

Coping with cold: The genome of the versatile marine Antarctica bacterium *Pseudoalteromonas haloplanktis* TAC125

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A considerable fraction of life develops in the sea at temperatures lower than 15°C. Little is known about the adaptive features selected under those conditions. We present the analysis of the genome sequence of the fast growing Antarctica bacterium *Pseudoalteromonas haloplanktis* TAC125. We find that it copes with the increased solubility of oxygen at low temperature by multiplying dioxygen scavenging while deleting whole pathways producing reactive oxygen species. Dioxygen-consuming lipid desaturases achieve both protection against oxygen and synthesis of lipids making the membrane fluid. A remarkable strategy for avoidance of reactive oxygen species generation is developed by *P. haloplanktis*, with elimination of the ubiquitous molybdopterin-dependent metabolism. The *P. haloplanktis* proteome reveals a concerted amino acid usage bias specific to psychrophiles, consistently appearing apt to accommodate asparagine, a residue prone to make proteins age. Adding to its originality, *P. haloplanktis* further differs from its marine counterparts with recruitment of a plasmid origin of replication for its second chromosome.

[Supplemental material is available online at www.genome.org. The sequence data from this study have been submitted to EMBL under accession nos. CR954246 and CR954247. The data are also available at the following Web sites: www.genoscope.cns.fr/agc/mage/psychroscope and <http://bioinfo.hku.hk/PsychroList/>

Three quarters of the Earth is covered by sea and >90% of its surface experiences yearly temperatures <15°C, asking for a remarkable adaptation of life to cold conditions. Several marine bacteria have been studied, but so far we possess only limited information about life in the sea at medium and low temperatures (Bartlett 1999; Raven et al. 2002; Thomas and Dieckmann 2002). Furthermore, heterotrophic bacteria in sea ice play a key role in carbon cycling, while little is known about their metabolic features, which are beginning to be deciphered (Moran et al. 2004). Challenges posed by cold to life stem from the slow pace of chemical reactions (for reviews, see Lonhienne et al. 2000; Feller and Gerday 2003; Weber and Marahiel 2003; Georgette et al. 2004), from the constraints induced by the stability of hydrogen bonds (the situation is particularly challenging for folded nucleic acids, i.e., RNA), and from increased solubility of gases

and stability of radicals. Phylogenetic studies based on 16S rRNA sequences indicated close relationships between marine bacteria within two bacterial divisions, Proteobacteria (in the genera *Alteromonas*, *Colwellia*, *Glaciicola*, *Octadecabacter*, *Pseudoalteromonas*, *Shewanella*, and *Vibrio*) and *Cytophaga-Flexibacter-Bacteroides* (*Cytophaga*, *Flavobacterium*, *Gelidibacter*, and *Polaribacter*) (Ivanova et al. 2004). *Pseudoalteromonas haloplanktis* TAC125 has been isolated from an Antarctic coastal sea water sample collected in the vicinity of the French Antarctic station Dumont d'Urville, Terre Adélie (66°40' S; 140° 01' E). By using genome sequencing, corroborated by *in silico* and *in vivo* analyses, we have uncovered exceptional genomic and metabolic features of this γ -proteobacterium compared with other bacteria from aqueous environments (Supplemental Table 1). These features, some of which explored by physiological experiments, account for their remarkable versatility and fast growth, showing adaptation to rare but periodic situations of abundance, making it an organism of choice for exploring heterologous protein production at low temperature (Tutino et al. 2001; Duilio et al.

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Table 1. General features of the *Pseudoalteromonas haloplanktis* genome

	Chromosome 1	Chromosome 2
Size (bp)	3,214,944	635,328
G+C percentage	41.0	39.3
Number of predicted CDSs	2942	546
Average size of CDSs (bp)	950	1013
Percentage coding	88.6	87.3
Number of rRNA operons (16S-23S-5S)	9	0
5S rRNA (extra copies)	1	0
Number of tRNAs	106	0
CDSs similar to known proteins	1123	157
Putative functions (limited homology/structural features)	759	251
Conserved hypothetical proteins	694	75
Orphan proteins	325	61
Doubtful CDS and gene remnant	41	2

2004). Although cold conditions are so prevalent on Earth, we do not possess at the moment a reference set of annotations for the genome of bacteria thriving in such conditions. The present genome sequence was therefore carefully annotated manually, and annotation will be, as much as possible, continuously refined. We endeavored to post to the community reference databases, allowing investigators to rapidly and efficiently retrieve relevant information while comparing it to what is known in other genomes. In the MaGe platform (www.genoscope.cns.fr/agc/mage/psychroscope) investigators can not only see gene annotations in parallel with synteny with the genomes they chose as relevant but compare the annotations with cognate data from the UniProt knowledgebase, as well as explore possible EC numbers, metabolic pathways reconstruction, clusters of orthologous genes, or membrane prediction properties. As a complement, a specialized database, PsychroList (<http://bioinfo.hku.hk/PsychroList/>), within the GenoChore suite (Fang et al. 2005), allows the user to search for patterns in DNA or protein sequences, taking into account a clustering of genes into formal operons as well as providing extra facilities to query sequences using predefined sequence patterns.

Results and Discussion

Genome organization

As in many marine γ -proteobacteria, the *P. haloplanktis* TAC125 genome is made of two chromosomes (Table 1; Supplemental Table 2). The replication origin of chromosome (chr) I maps near *dnaA* (McLean et al. 1998; Lobry and Louarn 2003) in a region that is highly conserved in γ -proteobacteria (Fig. 1). However, in remarkable contrast with the genomes of the vibrios (Okada et al. 2005), the second chromosome does not display a standard GC skew (Supplemental Fig. 1). The pattern observed is likely to be caused by unidirectional replication. To our knowledge, this is the first time that such a system would be uncovered in an authentic bacterial chromosome. This is supported by the signature of R1 plasmid replication (del Solar et al. 1998): the *tus* and *repA*-like genes, the *repA* and *dnaA* boxes, and the *parA* and *parB* genes (Fig. 1). In addition, *kisB* and *kidB* coding for a typical plasmid maintenance system have also been found in chrII. We chose the start at the centre of a TATATA palindrome near the genes coding for the partition system. The G+C content and gene density of chrII match that of chrI (Supplemental Table 2). It contains the

essential genes *hisS* and *gcpE*, in addition to a series of genes ubiquitous in γ -proteobacteria (Supplemental Table 3). A third of chrII genes have orthologs in *Escherichia coli*. Remarkably, the whole metabolism of histidine is coded in chrII, in a highly conserved gene cluster (Fig. 1). Nineteen percent of the *P. haloplanktis* chrII genes show high similarities with plasmid-encoded genes, further suggesting that this replicon was a plasmid recruited to become a chromosome encoding essential genes (Fig. 1; Supplemental Table 2).

