

# Genome sequence of the hyperthermophilic crenarchaeon *Pyrobaculum aerophilum*

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We determined and annotated the complete 2.2-megabase genome sequence of *Pyrobaculum aerophilum*, a facultatively aerobic nitrate-reducing hyperthermophilic ( $T_{\text{opt}} = 100^{\circ}\text{C}$ ) crenarchaeon. Clues were found suggesting explanations of the organism's surprising intolerance to sulfur, which may aid in the development of methods for genetic studies of the organism. Many interesting features worthy of further genetic studies were revealed. Whole genome computational analysis confirmed experiments showing that *P. aerophilum* (and perhaps all crenarchaea) lack 5' untranslated regions in their mRNAs and thus appear not to use a ribosome-binding site (Shine-Dalgarno)-based mechanism for translation initiation at the 5' end of transcripts. Inspection of the lengths and distribution of mononucleotide repeat-tracts revealed some interesting features. For instance, it was seen that mononucleotide repeat-tracts of Gs (or Cs) are highly unstable, a pattern expected for an organism deficient in mismatch repair. This result, together with an independent study on mutation rates, suggests a "mutator" phenotype.

*Pyrobaculum aerophilum* is a hyperthermophilic ( $T_{\text{max}} = 104^{\circ}\text{C}$ ,  $T_{\text{opt}} = 100^{\circ}\text{C}$ ) and metabolically versatile member of the crenarchaea (Fig. 1), which are predominantly anaerobic respirers. Unlike most hyperthermophiles, *P. aerophilum* can withstand the presence of oxygen, growing efficiently in microaerobic conditions, thus making it relatively easy to work with in the laboratory. Unlike most of its phylogenetic neighbors, the growth of *P. aerophilum* is inhibited by the presence of elemental sulfur, but it grows well anaerobically using nitrate reduction (1). Here we have determined the complete genome sequence of *P. aerophilum* IM2, which was isolated from a boiling marine water hole at Maronti Beach, Italy (1). We obtained the sequence by a low coverage random shotgun sequencing strategy, with gap closure and resolution of ambiguities aided by the creation of a genomic fosmid map (2). We present an overview of the features and content of the genome, including a possible explanation of the organism's intolerance to sulfur and evidence of a possible lack of mismatch repair activity. Studies of the genus *Pyrobaculum* provide important opportunities for understanding the boundaries of life in extreme habitats. In a recent molecular sampling of a deep subsurface geothermal water pool, the only organisms detected were hyperthermophilic archaeal members closely related to *Pyrobaculum* (3).

The *P. aerophilum* IM2 genome has 2,222,430 base pairs, 51% G + C. Two thousand five hundred and eight-seven protein-coding regions were designated, with an average length of 759 amino acids, covering 88% of the genome (Table 1). Circular and linear maps of genome features, including many features discussed below, are published as supporting information (Figs. 4 and 5) on the PNAS web site, [www.pnas.org](http://www.pnas.org).

## Methods

Fosmid and pUC18 libraries were constructed as described (2). Approximately 26,000 sequences were obtained from  $\approx 14,000$  pUC18 clones by using both Dye-terminator (18,000 sequences) and Dye-primer (8,000 sequences) chemistry on Applied Biosystems 373 and 377 sequencing machines. Twenty-three thousand four hundred and thirty-seven sequences were used to generate the

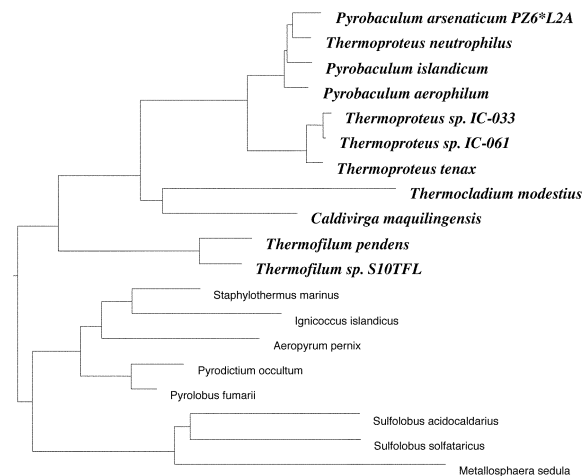


Fig. 1. Small subunit rRNA-based phylogenetic tree of the crenarchaea. Constructed by Harald Huber, Regensburg University, Regensburg, Germany.

