

Distribution, structure and diversity of “bacterial” genes encoding two-component proteins in the Euryarchaeota

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Summary The publicly available annotated archaeal genome sequences (23 complete and three partial annotations, October 2005) were searched for the presence of potential two-component open reading frames (ORFs) using gene category lists and BLASTP. A total of 489 potential two-component genes were identified from the gene category lists and BLASTP. Two-component genes were found in 14 of the 21 Euryarchaeal sequences (October 2005) and in neither the Crenarchaeota nor the Nanoarchaeota. A total of 20 predicted protein domains were identified in the putative two-component ORFs that, in addition to the histidine kinase and receiver domains, also includes sensor and signalling domains. The detailed structure of these putative proteins is shown, as is the distribution of each class of two-component genes in each species. Potential members of orthologous groups have been identified, as have any potential operons containing two or more two-component genes. The number of two-component genes in those Euryarchaeal species which have them seems to be linked more to lifestyle and habitat than to genome complexity, with most examples being found in *Methanospirillum hungatei*, *Haloarcula marismortui*, *Methanococcoides burtonii* and the mesophilic Methanosaecinales group. The large numbers of two-component genes in these species may reflect a greater requirement for internal regulation. Phylogenetic analysis of orthologous groups of five different protein classes, three probably involved in regulating taxis, suggests that most of these ORFs have been inherited vertically from an ancestral Euryarchaeal species and point to a limited number of key horizontal gene transfer events.

Keywords: histidine kinase, hybrid kinase, response regulator.

Introduction

Two-component systems are one of the key means by which bacteria respond to environmental changes (Hoch 2000, Stock et al. 2000, Alves and Savageau 2003, Hellingwerf 2005). They are assumed to be of bacterial origin, having radiated into archaea and some eukaryotes by horizontal gene transfer (HGT) (Koretke et al. 2000). Two-component systems consist of a sensor and a response protein. The sensor protein is char-

acterized by a histidine kinase (HK) made up of two main domains, a phosphoacceptor (HisKA) and a histidine kinase ATPase (HATPase) and, in many cases, other sensory domains are present (Galperin et al. 2001, Zhulin et al. 2003). The response protein (response-regulator, RR) is characterized by a response regulator domain that has a conserved aspartate residue. The histidine kinase autophosphorylates a conserved histidine residue in response to a signal and the phosphate group is then transferred to the conserved aspartate residue of the response-regulator. The transfer of the phosphate group to the response-regulator elicits a response causing a change in taxis, development or gene expression. Histidine kinases and response-regulators are sometimes found together in a single polypeptide known as a hybrid kinase (Hoch 2000, Stock et al. 2000).

The recognition of the Archaea as a distinct division of life has been strengthened by the availability of a number of complete genome sequences, representing three phyla (Euryarchaeota, Crenarchaeota and Nanoarchaeota). This has, in turn, enabled a more rigorous phylogenetic analysis based on the fusion of ribosomal protein sequences (Matte-Tailliez et al. 2002, Brochier et al. 2004, Baptiste et al. 2005, Makarova and Koonin 2005) and clusters of conserved orthologous genes (COGs) (Makarova and Koonin 2003). Analysis of genome sequences has revealed genes of bacterial origin in the genomes of archaea and vice versa (Nelson et al. 1999). The importance of HGT in the evolution of prokaryotes and the implications for phylogeny and definition of species is still being discussed (Ochman et al. 2000, Forterre et al. 2002, Boucher et al. 2003, Koonin 2003, Kurland et al. 2003, Lawrence and Hendrickson 2003).

For bacteria, it has been shown that the number of two-component genes possessed by an organism is related to the complexity of its genome, its physiology and the changeability of its habitat (Ashby 2004, Galperin 2005). The greater the value of any of those parameters, the greater the need for regulation of cellular activities.

The aim of this study was to analyze the complement of genes in archaeal genomes that could encode two-component proteins. The putative two-component proteins were classified by their domain structure, and the number of each class was

determined for each species. Potential orthologous groups and those that may be part of operons, with two or more two-component genes, are indicated. Phylogenetic analysis of possible orthologous groups representing five classes of protein, three associated with taxis, is shown.

Materials and methods

Genome sequence data

The list of publicly available Euryarchaeal genome sequences is shown in Table 1, along with brief details of the habitat,

physiology, genome size, putative number of open reading frames (ORFs) and the abbreviation used with gene sequences (Makarova and Koonin 2003). The sequences and annotations for the annotated sequences (up to October 2005) were accessed at the Integrated Microbial Genomes (IMG) server (<http://img.jgi.doe.gov/pub/main.cgi>) and HaloLex (<http://www.halolex.mpg.de/>). The identity of potential two component genes was determined by reference to the published assignments located at <http://www.tigr.org/tdb/> (Bult et al. 1996, Smith et al. 1997, Kawarabayasi et al. 1998, Klenk et al. 1997, Ng et al. 2000, Deppenmeier et al. 2002, Slesarev et al. 2002,

Table 1. Strain description of members of the Euryarchaea for which there are publicly available genome sequences as of October 2005. The gene name prefix is used with the gene designations to aid in identification of the species.