Genes around the origin of replication in chromosome display a high level of synteny with genes of other known proteobacteria. chrI codes for nine rDNA clusters (23S, 5S, and 16S RNAs, one operon has two copies of the 5S RNA gene), a large number compared with that found in most sequenced γ -proteobacteria (Ussery et al. 2004). The genome sequence of *P. haloplanktis* TAC125 shows some variability in the rRNA genes of ~1% (interestingly, most variations correspond to compensating mutations in regions coding for double stranded RNA) (data not shown). However, this probably does not influence phylogenies (Cilia et al. 1996), and the rRNA sequences are in line with the established phylogeny, placing TAC125 near vibrios and *Shewanella* (Ivanova et al. 2004). In the same way, the number of tRNA genes is quite high (106 genes), a feature in common with that in vibrios and in *Photobacterium profundum* (Table 1). These genes are organized in long runs of repeated sequences. The longest contains 19 tRNA genes in a row, seven of which coding for an identical tRNA^{Glu} (TTC anticodon), suggesting a slipped mispair mechanism of expansion in situations of rapid growth. Genomes with an origin of replication display a protein-coding gene distribution bias, with most essential genes located in the leading replication strand (Rocha and Danchin 2003). Interestingly, while the number of CDSs located in the leading strand of the chromosome is 61%, 72% of the tRNA genes are located in that strand. Because the speed of transcription/translation must be limited at low temperature, the large number of rRNA and tRNA genes may participate in the adaptation, allowing fast growth of the organism in the cold. TAC125 is similar to other bacteria in terms of number of tandem repeats. Several genes relevant to adaptation to cold conditions are clustered together in approximate repeats (Supplemental Fig. 2): genes coding for cold-shock proteins, nine paralogs of *cspA*, as in *E. coli* (four in chrII, three of them clustered together), as well as genes coding for a class of putative short secreted proteins that could bind calcium, next to a divalent metal exporter system, most likely used in calcium export. Calcium is known to be involved in cold adaptation and formation of exopolysaccharides (EPS) in bacteria (Kierek and Watnick 2003; Dominguez 2004). The duplication in chrII of the *ast* operon may be involved in adaptation to cold and high osmolarity.

Comparative genomics

Of the 3488 identified protein coding genes (CDSs) (Table 1), a biological function, based on a classification scheme adapted from Riley (Riley 1993; Fang et al. 2005), has been assigned for >65.6% (36.7% with a final assignment and 28.9% with a putative role assignment). More than 63% of the *P. haloplanktis* CDSs are similar to *Shewanella oneidensis* genes (Heidelberg et al. 2002), of which 47% are found in synteny groups, making the comparison with this aquatic organism particularly revealing (Fig. 2A). These mesophilic marine bacteria, together with *Vibrio vulnificus*, share with *P. haloplanktis* several sodium-type flagellar proteins

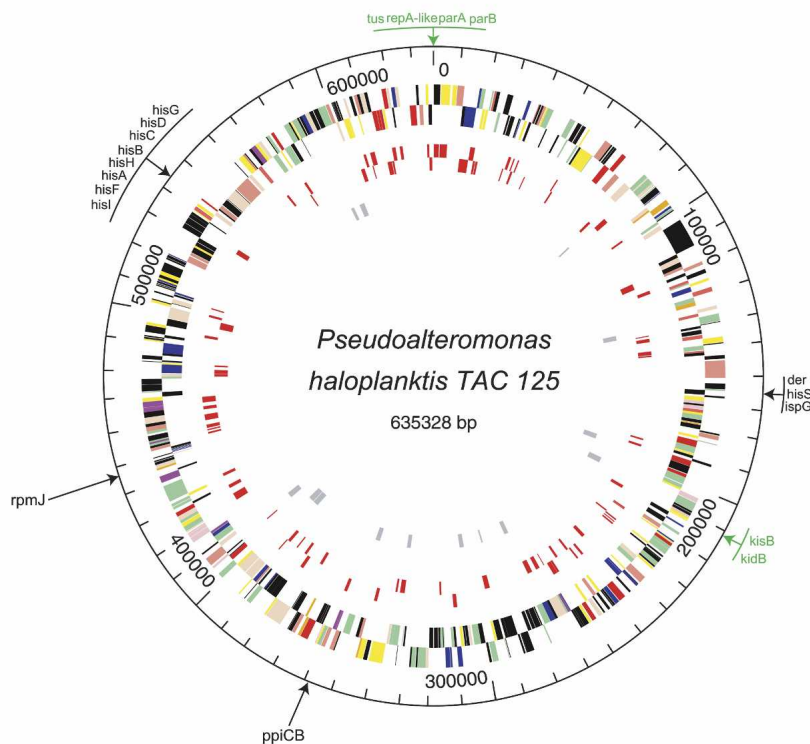
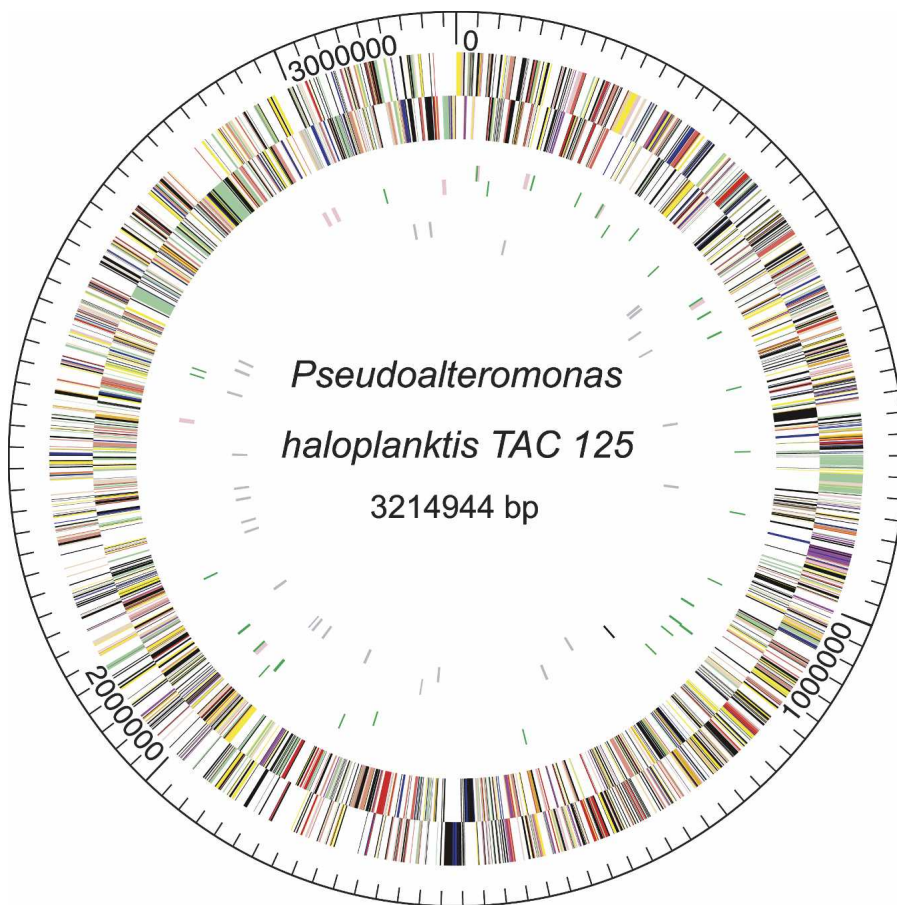


Figure 1. Circular representation of the *Pseudoalteromonas haloplanktis* genome. Circles display (from the outside): (1) predicted coding regions transcribed in the clockwise direction; (2) predicted coding regions transcribed in the counterclockwise direction. Genes displayed in 1 and 2 are color-coded according to different functional categories: salmon indicates amino acid biosynthesis; orange indicates purines, pyrimidines, nucleosides, and nucleotides; purple indicates fatty acid and phospholipid metabolism; light blue indicates biosynthesis of cofactors, prosthetic groups, and carriers; light green indicates cell envelope; red indicates cellular processes; brown indicates central intermediary metabolism; yellow indicates DNA metabolism; green indicates energy metabolism; pink indicates protein fate/synthesis; blue indicates regulatory functions; grey indicates transcription; teal indicates transport and binding proteins; and black indicates hypothetical and conserved hypothetical proteins. (3) tRNAs (green) and rRNA (pink) on chrI/genes similar to phage proteins (red) on chrII; (4) and *tonB* and *tonB*-like genes in grey. Chromosome II gene names similar to that of the R1 plasmid replication apparatus (unidirectional) are colored in green.