final assembly, giving  $\approx 4$ -fold coverage of the genome. Sequence assembly and editing were done by using PHRED/PHRAP/CONSED software (<http://www.phrap.org>). Sequence assembly was verified by comparison to sequence tagged sites of the published fosmid map (2) and PCR where necessary. After extensive manual inspection and resequencing,  $\approx 3\%$  of the genome was left covered by a single clone and  $\approx 6\%$  left unsequenced on both strands or with complementary sequencing chemistries.

Annotation was initially assisted by use of the MAGPIE suite of software for automated genome interpretation (4). Coding regions were identified by using GENEMARK (5), and exact start site predictions were refined with GENEMARKS (6). FASTA (7) and BLAST (8) programs were used to compare sequences against National Center for Biotechnology Information nonredundant protein and nucleotide databases. Predicted proteins were also compared against the PFAM database of protein domain hidden Markov models (9) by using HMMER (<http://hmmer.wustl.edu/>). Where possible, the most likely major function of each predicted protein and its associated category was determined by manual inspection of results from the above analyses with emphasis placed on matches to characterized proteins or genes. Regions of synteny between the *P. aerophilum* genome and other completed genomes (at the protein-coding level) were identified and used to help specify functions and functional categories. Transfer RNAs were predicted by tRNAscan-SE (v. 1.21) (10) and manually refined by Todd Lowe

Data deposition: The sequence reported in this paper has been deposited in the GenBank database (accession no. AE009441).

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**Table 1. Chromosomal coding sequences**

Similar to	No.	Av bp	Av pl
Known proteins	992 (38%)	965	8.4
Proteins of unknown function	577 (22%)	695	8.2
Only within <i>P. aerophilum</i> genome	302 (12%)	693	8.8
None	716 (28%)	553	8.5
Total	2,587		

(<http://rna.wustl.edu/tRNAdb>). Homologs of small nucleolar RNAs were identified by Todd Lowe (11). Repeat sequences were initially identified by using MIROPEATS (12). Transmembrane helices were predicted for the T/A box families by using TMHMM v. 2.0 (13)

## Replication and Repair

**Mismatch Repair and Mononucleotide Runs.** Some of the most intriguing discoveries coming from archaeal genome sequencing projects are those pertaining to genes involved in DNA repair. The two major mechanisms for avoiding mutations during DNA replication are immediate editing of the growing strand by the DNA polymerase and detection and correction of mismatches soon after replication by the mismatch repair system (see reviews in refs. 14–16). Homologs of the *Escherichia coli* proteins involved in mismatch repair have been found in humans, and damage to them is involved in inherited predispositions to colon (HNPCC), endometrial, and ovarian cancer (17–20). However, homologs of the mismatch repair proteins have not been detected in the *P. aerophilum* genome, nor have they been found in any of the six published genomes of thermophilic archaea [*Methanobacterium thermoautotrophicum* and *Archaeoglobus fulgidus* each have a divergent MutS (MutS2) homolog without a MutL homolog]. It remains to be seen whether mismatch repair activities can be detected in these organisms. However, there are indications that *P. aerophilum*, at least, may be mismatch repair deficient.

