover CL 1995. Ecology of Mid-Atlantic Ridge hydrothermal vents. *In* Parson LM, Walker CL, Dixon DR editors. Hydrothermal vents and processes. London: Geol Soc Spec Publi: 257-294.
- Vieille C, Zeikus GJ 2001. Hyperthermophilic enzymes: Sources, uses, and molecular mechanisms for thermostability. *Microbiol Mol Biol Reviews* 65(1): 1-43.
- Von Damm KL 1995. Controls on the chemistry and temporal variability of seafloor hydrothermal fluids. *In* SE Humphris RAZ, LS Mullineaux & RE Thomson editors. Seafloor Hydrothermal Systems: American Geophysical Union: 222-247.
- Von Damm KL, Bray AM, Buttermore LG, Oosting SE 1998. The geochemical controls on vent fluids from the Lucky Strike vent field, Mid-Atlantic Ridge. *Earth Planet Sci Let* 160: 521-536.

- Ward DM, Bateson MM, Weller R, Ruff-Roberts AL 1992. Ribosomal RNA analysis in microorganisms as they occur in nature. *Adv Microbiol Ecol* 12: 219-286.
- Ward DM, Weller R, Bateson MM 1990. 16S rRNA sequences reveal numerous uncultured microorganisms in a natural community. *Nature* 345: 63-65.
- Wery N, Lesongeur F, Pignet P, Derennes V, Cambon Bonavita MA, Godfroy A, Barbier G 2001a. Marinitoga camini gen, nov., sp nov., a rod-shaped bacterium belonging to the order Thermotogales, isolated from a deep-sea hydrothermal vent. Int J Syst Evol Microbiol 2: 495-504.
- Wery N, Moricet JM, Cueff V, Jean J, Pignet P, Lesongeur F, Cambon-Bonavita MA, Barbier G 2001b. Caloranaerobacter azorensis gen. nov., sp. nov., an anaerobic thermophilic bacterium isolated from a deep-sea hydrothermal vent. Int J Syst Evol Microbiol 51: 1789-1796.

a.es and de Environment Cambridge: Cambridge Einwerenty Frids Stener KO Huber 8, Biford E, Kure M Eiden RD Fiel der M. Cash R. Van. at 1993 Hyperiterrauphilie ar chara are thriving a deep NonteSea and Alaskan oil reservoirs. Naura 365 743-745

- Soment M. Bluess IA. 2001. A upped interochar non-netral the mid-ocean tridge subscalloor. Proc Natl Acod Sci. USA 98(5): 2158-63.
- Fekai K, Horikoshi K 1999, Genetic diversity of architer in deep sea hydrothermal yent environments. Cent. net 152, 1:285-07.
- Tatai K. Havkoshi K 2000 Thermmann phone (5.4) aox. at extremely ihermolohillo breterium isolinod from a deep-sea is doulismed went in Japan. Cifrementation at 1: 9-17.
- Teled R. Komatso T. Inagalo P. Boritoshi K 2001: Distribution of archaes in a black smoker chitanoy shireture. Anal Eavier Microbiol 67(8): 3018-29.
- Tatsei K. Sugar A. Itoli T. Honikoshi K. 2000. Palatororone terrophilus gen nov., sp. nov. a batopäiht, hyperthetinophilis architeon from a dean-sui hydrothermal van chimaty. Int J Spat Fred Micropiel 30: 499-500.
- Termichtle V 1938 Biogeography and evolution of avdiothermal-year found in the castern Pacific Ocean, here Royal Soc London IN 233, 347-366
- Van Dover (1, 1905) Isostopy of Mid. Allantic Ridge his derificienti vents. In Preson I M, Wilker (1, Discus DR editors, Electronicemal vents and processor. London: Gool Soc Free Public 137-294.
- Viellie C. Zeitas GJ 2001. Hyperhemophile enzymes: Sources, uses, and moleculit mechanisms for Incjmosability. Microfild Met Boll Review 67(1): 1-43.
- Von Dama KL 1994 Controls on the districting and temporal variability of scalloor hydrothermal fluids in SE Humphris RAZ, LS Multimeaux & Ro Thornson editors: Scalloor Hydrothermal Systems: Amoreau Grouphysical Union: 222-247
- Yon Dagun KL, Bay AM, Buttemore LG, Oosting ME 1998. The geochemical contraits on year Hidds from the Lucky Nirtke year field. Mid-Affantic Ridges Earth Physics Soc Let 160: 521-536.

- White O, Eisen JA, Heidelberg JF, Hickey EK, Peterson JD, Dodson RJ, Haft DH, Gwinn ML, Nelson WC, Richardson DL *et al* 1999. Genome sequence of the radioresistant bacterium *Deinococcus radiodurans* R1. *Science* 286: 1571-1577.
- Wiegel J, Adams MWW editors. 1998. Thermophiles: The Keys to Molecular Evolution and the Origin of Life? Athens, USA: Taylor & Francis,346 p.
- Woese CR, Fox GE 1977. Phylogenetic structure of the prokaryotique domain: the primary kingdoms. *Proc Natl Acad Sci USA* 74: 5088-90.
- Zierenberg RA, Adams MWW, Arp AJ 2000. Life in extreme environments: Hydrothermal vents. *Proc Natl Acad Sci USA*. 97(24): 12961-12962.

Reçu le 30 août 2001; received August 30, 2001 Accepté le 5 novembre 2001; accepted November 5, 2001

- Litem K.C. Clayton RA. Gill SP. Owino MI., Dudson RI, H.H. EH, Hickev RK, Parerson JD. Netson W.C. JSCI: bina KA et al. 1999 Evidence for Interal gene in faster between Archaea and bacteria from genome sequence of Thermotogia matriana. Materie 3991 37:19
- benace A. Jaansch HW, Bolkin S, Stoner KU 1990. Thermocuccus huoratis sp hov : a now species of estremely thermophile marine archaebaciefia Area Me contine 101. 203-1.
- Peak MJ, Robb FT, Peak 4G 1995. Extreme existance to thermally induced DNA backbone bunchs in the by performability exchange formercean firetures. J Received 177-011: 6316 6315.
- Pledger RJ, Baross JA 1991, Preliminary description and agritudiat characterization of a chemoorganotrophic erclipediateration growing at compositioned of up to 110°C isolated from a submarge bydeoflorenal west environment. J Gen Microbiol 137(1): 103-211
- Play U. Schieka J. Gambacorta A. Jamasch HW. Fricke H. Bernhardt R. Stetter KO 1991. Production objects so nov., represents a novel futerofrephic marine arciaccal hyperhemophilic growing at 110°C. South April March 2001 [4: 235-253.
- Reventration AL, Longneder K, Kurduen J 2020, Novel parternation functional lineage from in 19 year prostin opariter-leadoved at a Mid-Atlantic folder hearther and vents Ame Frances Microbial 60(9), 295-806, Morb FT, Macder DL, Brown JK, Diffinggioro 1, Stung
- ath. Yeb & Yours Rb. Dunit DM 2001. Conomic sequence of hypertitermentalic. Proceeding Intromus on invication: or physiology and envymology. Merthous antivinol. 330: 114-57.
- Seamona H. Con attri V. Chiamidde R. Politi L. Eneri PC 2001. Protein Scientify in extremophilic archaea. Prost. Epise 17:277-05.