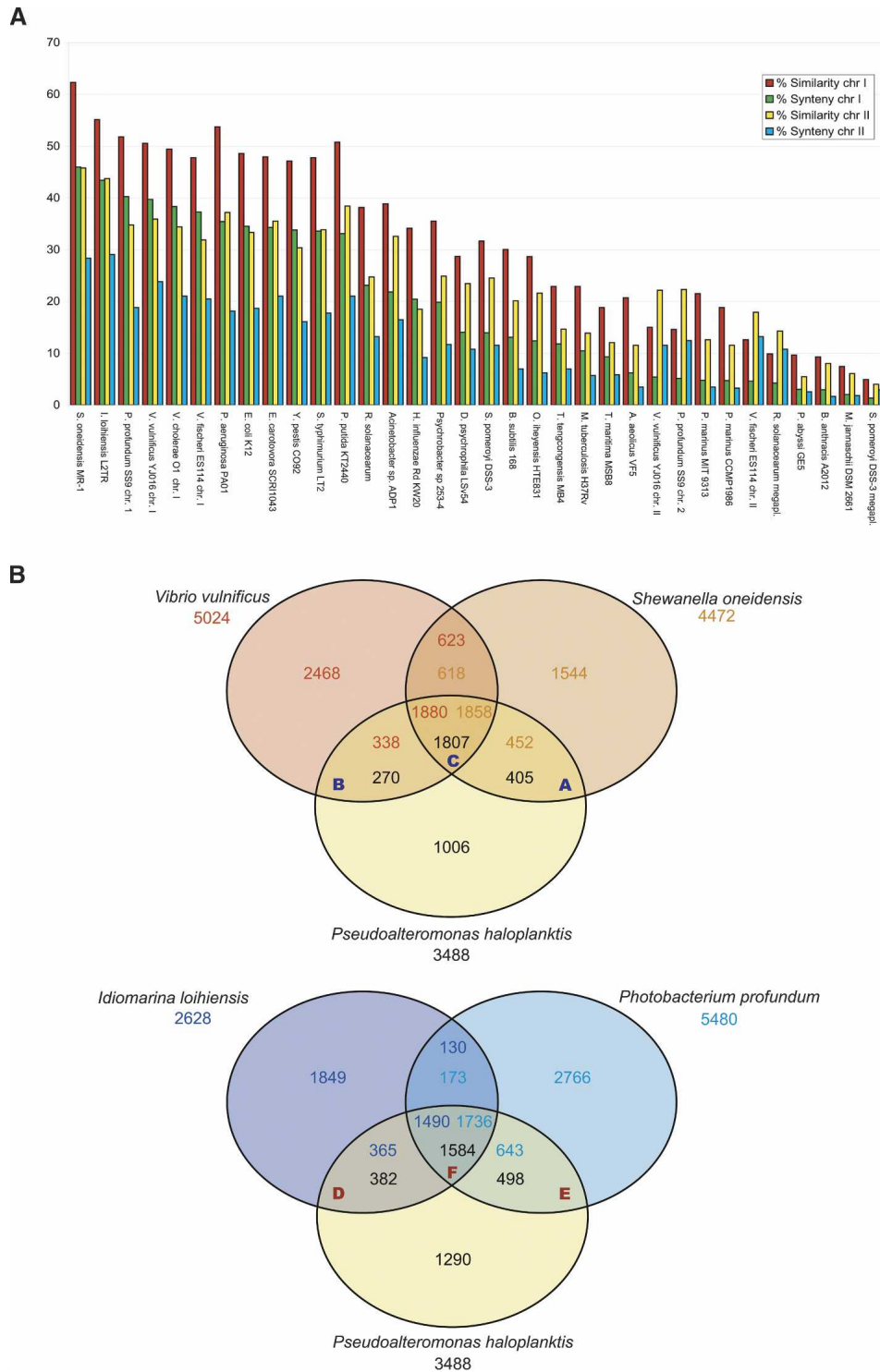


Figure 2. Putative orthologs and synteny between the genome of *P. haloplanktis* and the genome of related bacteria. The alphabetic letters A to F refer to the Supplemental Table 4 (T4, A to F). (A) The percentage of *P. haloplanktis* genes homologous to a selection of 34 complete bacterial proteomes (i.e., 30% identity and a ratio of 0.8 of the length of the smaller protein to that of the larger one) is represented by red bars for chrI and by yellow bars for chrII. The percentage of *P. haloplanktis* genes in synteny groups with a selection of 34 complete bacterial genomes is represented by green bars for chrI and by blue bars for chrII. The closest organism is *S. oneidensis*. (B) Comparison of the gene content of *Pseudoalteromonas haloplanktis*, *Shewanella oneidensis*, *Vibrio vulnificus*, *Photobacterium profundum*, and *Idiomarina loihiensis*. Putative orthologs are defined as genes showing a minimum of 30% identity and a ratio of 0.8 of the length of the smaller protein to that of the larger one, or as two genes included in a synteny group. The intersections between the three circles give the number of genes found in the two or three compared species. Genes outside these areas are specific to the corresponding organism. The total number of annotated genes is also given under each species name.

(*mot* genes), sodium-dependent transporters, gene clusters of type IV pilus and flagellin proteins, and some iron ABC transporter and *tonB* system-dependent transport proteins (Fig. 2B; Supplemental Table 4C). A putative cold-shock RNA methyltransferase (PSHAb0516) and a cold-shock regulated carbon starvation protein A (PSHAb0210) have also been found in *S. oneidensis* and *V. vulnificus*. Other *P. haloplanktis* TonB-dependent receptors and calmodulin-like proteins have counterparts in *S. oneidensis* only, while *V. vulnificus* shares with *P. haloplanktis* several integrases/transposases, putative potassium channel proteins and metabolite exporters (Supplemental Table 4A,B). Gene content comparisons have also been performed with the two closest psychrophilic γ -proteobacteria, *Idiomarina loihiensis* and *Photobacterium profundum* (Fig. 2B): apart from the set of genes shared also with the marine bacteria, many TonB-dependent receptors are specific to *P. haloplanktis* and *I. loihiensis*, and two sodium-dependent carbohydrate transporters (a permease and a transporter) have been found in *P. profundum* only (Supplemental Table 4D,E). The set of genes common to *P. haloplanktis* and these two psychrophilic bacteria contain two copper resistance protein genes (*copA* and *copB* genes in chrII) (Supplemental Table 4F). These genes belong to a synteny group made of six genes with *I. loihiensis* and, interestingly, with the megaplasmid of *Ralstonia solanacearum*. They are also found in other distant cold marine bacteria such as *Psychrobacter* sp. and *Silicibacter pomeroyi* (in its megaplasmid; Supplemental Table 5). Two synteny groups (a potassium efflux system and an urease operon) are found with *S. pomeroyi* only. A total of 133 *P. haloplanktis* genes (73 hypothetical), including several insertion sequences and genes coding for TonB-dependent receptors, have not been found in the genome of the cold-adapted marine bacteria used for comparison (Supplemental Table 6). In addition, a prophage region (50 kb long, between genes PSHAA1505 and PSHAA1558) is specific to *P. haloplanktis*. We also identified one specific region in chrI coding for several calcium-dependent proteins, as well as a specific gene in chrII that may regulate cell volume and resistance to cold conditions (PSHAb0555).