During assembly and editing of the *P. aerophilum* genome, 10 regions were identified where high-quality sequences from different clones gave varying lengths of a mononucleotide run (Table 2). Further characterization of cultures grown from single colonies

**Table 2. Mononucleotide runs**

Genome position	Base	Lengths*	Context (intergenic region), bp <sup>†</sup>
40977	G	11,15	Intergenic (1,176)
93514	G	11,11,11	Intergenic (121)
159685	A	9	Intergenic (400)
221366	C	11,12,12,16	Intergenic (548)
380761	G	12,13,14,14	Intergenic (988)
463202	C	14,15	Within gene
496647	G	11,11,13	Within gene
832639	T	8,10	Intergenic (122)
839489	A	9,9	Intergenic (121)
909549	C	7,8,8,9	Within gene
1028272	T	11,11,11	Intergenic (253)
1059643	A	9,9,9,9,9	Intergenic (782)
1110478	C	9,9,10,10,10,13	Within gene
1181593	A	9,9	Intergenic (226)
1217896	C	12	Within gene
1225895	C	12,15	Intergenic (126)
1293892	A	9,9,9,9,9	Intergenic (85)
1818426	C	11,12,15	Intergenic (140)

All mononucleotide runs of nine or more bases found in the *P. aerophilum* genome are listed.

\*Each of the lengths listed corresponds to sequences (one or more) from one genomic library clone.

<sup>†</sup>The size of each intergenic region is given in brackets. The average size of intergenic regions in the genome is 121 bp.

confirmed that these variable length mononucleotide runs were because of rapid fluctuations in the genome (M. M. Slupska, A. Conrad, L. Garibyan, S.T.F., J. Chiang, J. Pan, H. A. Nguyen, and J.H.M, unpublished work). Mononucleotide runs are particularly susceptible to strand slippage, and frequent frameshifting at such runs has been demonstrated in mismatch repair-deficient strains for both *E. coli* and yeast (21–23). An increased rate of variation of mononucleotide run lengths was also discovered during the genome sequencing project of *Campylobacter jejuni* (24). This genome has only a divergent MutS2 and no detectable MutL homolog. The data in Table 2 indicate that repeat-tracts are more unstable in runs of Gs (or Cs) than in runs of As (or Ts), as has been found in other systems (see, for instance, ref. 21). In *C. jejuni*, the repeat-tract sequences are proposed to have a function in producing variable surface structures that can be important for virulence and rapid adaptation to changing environments. It does not appear that the unstable mononucleotide runs in *P. aerophilum* have any obvious function, because they are distributed within tolerant regions, such as suspected pseudogenes and long intergenic regions. Rather, it seems that the repeat-tract instability is a consequence of the lack of a mismatch repair system. If *P. aerophilum* is indeed lacking mismatch repair, then an increased rate of frameshifting would be expected. During annotation, 32 genes were clearly found to have one or more frameshifts, many occurring at mononucleotide runs of from four to seven bases. Undetected frameshifts near the ends of genes, or in less conserved proteins, could double or triple this number.

A study of specific base substitution rates in *P. aerophilum* (M. M. Slupska, A. Conrad, L. Garibyan, S.T.F., J. Chiang, J. Pan, H. A. Nguyen, and J.H.M, unpublished work) reveals that transitions occur at the level expected of a mismatch repair deficient mutator. Although lack of a mismatch repair system is advantageous under certain selective conditions (see, for instance, ref. 25) and as a way of generating diversity and responding to changing environments (26–30), a permanent mutator lifestyle, in which the absence of the function generates high rates of mutation (a mutator phenotype), can have serious consequences as deleterious mutations accumulate (22). Other enzymes might partially compensate for the lack of a canonical mismatch repair system. Higher fidelity polymerases or editing functions might be used during replication, or additional as yet uncharacterized systems might operate. However, the repeat-tract instability observed here indicates that like *C. jejuni* (24), the high-temperature archaea such as *P. aerophilum* might be extraordinary examples of organisms that can survive as permanent mutators (we cannot at this stage rule out the possibility that the *P. aerophilum* species is not a mutator, but the IM2 isolate is a mutator variant). Studies with *Mycobacterium smegmatis* have also led to the idea that *M. smegmatis* and *Mycobacterium tuberculosis* might often have a mutator lifestyle (31).