General features of the proteome

Global properties of the proteins at the level of individual amino acids and motifs integrate all kinds of selection pressure associated to adaptation to cold. The pattern of amino acid distribution in γ -proteobacteria from different biotopes (Supplemental Fig. 3) displays an overall trend similar in the various genomes of interest, with leucine (L) being most abundant, while tryptophan (W), cysteine (C), histidine (H), and to a lesser extent methionine (M) are used infrequently. The proteome of the thermophilic genomes looks significantly different from that of the mesophilic and psychrophilic counterparts (strong avoidance of glutamine [Q] in thermophilic species, preference for alanine [A] in mesophilic and psychrophilic species, except in *Oceanobacillus iheyensis*). The amino acid distribution in mesophilic and psychrophilic species display a few noteworthy differences specifically relevant to growth in the cold (in particular in the relative abundance of N and Q) (Supplemental Fig. 3). Using correspondence analysis (CA) (Benzécri 1984) coupled to dynamic clustering (Delorme and Henaut 1988), to identify subtle differences in this cluster of related objects, five classes with close amino acid composition were found (Fig. 3): (1) integral inner membrane proteins (~12%); (2) proteins involved in the metabolism of small molecules (25%); (3) associated to information transfer pathways (21.5%);

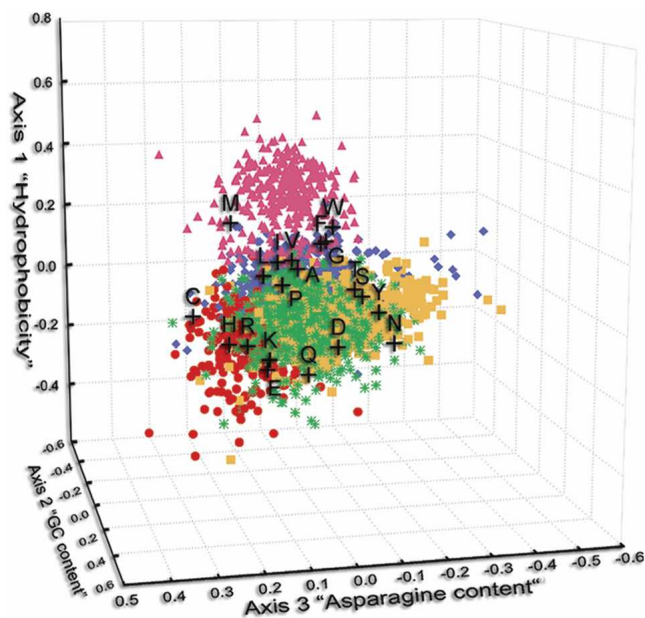


Figure 3. Distribution of the protein sequences on the CA space determined by the three first factors. The first one discriminates proteins by their hydrophobicity; the second one, by the GC content of genes coding for them; and the third one, by their asparagine content. Five classes of proteins are found by clustering method (see "General Features of Proteome") and are represented: (1) IIMP by pink triangles, (2) proteins involved in the metabolism of small molecules by blue diamonds, (3) proteins associated to information transfer pathways by red circles, (4) proteins associated to the outer membrane or secreted by green stars, and (5) proteins with unknown functions, or likely to be of phage origin, by yellow squares. Amino acids are represented by black pluses.

(4) associated to the outer membrane or secreted (21.5%); and (5) with unknown functions, or likely to be of phage origin (20%). This biological consistency demonstrates that a relationship exists between amino acid composition and the role of the protein inside the cell. The two first biases have been previously identified: They are driven by the membrane compartmentalization of some proteins and by the G+C-content of codons (Pascal et al. 2005). In contrast, the bias scattering proteins along the third factorial axis was unexpected. It discriminated proteins in *P. haloplanktis* TAC125 according to their asparagine (N) content. This small hydrophilic uncharged amino acid carries an amide group that is often thermolabile (Zhou et al. 2000; Stratton et al. 2001; Weintraub and Manson 2004). Analysis of this remarkable N-driven bias was further submitted to CA using the pool of two psychrophilic (*Desulfotalea psychrophila* and *P. haloplanktis*), two mesophilic (*E. coli* K-12 and *Bacillus subtilis*) and two thermophilic (*Aquifex aeolicus* and *Thermotoga maritima*) bacteria. An N-driven bias was observed in proteins from the psychrophiles, differentiating them from their mesophilic and thermophilic counterparts (Supplemental Fig. 4). The proteins responsible for the bias belong to the following structures or processes: motility of the organism, cell wall, outer membrane, transport, sensor activity, adaptation to atypical conditions, and secretion. Some proteins of DNA metabolism (replication, packaging, segregation, restriction/modification, proof-reading, and repair) and RNA metabolism are also present. Asparagine residues often undergo deamidating cyclisation, a process extremely sensitive to temperature (Daniel et al. 1996). The *P. haloplanktis* TAC125 proteome is enriched in N residues compared with counterparts that

prefer growth at higher temperature, making it an organism of choice for foreign protein production when deamidation ought to be put to a minimum (Weintraub and Manson 2004).

RNA folding and metabolism

The genome of *P. haloplanktis* TAC125 contains 19 genes presumably coding for known RNA binding proteins or RNA chaperones. The most unexpected feature of the genome is the prominent absence of a RNA/nucleoid-associated cold-shock gene ubiquitous in γ -proteobacteria, *hms*. Analysis in silico failed to uncover a H-NS-like protein gene even when using the highly relaxed comparison criteria that uncovered counterparts in bacteria phylogenetically distant from *E. coli* (Tendeng and Bertin 2003). In contrast, the genes coding for all other nucleoid-associated proteins such as HU, IHF, FIS, and Hfq are present in the genome. The existence of a complete set of genes (including *hms*) was also observed in other marine and cold-adapted bacteria, including *S. oneidensis* (Heidelberg et al. 2002) and *D. psychrophila* (Rabus et al. 2004). In vivo complementation of a *hms* defect of *E. coli* at room temperature led us to repeatedly isolate clones coding for the counterpart of *csrA* as an efficient complementation of the *hms* defect of *E. coli*. The lack of the *hms*-encoding gene in the psychrophilic *P. haloplanktis* genome shows that H-NS is not sufficient to promote growth at low temperature and that its role is connected to that of the regulatory protein/RNA complex CsrA/CsrB (Fig. 4). The importance of control of RNA folding and degradation at low temperature is visible in the presence of many RNA helicases (three copies of *rhIE*, two in *chrI* and one in *chrII*, and possibly a fourth one, PSHAa0641, and two copies of *srnB*, instead of one in *E. coli*). Interestingly, in contrast with the situation reported for *Oleispira antarctica* (Ferrer et al. 2004) the *groES* *groEL* genes from TAC125 did not permit growth of *E. coli* at low temperature. Other factors are therefore important for cold adaptation in the present organism.