**Base Excision Repair.** Repair of damaged or altered bases often begins with excision of the altered base by DNA glycosylases and abasic-site endonucleases. The major enzyme in *P. aerophilum* cell free extracts responsible for removing uracil from DNA has been identified and characterized (32). It is similar in sequence to the recently characterized uracil DNA glycosylases of *Thermotoga maritima* and *A. fulgidus* (33). Most identified archaeal DNA glycosylases are in the MutY/Nth family, with every archaeal genome having from one to three homologs. *P. aerophilum* has three MutY/Nth homologs, two of which have been functionally characterized. Pa-MIG has been shown to have DNA glycosylase activity specific to U/G and T/G mismatches as well as an uncoupled AP lyase activity (34). The second characterized *P. aerophilum* DNA glycosylase (PaNth) recognizes and begins repair of pyrimidine adducts in a manner similar to *E. coli* endonuclease III (EcNth). It has DNA glycosylase/ $\beta$ -lyase activity with the modified pyrimidine base 5,6-dihydrothymine (DHT), which is enhanced when the DHT is paired with G (35).



DNA polymerases, one of each of the B1, B2, and B3 subfamilies (38). The *P. aerophilum* B3 DNA polymerase shares 78% amino acid identity with a recently cloned and characterized DNA polymerase from the closely related *Pyrobaculum islandicum*. This *P. islandicum* enzyme was shown to have 3' to 5' exonuclease activity and, under suitable assay conditions for PCR, to amplify DNA fragments of up to 1,500 bp (37).

As with all archaea, the other replication factor homologs detected in the genome are more similar to their eukaryotic counterparts than bacterial. They include two copies of the sliding clamp processivity factor (proliferating cell nuclear antigen-like), one large subunit, and two copies of the small subunit of the clamp loading protein (replication factor C), DNA ligase, minichromosome maintenance protein, and a possible origin recognition protein (*orc/cdc6*). Single strand-binding protein (replication factor A) was not detected.

**Cell Division.** Surprisingly, although FtsZ homologs are found in the euryarchaea and FtsZ ring structures have been visualized in *Haloflex mediterranei* (39), no homologs of FtsZ have been found within the crenarchaea, including *P. aerophilum*. A type of “snapping division” has been visualized for *Thermoproteus tenax* (40), which may be an example of a crenarchaeal FtsZ independent mechanism (41).

**Transcription and Translation.** It has recently been shown experimentally and computationally that *P. aerophilum* mRNAs generally lack 5' untranslated regions, and it was shown computationally that *Aeropyrum pernix*, the other crenarchaea with an available complete genome, probably also uses mainly leaderless transcripts (42). Thus the mechanism for translation initiation at the 5' end of transcripts must be fundamentally different for these crenarchaea from the mechanism generally used by the characterized euryarchaea in which the ribosome binds to a Shine–Dalgarno site upstream of the translation start site. One might expect to find similarities to the eukaryotic translation initiation system in which the ribosome is first directed to a cap structure on the 5' end of the mRNA. However, the genome in this case has not provided us clues as to the possible mechanism. The set of ribosomal proteins and translation initiation factors found in the genome is similar to those found in euryarchaeal genomes.

*P. aerophilum* has a similar set of translation factors as have been identified in other archaeal genomes. Aminoacyl tRNA synthetases have been identified for all amino acids except glutamine. Glu-tRNA (Gln) amidotransferase subunits are found; thus, as seen in other archaea and Gram-positive bacteria (43), glutamine tRNAs are probably amino-acylated with glutamic acid, followed by a transamidation to yield the glutamyl-tRNA. Two apparent asparaginyl-tRNA synthetases are found, as well as two tyrosyl-tRNA synthetases. The C-terminal section of the canonical methionyl-tRNA synthetase is found as a separate ORF in the *P. aerophilum* genome, 1,334 bases upstream of the N-terminal gene.

No evidence was found for the use of the 21st amino acid selenocysteine. No matches were found to the set of genes coding for the proteins involved in the selenocysteine incorporation system. No selenocysteine tRNA was found, and no UGA codons were found in known selenocysteine positions.

**Nonprotein Coding RNA.** After careful manual inspection of the results from the tRNAscan-SE program, 46 tRNAs and 1 pseudogene were identified (T. M. Lowe, personal communication). This corresponds to exactly one copy of each expected tRNA necessary to decode all 61 canonical codons. The most interesting feature of the *P. aerophilum* tRNAs is the prediction of large numbers of noncanonically placed introns (at least 11 of the 15 detected introns). This was also observed in the *A. pernix* genome (T. M. Lowe, personal communication) and is surprising in that it contrasts with the high consistency of intron placement among the vast

**Table 3. T/A box families**

Genome position	Run	Genome strand	No. bases from start codon	Gene identifications
1864462	8Ts	–	–24	PAE3129,3125
1895538	8Ts	+	–7	PAE3182
1957470	8Ts	–	–7	PAE3261,3260
159693	9Ts	–	–5	PAE0282,0280
163987	8T/As	+/-	-20/-5	PAE0293/0292
577313	8Ts	–	–3	PAE0985,0984
624765	8Ts	+	–9	PAE1067,1068
633675	8Ts	+	–6	PAE1079,1081
639492	8Ts	+	–6	PAE1089,1092
806449	8Ts	+	–4	PAE1358
839489	9As	+	–22	PAE1407
1007991	8As	+	–21	PAE1720
1013101	8As	–	–21	PAE1727
1028282	11As	–	–20	PAE1756,1755
1049571	8As	–	–17	PAE1783
1059643	9As	+	–15	PAE1802,1803
1071224	8As	+	–21	PAE1819
1181593	9As	+	–22	PAE2005,2006
1293892	9As	+	–20	PAE2186
1736810	8As	–	–24	PAE2924

Intergenic runs of eight or more As or Ts and their associated gene families are listed in the order in which they appear on the circular genome, starting at the switch between As and Ts. The eight runs that are not listed either fell within genes or were 73 bases or more away from a start codon. The run at 163987 is upstream of two start codons, one on each strand. Listed pairs of genes have the run upstream of the first gene and are always immediate neighbors (and in the same orientation), even if the gene identifiers are not contiguous numbers.

majority of both archaeal and eukaryal tRNAs. Current tRNA annotations are available in the Genomic tRNA Database (T. Lowe, <http://rna.wustl.edu/tRNAdb/>).

Homologs of the eukaryal small nucleolar RNAs have been identified in archaea, with generally increasing numbers found in the genomes of organisms with higher growth temperatures (11). This may be because of an increased need for methylations of structural RNAs in higher-temperature environments (11). A record 53 homologs were identified in *P. aerophilum* by computational methods (T. M. Lowe, personal communication), most of which could be experimentally confirmed by primer extension (T. M. Lowe, A. Omer, L. Garibyan, S.T.F., J.H.M., and P. P. Dennis, unpublished work).

A 20-mer (ggcggcggcgtgggggttt) inverted repeat was found surrounding the single rRNA operon, 51 and 15 bases upstream and downstream, respectively. The single 5S rRNA gene was not near the 16S-23S operon. Concurring with a previous report for *P. aerophilum* IM2 (44), a 713-bp intron was found within the 16S rRNA gene. Otherwise, introns were found within 15 tRNAs and not within any other genes.

**Metabolism.** *P. aerophilum* is metabolically versatile, able to grow using organic and inorganic substrates for both aerobic and anaerobic respiration. Maximum cell densities are obtained with complex organics as substrates, although in the absence of organic material, *P. aerophilum* will grow chemolithoautotrophically using molecular hydrogen or thiosulfate (1). A set of genes for a complete citric acid cycle (TCA cycle) using 2-oxoacid/ferredoxin oxidoreductase was found.

**Glyoxylate Cycle.** Consistent with *P. aerophilum*'s growth on acetate (1), candidates for the key enzymes of the glyoxylate cycle, isocitrate lyase and malate synthase, have been found in the genome. The glyoxylate cycle is required to synthesize precursors for carbohydrates only when the carbon source is a C2 compound. To date, the only demonstrated occurrence of the glyoxylate cycle within the archaea is for a halophilic euryarchaeota, *Haloflex volcanii* (45).