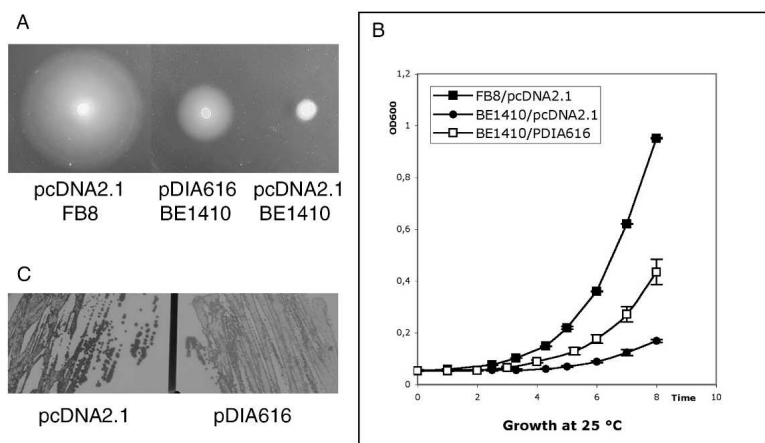


Figure 4. *hms* complementation in *E. coli* with *P. haloplanktis* *csrA*. A DNA fragment encompassing *P. haloplanktis* gene *csrA* with 140 bp of its upstream region was cloned into plasmid pcDNA2.1 (pDIA616) and the phenotypes of the resulting *E. coli* *hms* transformants were compared to the *hms* mutant (BE1410) and to the wild-type FB8 parental strain. (A) Motility assay: Partial motility is restored with the *csrA* gene of *P. haloplanktis*. Other phenotypes such as serine sensitivity of the *hms* mutant are restored by the *csrA* gene as well. (B) Growth at 25°C: overnight cultures were diluted to 0.05 OD₆₀₀ in LB medium with ampicillin 100 μ g/mL, and growth was monitored (as in Dersch et al. 1994). Significant improvement of growth is witnessed with the *csrA* gene of *P. haloplanktis*. (C) CsrA-dependent storage of glycogen in *E. coli* MG1655. Expression of *P. haloplanktis* *csrA* inhibits glycogen storage (light iodine color, right panel).

Several RNA motifs indicating the presence of RNA-coding genes have been found in *chrI* using the Rfam databank (see Methods): a tmRNA at position 2,231,297 bp, the RNA component of RNase P at position 2,674,641 bp, the t44 RNA (of unknown function), and three riboswitch structures (Nudler and Mironov 2004), a lysine riboswitch just in front of *dapA* gene (at position 189,092 bp); the RFN element, located at 1283,710 near *ribB* gene; and the “ubiquitous” TPP riboswitch (THI element) at 496,922 bp (upstream of the *thiC* gene).

Response to oxygen and reactive oxygen species

The solubility of gasses increases rapidly at low temperature. This is the case of dioxygen, which is a very reactive molecule. We expected that the proteome would comprise a vast arsenal of enzymes active against H₂O₂ and superoxide. This was the more so because the organism is indeed remarkably resistant to H₂O₂ (Supplemental Fig. 5). Surprisingly, we found only the gene counterpart coding for the iron superoxide dismutase (*sodB*) and only one clear catalase (*katB*) located in *chrII*, with a possible paralog in *chrI* (PSHAa1737). Furthermore, while the oxygen responding OxyR control is present, the SoxR regulation is absent.

This unexpected finding was explained when we discovered that *P. haloplanktis* TAC125 lacks a series of activities that result in reactive oxygen species (ROS) production. Despite the availability of molybdate in sea water (Hille 2002), *P. haloplanktis* TAC125 entirely lacks molybdopterin metabolism: not only are the biosynthetic and transport gene absent but genes coding for enzymes using the cofactor are also missing (e.g., TMAO reductase, xanthine oxidase, biotin sulfoxide reductase, or the novel oxido-reductase YedY) (Loschi et al. 2004). All related organisms such as the vibrios or *Shewanella* have molybdopterin metabolism, as do almost all Bacteria and Eukarya. The cells, however, must cope with increased oxygen solubility and inevitable interaction with reduced iron, leading to the deleterious Fenton reaction and

ROS. A way for the cells to protect their metabolism against those is to use dioxygen directly. This is seen in the large number of putative dioxygenases present in both chromosomes (Table 2). To this list one should add the fatty acids desaturases (see below) that also play a role to increase membrane fluidity at low temperature.

Furthermore, the protective role of methionine against ROS is enhanced by the existence of two MsrA proteins, instead of one as is usual, together with a fused MsrA-MsrB protein, a rare situation, found in *Nitrosomonas europaea*. Interestingly, one MsrA copy, PSHAa2274, is present in all related genomes, while the second copy, PSHAa1583, not present in *P. profundum*, exists in *Shewanella* and vibrios, and two copies in *D. psychrophila* that develop preferentially at low temperature. A further sign that the bacteria cope with the specific problems posed by ROS is the number of proteins involved in scavenging chemical groups affected: peroxiredoxins such as alkyl hydroperoxide reductase AhpC, AhpCB,

Table 2. Proteins involved in cold/salt adaptation

Locus	Proteins size	Gene name	Chr.	Description
Cold-shock protein				
PSHAa0109	463	<i>dbpA</i>	1	Putative ATP-dependent RNA helicase; putative cold-shock protein
PSHAa0114	421	<i>rhlB</i>	1	ATP-dependent RNA helicase, putative cold-shock protein
PSHAa0990	617	<i>deaD</i>	1	ATP-dependent RNA helicase, cold-shock protein A
PSHAa1184	68	<i>cspC</i>	1	Cold-shock protein, transcription antiterminator
PSHAa1222	209	<i>grpE</i>	1	Putative member of the DnaK/DnaJ/GrpE foldase complex (heat/cold-shock protein; involved in thermal regulation of folding)
PSHAa1600	204		1	Putative cold-shock DNA-binding domain
PSHAa1726	73	<i>cspD</i>	1	Nucleic acid-binding domain, cold-shock RNA chaperone
PSHAa2978/2979/2980	70	<i>cspE</i>	1	RNA chaperone, transcription antiterminator
PSHAa2981	750		1	Cold-shock RNase R
PSHAb0078	63	<i>cspX</i>	2	Cold-shock protein
PSHAb0210	491	<i>cstA</i>	2	Cold-shock regulated carbon starvation protein A
PSHAb0384/0386/0387	70	<i>cspX</i>	2	Cold-shock protein
PSHAb0516	171		2	Putative cold-shock RNA methyltransferase
Dioxygenases				
PSHAa0187	235		1	Conserved protein of unknown function; putative dioxygenase
PSHAa0570	140		1	Conserved protein of unknown function; putative glyoxalase domain
PSHAa0900	297	<i>tesB</i>	1	Putative dioxygenase
PSHAa0904	372		1	Putative dioxygenase
PSHAa2137	365		1	Putative protein with ferredoxin subunits; putative dioxygenase
PSHAa2147	309		1	Putative taurine dioxygenase
PSHAa2168	351	<i>melA</i>	1	4-hydroxyphenylpyruvate dioxygenase (4HPPD) (HPD) (HPPDase)
PSHAa2449	158		1	Conserved protein of unknown function; putative dioxygenase domain
PSHAb0029	167		2	Putative enzyme; dioxygenase superfamily
PSHAb0041	119		2	Putative enzyme; dioxygenase superfamily
PSHAb0115	128		2	Putative enzyme; dioxygenase superfamily
PSHAb0338	434	<i>hmgA</i>	2	Homogentisate 1,2-dioxygenase
Fatty acids (desaturase)				
PSHAa0567	271		1	Conserved protein of unknown function; putative sterol desaturase family protein
PSHAa1269	351		1	Putative fatty acid desaturase
PSHAa2897	378		1	Putative long chain acyl-CoA desaturase
PSHAb0225	264		2	Putative C-5 sterol desaturase
Salt adaptation				
PSHAa0020	459	<i>trkA</i>	1	Potassium transport inner membrane protein subunit; involved in osmoprotection
PSHAa0208	257	<i>cysQ</i>	1	3',5' adenosine diphosphate 3' phosphatase, sodium sensing
PSHAa0325	188	<i>osmY</i>	1	Putative hyperosmotically inducible periplasmic protein
PSHAa0326	53		1	Putative low-temperature and salt-responsive protein
PSHAa0396	273	<i>mscS</i>	1	Putative mechanosensitive channel protein; protection against hypo-osmotic shock
PSHAa0687	255	<i>surE</i>	1	Putative acid phosphatase SurE; protection against osmotic shock
PSHAa0833	512	<i>putP</i>	1	Major sodium/proline symporter; protection against osmotic shock
PSHAa1072	522		1	Putative glycine betaine transporter
PSHAa1226	439		1	Putative Na ⁺ /H ⁺ antiporter
PSHAa1436	531		1	Putative choline/betaine transporter
PSHAa1625	403		1	Putative sodium ABC exporter, permease component
PSHAa1626	263		1	Putative sodium ABC exporter, ATP-binding component
PSHAa1678	675	<i>prc</i>	1	Periplasmic C-terminal protease with specificity for nonpolar C termini; protection against osmotic shock
PSHAa1679	213	<i>proQ</i>	1	Putative post-translational activator of ProP expression; sensing osmotic shock
PSHAa2041	676		1	Putative choline/betaine transporter
PSHAa2202	144	<i>osmC</i>	1	Osmotically inducible protein C; protection against oxidative stress
PSHAa2243	102	<i>bolA</i>	1	Regulator involved in adaptation to osmotic shock
PSHAa2252	561		1	Putative voltage-gated ClC-type chloride channel ClcA
PSHAa2274	660	<i>mdoB</i>	1	Putative phosphoglycerol transferase; shape adaptation to osmotic shock
PSHAa2849	434	<i>envZ</i>	1	Sensor histidine kinase, senses osmolarity
PSHAa2850	240	<i>ompR</i>	1	Response regulator for adaptation to osmolarity
PSHAb0106	576	<i>cvrA</i>	2	Putative Na ⁺ /H ⁺ antiporter; adaptation to hypo-osmotic shock
PSHAa0176	289		2	Putative mechanosensitive channel protein; protection against hypo-osmotic shock
PSHAb0127	1536	<i>glbB</i>	2	Glutamate synthase, large subunit, GOGAT
PSHAb0128	497	<i>glbD</i>	2	Glutamate synthase, small subunit
PSHAb0261	534	<i>betC</i>	2	Putative choline dehydrogenase
PSHAb0365	639	<i>dnaK</i>	2	Chaperone protein DnaK; involved in protection
Against osmotic shock				
PSHAb0381	129	<i>betC</i>	2	Putative osmC-like protein
PSHAb0418	556	<i>betA</i>	2	Putative choline dehydrogenase
PSHAb0419	500	<i>betB</i>	2	NAD ⁺ -dependent betaine aldehyde dehydrogenase
PSHAb0420	197	<i>betI</i>	2	Putative transcriptional repressor for the cellular response to osmotic stress

(continued)

Table 2. Continued

Locus	Proteins size	Gene name	Chr.	Description
Against osmotic shock				
PSHAb0426	489		2	Putative succinylglutamic semialdehyde dehydrogenase; may be involved in adaptation to high osmolarity
PSHAb0427	345		2	Putative arginine succinyltransferase; may be involved in adaptation to high osmolarity
PSHAb0428	402		2	Putative acetylornithine transaminase; may be involved in adaptation to high osmolarity
Helicases [DEA(DH)]				
PSHAa0109	463	<i>dbpA</i>	1	Putative ATP-dependent RNA helicase; putative cold-shock protein; could be specific for rRNA folding
PSHAa0114	421	<i>rhIB</i>	1	ATP-dependent RNA helicase with nucleoside triP hydrolase domain, putative cold-shock protein
PSHAa0510	409	<i>srmB</i>	1	ATP-dependent RNA helicase
PSHAa0641	433		1	ATP-dependent RNA helicase (rhIE-like); DEAD-box protein family
PSHAa0990	617	<i>deaD</i>	1	ATP-dependent RNA helicase, cold-shock protein A
PSHAa1144	1298	<i>hrpA</i>	1	Helicase, ATP-dependent
PSHAa1432	441	<i>srmB</i>	1	ATP-dependent RNA helicase, DEAD box family
PSHAa1522	415		1	Conserved protein of unknown function
PSHAa1930	467	<i>rhIE</i>	1	Putative ATP-dependent RNA helicase with P-loop hydrolase domain
PSHAa1991	692	<i>dinG</i>	1	Putative ATP-dependent helicase DinG
PSHAa2216	809	<i>hrpB</i>	1	ATP-dependent helicase
PSHAa2480	831	<i>rnr</i>	1	RNase R, 3'-5' exoribonuclease
PSHAa2572	608	<i>recQ</i>	1	ATP-dependent DNA helicase
PSHAa2624	965	<i>hepA</i>	1	RNA polymerase associated protein (ATP-dependent helicase HepA)
PSHAa2762	676	<i>rep</i>	1	Rep helicase, a single-stranded DNA-dependent ATPase
PSHAa2983	435	<i>rhIE</i>	1	DEAD-box protein family; putative ATP-dependent RNA helicase with P-loop hydrolase domain
PSHAb0003	1049		2	Putative DNA helicase with DEAD/DEAH box helicase domain
PSHAb0039	1352		2	Putative protein with DEAD/DEAH box helicase domain
PSHAb0119	649		2	Putative ATP-dependent DNA helicase (recQ-like)
PSHAb0411	434	<i>rhIE</i>	2	Putative ATP-dependent RNA helicase with P-loop hydrolase domain; DEAD-box protein family
PSHAb0497	639	<i>yoaA</i>	2	Putative ATP-dependent helicase YoaA with nucleotide triphosphate hydrolase domain, SOS repair, DinG family

thiol-specific cytoplasmic peroxidase BcpA; thiol-specific periplasmic peroxidase Tpx, and their coupled flavoproteins: AhpF, TrxB, PSHaA0892, Gor. All these activities would protect against the inevitable consequences of the Fenton reaction, and the needed sulfur metabolism genes, involving glutathione and S-adenosylmethionine, are all present. Furthermore, a control system that would limit the concentration of copper in the cell (Cu^{2+} is particularly reactive toward oxygen) involves at least two transport systems to expel copper out of the cell (CusA, CopAB, in chrI and PSHAb0008, PSHAb0009, CopB, CopA, and PSHAb0325 in chrII), a periplasmic disulfide reductase (DipZ), and a periplasmic putative laccase (PcoA) that could chelate copper ions while also acting as a dioxygenase.

Finally, an interesting gene cluster involved in fatty acid metabolism PSHAa0894–0910 is absent from other γ -proteobacteria, but is largely present in *Mycobacterium tuberculosis* (where it may have a protective role against ROS, contributing to virulence). In short, *P. haloplanktis* TAC125 is remarkably well adapted to protection against ROS under cold conditions, a feature that could be very useful for expression of foreign proteins in the cold.