PAE3129 GGAAAAGGTATTTAAATCCTTTTTTTTGGAGTATTATGTCATAAAGGGTATATG  
 PAE3182 AGCATATGAGCAGTGGACTGTAAGTTTTTTAAACTCTTTTTTTTGAACGTATG  
 PAE3261 CACGGATAGATTCCGCTGTGGTAAATTTTTTAAACTCTTTTTTTTGGAGTACATG  
 PAE0282 TCAGAGCCTATTCGCCCTCAAGATTTTATAAATCCGTTTTTTTTTGCACGCATG  
 PAE0293 TTTCTGTAGGCTGGGACATAAGCTTTTTTTTCTGTATTTTTAAGCCTTATG  
 PAE0985 AGCTAACGGCTTTACTGAAGTAAACTTAAAAATACCAGTTTTTTTTTGCATG  
 PAE1067 AGATTATCCAGTGAGAGTAAATTTTTTAAAGCTGTTTTTTTTTGACACTGTTG  
 PAE1079 GAATTCACCTGCCGCTAGCGTAATGTTTATAAAGCCGTTTTTTTTTGACATATG  
 PAE1089 TAGGAAATACTACTATAGCGCAATGTTTATAAATCCGTTTTTTTTTGAGGCATG  
 PAE1358 TGGCGCAGCCGCTGGATAAATTTTTTAAAGGAAATAGCTCTTTTTTTTCCAATG  
 PAE1407 TTTCTGTAAGTTTTACAGCGAAAAAAATACTGGGAGTTTTCTCTCTATATG  
 PAE1720 ACTATTCGACAATAAATAACGAAAAAAGACTGTAATTTCTTGGGACTATG  
 PAE1727 AATTACGACAAAACGTGAAAGTAAAAAAGACTGGAAAAATCAGTAAAAATATG  
 PAE1756 TTTAGTAAAGTTGATGAAAGTAAAAAAGACGGAGCTCCCGTCTCCTATG  
 PAE1783 ATATAAGCTAAAGCGCGCAAAATTTTTAAAAAAGATTTTCAACAAAATAAGT  
 PAE1802 AGACTGTACAGAATACGAGAAAAACTTAAAAAAGGTTTCTAATCCTTATG  
 PAE1819 CTTTTCATTTTTTGGCTAACGCAAAAAAAGACCGGAATTACATGACGCCATG  
 PAE2005 AATTGCTATTGGCTAAAGTAAAAAAGGTTTAGGACTTACCGCCATTATG  
 PAE2186 ATATCTCAGTTGAACAACAGTGAAAAAATACTTTGTCTAATTACTACATG  
 PAE2924 ATACGCGAAAAATCAACGAAAAAATAATACGAAAAAATACTTACTTATG  
 -40            -30            -20            -10

**Fig. 3.** Upstream regions of T/A box family genes. T/A boxes are underlined. Start codons for the associated genes are in bold. The second possible associated start codon, on the reverse strand, is shown underlined for PAE0293.

Interestingly, the *H. volcanii* malate synthase falls into a distinct class separate from the previously known sequence-based classes (A and G) (45). However, this class may not be typical of the archaea, as the *P. aerophilum* putative malate synthase falls clearly within class A, sharing 30% amino acid identity (BLAST Expect =  $e^{-23}$ ) with a yeast glyoxysomal malate synthase over the first two-thirds of the yeast protein length. Further, a database search revealed a putative malate synthase in the recently published *Sulfolobus solfataricus* genome (46), which shares 60% amino acid identity over the full length (813 amino acids) of the *P. aerophilum* protein.

**2-Oxoacid Dehydrogenase.** Another metabolic similarity between *P. aerophilum* and *H. volcanii* is the presence of genes with highly significant homology to 2-oxoacid dehydrogenase multienzyme complexes. 2-Oxoacid dehydrogenases are used in eukarya, and most aerobic bacteria to convert pyruvate to acetyl-CoA, whereas archaea are generally known to use a simple pyruvate oxidoreductase (47). An operon containing genes homologous to 2-oxoacid dehydrogenase enzymes: E1 (2-oxoacid decarboxylase), E2 (dihydrolipoamide acetyltransferase), and E3 (dihydrolipoamide dehydrogenase) has been characterized for *H. volcanii*; however, the enzymatic activity was not detected in the organism (47). The *P. aerophilum* genome contains an apparent operon of not only the E1, E2, and E3 genes but also genes for lipoic acid synthetase and lipoate protein ligase B, known in eukarya and bacteria to act in the biosynthesis and attachment (to the E2 subunit) of lipoic acid cofactors (47).