Metabolic features

Marine bacteria are facing a medium generally unbalanced in terms of carbon, nitrogen, and phosphorous supply (Moran et al. 2004) but are not depleted in sulfur sources. Strain TAC125 is adapted to fast growth, suggesting that it regularly encounters a fairly rich medium (this is probably due to its propensity to make a water/air biofilm [see below], allowing it to live in region full

of plankton debris). Excess of several easily metabolized carbon sources present simultaneously is unlikely, making catabolite repression the exception rather than the rule. Indeed, *P. haloplanktis* TAC125 is lacking the cAMP-CAP complex that regulates carbon availability in related organisms such as vibrios and *Shewanella*. Furthermore, in contrast to many γ -proteobacteria (including vibrios), it does not possess a phosphoenolpyruvate-dependent phosphotransferase system for the transport and first metabolic step of carbohydrate degradation. This accounts for its lack of growth on glucose, likely to be phosphorylated by glucokinase (PSHAa1364). When oxygen is present at a high level, the presence of glucose-phosphate isomerase might drive the Embden Meyerhof pathway and subsequently activate NADPH-dependent aldose reductase (PSHAa2392). This would affect the polyol pathway, lowering the amount of NADPH available for the reduction of oxidized glutathione by glutathione reductase. Indeed, growth is inexistent on glucose unless tyrosine is supplemented to the medium (data not shown). The counterpart of PSHAa2392 has been implicated in a process controlling tyrosine bradytrophism in *E. coli* (Timms and Bridges 1998).

An essential step for biomass construction is formation of pyruvate, which, because of the absence of the PTS, must go through an alternative pathway starting with pyruvate kinase. In contrast to *E. coli*, with two such enzymes, and vibrios, with three, TAC125 possesses only one pyruvate kinase. Interestingly, it is homologous to the cold-adapted PykA enzyme of *E. coli*. This is further in line with gluconate as a preferred carbon source, providing pyruvate directly through the Entner-Doudoroff pathway (Edd, Eda), which also provides the level of NADPH needed

for protection against oxygen toxicity. As in vibrios, phosphoenolpyruvate synthase (*ppsA* gene) and phosphoenolpyruvate carboxylase (*ppc* gene) genes are located in chrII. However, *ppsA* is located in chrI in *P. profundum*.

TAC125 grows in minimal medium under anaerobic conditions (data not shown), in line with the presence of the *fnr* gene, while the putative aspartate ammonia lyase PSHAA0048 would provide the needed fumarate electron acceptor.

In contrast, the metabolism of nitrogen appears to be highly similar to that in phylogenetically related organisms and be controlled by a phosphorylation cascade involving PtsP (a homolog of the PTS enzyme PtsI) that phosphorylates PtsO and the regulator PtsN, controlling all sigma54-dependent operons. Arginine catabolism could provide a direct source of ammonia under nitrogen-limiting conditions, through the AST pathway, while providing metabolites for adaptation to cold (Schneider et al. 1998). The organism can metabolize *N*-acetyl-glucosamine, a carbon and nitrogen source ubiquitously present in marine environment (Riemann and Azam 2002). In the same way, phosphate input in metabolism is controlled by the counterparts of PhoB, PhoR, and PhoU, with several putative transport systems, including one of high affinity.

Most coenzymes can be synthesized in the organism except coenzyme B12 and, as discussed, molybdopterin. There does not appear to exist a selenium metabolism, in line with the high reactivity of that atom toward oxygen. Other marine bacteria (including the closest one, *S. oneidensis*) do have a selenium metabolism.

Growth, yield, and adaptation to salt

A remarkable feature of TAC125 is that, when provided with sufficient nutrients and aeration, it grows to very high density under laboratory settings, even at 0°C. The very high growth yield indicates that respiration must be particularly efficient in this bacterium. The cells are well adapted to salt, and although they can grow in low osmolarity media, optimal growth is between 1.5% and 3.5% NaCl. We looked for systems that would account for controlling osmolarity in the cell. The common trehalose system does not seem to be present. In contrast, chrII harbors two copies of a choline dehydrogenase (PSHAb0261 and PSHAb0418) for synthesis of glycine betaine, an extremely efficient osmoprotectant (Felitsky et al. 2004). Along the same line, GOGAT glutamate synthase is coded in that same chromosome, allowing synthesis of glutamate that, as potassium glutamate, is the most common response of bacteria to increased osmolarity (Table 2; Lee and Gralla 2004).

All autonomous organisms have at least one pathway to degrade *S*-adenosylhomocysteine (AdoHcy). The MtnN(Pfs)/LuxS pathway leads to synthesis of the quorum sensing (QS) effector autoinducer-2 (AI-2). Xanthomonadales and Pseudomonadales aside, all γ -proteobacteria sequenced use that pathway. LuxS, responsible for the last enzymatic step of AI-2 synthesis is present in bacteria closest to TAC125, and AI-2 produced by heterologous organisms triggers luminescence in reporter strains. The *mtnN* gene is present in TAC125. We failed, however, to identify the *luxS* gene. We looked further for other genes involved in AdoHcy degradation and recycling to homoserine lactones or autoinducer CAI-1. Among other systems, involving quinolones, cyclic dipeptides, or indole, none appears to be present, and in assays, using *Photobacterium luminescens* as reporter TAC125 supernatant did not trigger luminescence. This does not, however, exclude the presence of less well known QS systems: gene PSHAA0159

codes for a multidomain putative aconitate hydratase that may use aconitate as an iron-sulfur cluster-dependent signal in stationary phase (Kiley and Beinert 2003). Furthermore, TAC125 possesses several enoyl-CoA hydratase-like genes that may be involved in synthesis of a diffusible signaling factor as in plant pathogenic γ -proteobacteria (Barber et al. 1997).

Membranes, motility, biofilms, and secretion

P. haloplanktis TAC125 deals with the membrane fluidity challenge at low temperature by lipid desaturases, which simultaneously protect against dioxygen (Table 2). Two clusters of genes, absent in the closest genomes, may be involved in the degradation of steroids or hopanoids, membrane rigidifying molecules present in the environment of heterotrophic bacteria. Protein export from the cytoplasm is similar to that of proteobacteria, with the long form of SecE. Type II secretion is functional (GSP proteins are present) as is the TAT export system. In contrast, type III secretion is absent. TAC125 possesses two gene clusters in chrI and one in chrII for the biosynthesis of type IV pili and of curli, respectively. Furthermore, chrII encodes elements of the new pathway involved in secretion of a specific amylase composed of a signal peptide, the mature enzyme, and a long C-terminal propeptide without foldase function or action on amylase activity (Claverie et al. 2003). Amylase secretion required two accessory proteins, PSHAb0130 (possibly an outer membrane associated protein) and PSHAb0132, coding for a conserved secreted protein present in several phytopathogenic γ -proteobacteria. Interestingly, the neighboring glutamate synthesis genes were essential to allow amylase secretion when reconstructed in *E. coli*.