**Glycolysis/Gluconeogenesis.** As with the other completed archaeal genomes, several of the genes involved in glycolysis/gluconeogenesis have been identified; however, no ORFs with sequence similarity to phosphofructokinase or fructose biphosphate aldolase are found. On the basis of a 25-aa N-terminal sequence from a recently characterized ATP-dependent 6-phosphofructokinase from the hyperthermophilic archaeon *Desulfurococcus amylolyticus* (48), we have annotated a *P. aerophilum* ORF (PAE0835) as a possible phosphofructokinase (Pfk). This candidate Pfk has a high-scoring BLOCKS (49) match to the INTERPRO (50) PfkB family of carbohydrate kinases (Entry IPR002173) and is of similar molecular mass (32 kDa) to the *D. amylolyticus* Pfk (33 kDa). PAE0835 has a reasonable match at the N-terminal to the *D. amylolyticus* N-terminal fragment (7 of 25 residues are identical with no gaps); however, a strong homolog of PAE0835 ( $E = 10^{-27}$ ), in the genome of the crenarchaeon *A. pernix* (APE0012), has a more conclusive

match to the *D. amylolyticus* N-terminal (11 identities of 25 with no gaps). Homologs of APE0012 and PAE0835, generally annotated as sugar kinases, are found in all of the completed archaeal genomes as well as in many bacteria and eukaryotes.

**Respiratory Proteins.** Although the cultivated thermophilic crenarchaeota are almost exclusively anaerobic respirers, the three crenarchaeons with genome sequences available, *P. aerophilum*, *A. pernix* (51), and *S. solfataricus* (<http://www-archbac.u-psud.fr/projects/sulfolobus/>) are able to grow aerobically. Numerous respiratory chain proteins are found in the *P. aerophilum* genome, including a cluster containing cytochrome C oxidase subunits and assembly factor, cytochrome *b/b6*, *senC*, *nosD*, and two iron sulfur proteins. Three putative Rieske iron-sulfur proteins were identified in the genome, one of which, designated ParR, has been expressed and partially characterized (52). *P. aerophilum* can also grow anaerobically, by nitrate reduction, using the denitrification pathway (refs. 1 and 53). Five putative nitrate reductase subunits were found in the genome, with the major ( $\alpha$  and  $\beta$ ) subunits sharing 50–55% amino acid identity to their *Bacillus subtilis* homologs. Three putative nitrite reductase subunits were found as well as a nitric oxide reductase subunit with 28% amino acid identity to the NorZ of the  $\beta$  proteobacteria *Ralstonia eutropha*. Surprisingly, a set of nitrate reductase genes similar to the *P. aerophilum* genes (60% amino acid identity for the major subunits) were found in the *A. pernix* genome, although *A. pernix* has been reported to be strictly aerobic (54).

**Sulfur Metabolism.** The presence of elemental sulfur is lethal to cultures of *P. aerophilum* (1). This feature distinguishes *P. aerophilum* from its close phylogenetic neighbors, including two other cultured species of *Pyrobaculum* (*islandicum* and *organotrophum*). *P. aerophilum* does have a set of three sulfur metabolism genes that, when intact, could be used for either sulfate reduction or sulfur oxidation. However, both of the adenylylsulfate reductase subunits, for the middle step, are disrupted. The  $\alpha$  subunit has a premature stop codon three quarters into the gene, and the  $\beta$  subunit has a frameshift near the middle of the gene. The proteins involved in this potentially bidirectional pathway (sulfate adenylyltransferase, adenylylsulfate reductase, and sulfite reductase) are well conserved across phylogenetically distant prokaryotes. It is possible that the loss of function of the adenylylsulfate reductase is responsible for the organism's sensitivity to elemental sulfur, and that introducing a functional replacement enzyme could restore resistance and/or add a new definable metabolic capacity. In this case, these genes could provide a plasmid-borne selectable marker that would be a significant step toward the development of a genetic system. The closely related *P. islandicum* (Fig. 1) is a likely source of the genes, as sequences of the three subunits of the neighboring sulfite reductase have been reported (55) (66, 69, and 78% nucleotide identity to *P. aerophilum*).