P. haloplanktis has several genes and operons that may play an important role in colonization of both biotic and abiotic surfaces. In particular, up to 16 genes involved in the synthesis of mannose-sensitive agglutinin are located on chrI. These genes have been recently demonstrated to facilitate adhesion to the chitin surface in *V. cholerae* (Meibom et al. 2004). When investigating the formation of biofilms on solid surfaces (many genes are compatible with synthesis of a biofilm such as production of EPS), we did not observe synthesis of a strong biofilm. In contrast, the air-water interface was rapidly occupied by a dense layer of compact cells (Supplemental Fig. 6), suggesting that, for this organism, this is the normal way to concentrate cells and occupy a biotope. The formation of such a biofilm compatible with life in water and scavenging of organic particles that concentrate in the foam of waves has been recently demonstrated in *V. parahaemolyticus* (Enos-Berlage et al. 2005). Genes for the synthesis of polar flagellum are present, and these appendages are visible under the microscope. In salty water the cells are highly motile. However, in contrast to the situation with several vibrios, pseudomonads, and related bacteria, the cells have a reduced motility in low salt media, while they still grow well under such conditions (Supplemental Fig. 7). In minimal medium, the strain grew in a large range of NaCl concentration (0% to up 11% NaCl). In rich media, however, the growth of the strain is remarkably sensitive to the presence of salt. At 20°C, no growth occurs in the absence of NaCl. In remarkable contrast, slow but significant growth is observed at 4°C in the absence of salt, suggesting some adaptation to ice or melting ice water.

Conclusions

P. haloplanktis TAC125 has found many unexpected solutions to cope with cold. Not only does it grow fast under such conditions,

but it displays remarkable resistance to ROS. Moreover, as seen in silico with its proteome composition, it provides a way to resist to the aging features involving asparagine cyclisation and deamidation. This makes this bacterium not only a model for the study of adaptation to cold marine conditions but also a promising tool for biotechnology production of proteins.

Methods

Bacterial strains, growth media, and assays

The *P. haloplanktis* strain TAC125 is deposited and available at the Institut Pasteur Collection (CIP). *Escherichia coli* strains used in this work are strains MG1655, FB8, and its *hns* defective counterpart BE1410 (Laurent-Winter et al. 1997). TAC125 bacteria were grown in TYP rich media: 16 g/L yeast extract (DIFCO) and 16 g/L bacto-tryptone (DIFCO) supplemented with NaCl as required. For growth on specific carbon sources (0.4%), the following minimum medium was used: 10 g/L Na₂HPO₄, 3 g/L KH₂PO₄, 1 g/L K₂SO₄, 20 g/L NaCl, 0.4 g/L MgSO₄ · 7H₂O, 0.1 g/L CaCl₂, 0.018 g/L FeSO₄ · 7H₂O, and 3 g/L NH₄Cl (pH 7). Motility assay of *E. coli* cells with plasmids carrying *P. haloplanktis* genes was performed on semisolid plates with 0.5% NaCl 16 h. For CsrA-dependent storage of glycogen analysis, wild-type MG1655 *E. coli* colonies growing on Kornberg medium with 1% glucose and 100 µg/mL ampicillin with (pDIA616) and without (pcDNA2.1) *csrA* were assayed twice for glycogen accumulation. The plates were stained with iodine vapor after 48 h growth at 25°C as in Liu and Romeo (1997). Oxidative stress adaptation was assayed as follows: after overnight growth in TYP medium with 1% NaCl, bacteria are washed in fresh TYP medium with 1% NaCl and diluted to OD₆₀₀ = 0.15. When the OD₆₀₀ reached 0.5, the culture was separated into four equal parts and then exposed to 0, 15, 20, and 25 mM of H₂O₂. The OD₆₀₀ was measured at various times during growth. Experiments were performed twice at 37°C for MG1655 *E. coli* strain and at 15°C for *P. haloplanktis* TAC125. For electron microscopy cells were stained with 0.1% (v/v) osmium tetroxide prepared in water: 20 µL cells were deposited onto 300-mesh copper grids coated with Formvar (Electron Microscopy Sciences). Excess sample was removed by using Whatman 3MM paper. Bacteria were examined at 75 keV under a Hitachi H600 transmission electron microscope. Image's acquisition was performed with a CCD Advantage HR Hamamatsu camera and the AMT 542 software (Advanced Microscopy Techniques).

Genome sequencing, assembly, and annotation

DNA was isolated from *P. haloplanktis* TAC125 grown in rich medium supplemented with 20 g/L NaCl. The complete genome sequence was determined by using the whole shotgun method (10× coverage, using two plasmid libraries and one BAC library to order contigs). Finishing was performed by PCR amplification from contig's extremities. All rDNA clusters were sequenced individually. After a first round of annotation, regions of lower quality as well as regions with putative frameshifts were resequenced from PCR amplification of the dubious regions.

A first set of potential coding sequences (CDSs) was identified by using the AMIGene software (Annotation of Microbial Genes) (Bocs et al. 2003) trained with a set of CDSs >500 bp from the genomic sequence. Three gene models, computed from the three gene classes identified by codon usage analysis (see below), were then subsequently used together in the core of AMIGene with minimum CDSs length set to 60 bp. This second set of putative genes (made of 3488 CDSs) was submitted to functional

annotation: exhaustive BLAST searches against the UniProt databank were performed to determine significant homology. Protein motifs and domains were documented by using the InterPro databank. In parallel, genes coding for enzymes were classified by using the PRIAM software (Claudel-Renard et al. 2003). PRO-TMHMM and PRODIV-TMHMM were used to identify transmembrane domains (Viklund and Elofsson 2004), and SignalP 3.0 was used to predict signal peptide regions (Bendtsen et al. 2004). Finally, tRNAs were identified by using tRNAscan-SE (Lowe and Eddy 1997).

Sequence data for comparative analyses were obtained from the National Center for Biotechnology Information (NCBI) databank. Putative orthologs between *P. haloplanktis* and the 228 other genomes were defined as genes showing a minimum of 30% identity and a ratio of 0.8 of the length of the smallest protein. Orthology relations were strengthened by synteny detection (i.e., conservation of the chromosomal colocalization between pairs of orthologous genes from different genomes) using the Syntony software, in which all possible kinds of chromosomal rearrangements are allowed (inversion, insertion/deletion). A "gap" parameter, representing the maximum number of consecutive genes not involved in a synteny group was set to five genes. Species-specific genes were identified as having no ortholog in the compared species. Specific regions are defined by at least two consecutive specific genes. Insertion of genes with similarities in the compared species was allowed. A gap parameter, representing the maximum number of consecutive genes with similarities, was set to two genes.

All the data (i.e., syntactic and functional annotations, and results of comparative analysis) were stored in a relational database (using the MySQL DBMS software). Each predicted gene was assigned a unique identifier prefixed with "PSHAA" for chrI and "PSHAB" for chrII. Manual validation of the automatic annotation was performed by using the Web interface MaGe (Magnifying Genomes), which allows graphic visualization of the *P. haloplanktis* annotations enhanced by a synchronized representation of synteny groups in other genomes chosen for comparisons. Translation start codons were corrected based on protein homology, proximity of ribosome-binding site, and relative position to predicted signal peptides when present. To this purpose, we used the Artemis sequence Viewer connected to the MaGe system. The *P. haloplanktis* nucleotide sequence and annotation data have been deposited at EMBL databank under accession number CR954246 and CR954247. The PseudoList database is constructed by using the MySQL DBMS, as previously described (Fang et al. 2005).

Correspondence analysis (Benzécri 1984) was used to analyze the data table with the relative synonymous codon usage values of each annotated gene as well as the table of distribution of amino acids in the proteome of *P. haloplanktis*. Clustering into consistent classes used a second method (dynamic clouds) (Delorme and Henaut 1988) that automatically clusters the objects located close to one another.

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