These sulfur metabolism genes are all found within a 100-kb region of the *P. aerophilum* genome. Putative enzymes thiosulfate sulfurtransferase and phosphoadenosine phosphosulfate reductase (PAPS reductase) were also found in this region.

**PaRep2 Family.** *P. aerophilum* does not have any large families of exceptionally well conserved (98–100% nucleotide identity) repeats with ORFs, reminiscent of active transposons or insertion elements. However, the genome does have a very large and unusual family, comprising 3.3% of the genome, whose members are remarkably variable in both length and level of conservation. The repeat generally carries three ORFs (paRep2a2, 2a1, and 2b) divergently oriented from a central point between 2a1 and 2b. There are 22 copies that contain both 2a and 2b regions, 9 with only 2a regions, and 14 with only 2b regions. The highest conservation of each family member with another ranges from 100% nucleotide identity over 876 bases (and 96% identity over 3,770 bases) to less than 30%

amino acid identity. They are extremely variable in size, ranging from 98 to 5,532 bases, primarily because of deletions from the outer 3' ends of the paRep2as and the central 5' ends of the paRep2bs. Multiple alignments (Fig. 6, which is published as supporting information on the PNAS web site) show numerous frameshifts and internal stop codons, perhaps indicating that most family members are no longer functional and are undergoing degradation. Nucleotide alignment of the three largest and most conserved copies (Fig. 6c) reveals several interesting features of this family. The unequal pattern of conservation across the length of the alignment (ranging from 55% in some regions to 100% in others) suggests that partial recombination has occurred between at least two of these sequences. All of the eight small insertion/deletions between the three sequences, found distributed over the 4,700-bp shared region, are in multiples of 3 (thus not causing frameshifts), suggesting that a selective pressure has maintained the ORFs because of the duplication/recombination events that created them. Surprisingly, however, all three of the sequences share an identical frameshift within the 4,050-bp paRep2b ORF (at approximately position 1020). A graphical outline of the 45 paRep2 regions (Fig. 7, which is published as supporting information on the PNAS web site) shows further the surprising patterns and variation among this family.

**T/A Box Families.** A few distantly related families of *P. aerophilum* specific proteins have runs of 8–11 As or Ts in the 24 bases preceding their start codons (Table 3 and Fig. 3). Remarkably, whether the run is of As or of Ts (on the strand coding for the gene) seems to depend on its location in the genome. There are 10 upstream A runs found starting at genome positions 839489 until the last at position 1864462, covering roughly half of the genome, whereas the 10 upstream runs in the rest of the circular genome are all of Ts. The pattern does not depend on genome strand, as the genes fall on both strands within each region (Table 3). Many of the associated genes share similar characteristics and are part of a gene pair in which the downstream genes also share similar character-

istics with each other (Table 4, which is published as supporting information on the PNAS web site). The first gene of each of the pairs codes for a large protein of between 77 and 90 kDa and usually has a single transmembrane helix predicted at the N terminus. The second gene of each of the pairs codes for a small protein of between 16 and 23 kDa and always has from four to six predicted transmembrane helices. There are other members of these gene families that do not have T or A runs upstream. No gene pair families with similar molecular weights and numbers of transmembrane helices were evident in any of the published microbial genomes. The reason for the striking dichotomous distribution of the T/A boxes is unclear. Further characterization of these interesting families could possibly lead to insights about genome organization.

A genome sequence is important, because it provides both an invaluable tool for ongoing biological research and a generator of new hypotheses for further research. The *P. aerophilum* genome not only provides valuable basic data in an understudied branch of life but also poses many questions warranting several new lines of research.

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