Strain	Genus	Gene name prefix	Temp opt. (°C)	Physiology	Genome size (MB)	No. protein-coding genes
<i>Archaeoglobus fulgidus</i> DSM 4304	Archaeoglobales	Aful	83	Anaerobic, sulphate-reducing chemolitho- or chemorganoheterotroph	2.18	2,456
<i>Ferroplasma acidarmanus</i> Incomplete	Thermoplasmatales		40	Acidophile	1.97	1740
<i>Haloarcula marismortui</i> NRC-1	Halobacteriales	Hma	37	Aerobic halophilic chemorganotroph	4.3	4,242
<i>Halobacterium</i> sp.	Halobacteriales	Halo	37	Aerobic halophilic chemorganotroph	2.57	2,656
<i>Methanocaldococcus jannaschii</i> DSM 2661	Methanococcales		85	Chemolithoautotroph, anaerobe, methanogen	1.66	1,758
<i>Methanococcus maripaludis</i> S2	Methanococcales	Mmar	37	Mesophilic anaerobe, hydrogenotrophic methanogen	1.67	1,742
<i>Methanopyrus kandleri</i> AV19	Methanopyrales		110	Chemolithoautotroph, anaerobe, methanogen	1.69	1,691
<i>Methanococcoides burtonii</i> DSM 6242 Incomplete	Methanosaecinales	Mbur	~0	Anaerobe, psychrotolerant methanogen	~2.6	~2872
<i>Methanosaecina acetivorans</i> C2A	Methanosaecinales	Mace	37	Chemoorganoheterotroph, anaerobe, N ₂ -fixing, methanogen	5.75	4,540
<i>Methanosaecina barkeri</i> str. fusaro	Methanosaecinales	Mbar	37	Chemoorganoheterotroph, anaerobe, methanogen	4.87	3,380
<i>Methanosaecina mazei</i> Go1	Methanosaecinales	Mmaz	37	Chemoorganoheterotroph, anaerobe, N ₂ -fixing, methanogen	4.1	3,371
<i>Methanospirillum hungatei</i> JF-1	Methanomicrobiales	Mhun	37	Chemoorganoheterotroph anaerobe, methanogen	3.53	3,356
<i>Methanothermobacter thermautotrophicus</i> str. Delta H	Methanobacteriales	Mthe	65	Chemolithoautotroph, anaerobe, N ₂ -fixing methanogen	1.75	1,914
<i>Natronomonas pharaonis</i>	Halobacteriales	Nph	20	Haloalkaliphile	2.75	2,843
<i>Picrophilus torridus</i> DSM 9790	Thermoplasmatales		50	Extreme acidophile	1.58	1,582
<i>Pyrococcus abyssi</i> GE5	Thermococcales	PAB	96	Anaerobic heterotroph	1.76	1,769
<i>Pyrococcus furiosus</i> DSM 3638	Thermococcales		96	Anaerobic heterotroph	1.91	2,115
<i>Pyrococcus horikoshii</i> OT3	Thermococcales	PH	98	Anaerobic heterotroph	1.74	2,110
<i>Thermococcus kodakaraensis</i>	Thermococcales	Thermoco	102	Anaerobic heterotroph	2.09	2,358
<i>Thermoplasma acidophilum</i> DSM 1728	Thermoplasmatales		59	Fac. anaerobe, chemorganotroph, thermoacidophile	1.56	1,482
<i>Thermoplasma volcanium</i> GSS1	Thermoplasmatales		60	Fac. anaerobe, chemorganotroph, thermoacidophile	1.55	1,499

Galagan et al. 2002, Cohen et al. 2003, Baliga et al. 2004, Falb et al. 2005). This was supplemented by BLASTP (Altschul et al. 1997) searches of each genome with a battery of two component domains (domains used include receivers from CheY and OmpR, HisKA/HATPase and Hpt; see Table A1) from *Methanosarcina acetivorans* and *E. coli* K12 at IMG (<http://img.jgi.doe.gov/pub/main.cgi>), The Integrated Genome Resource (TIGR, <http://www.tigr.org/tdb/>) or the National Center for Biotechnology Information (NCBI, <http://www.ncbi.nlm.nih.gov/>).

Bioinformatic analysis

Putative two-component domains were initially assigned in ORFs by Pfam batch analysis (<http://www.sanger.ac.uk/Software/Pfam/>; Bateman et al. 2004). Domains were recorded for each two-component gene only if they were scored as “Pfam’s trusted match thresholds.” Domain assignments were checked and modified using the more extensive domain assignments at InterPro (<http://www.ebi.ac.uk/interpro/>). The results were used to classify the putative two-component proteins by domain organization using a nomenclature adapted from Ohmori et al. (2001), with each group subdivided by the organization of the identified signalling domains. The cartoon style diagrams that present the domain organization of the deduced sequences were constructed from these data, with the sizes of the domains roughly in proportion to each other. For clarity, each gene name begins with a four character acronym (except for *Haloarcula marismortui*, *Pyrococcus abyssi* GE5 and *Pyrococcus horikoshii* OT3, see Table 1) followed by either the locus tag that can be found at IMG or HaloLex (<http://img.jgi.doe.gov/pub/main.cgi> and <http://www.halolex.mpg.de/>) or the gene object identifier if the sequencing or annotation is incomplete.

To determine orthologous groups, orthology information, based on the bidirectional best hits from BLASTPs of each organism against each other organism polypeptide, is accessible at IMG (<http://img.jgi.doe.gov/pub/main.cgi>). This definition is not completely accurate, but it provides a useful approximation as it is not always possible to know whether the polypeptides arose from a single gene present in the last common ancestor (orthologues) or from a gene duplication within a genome (paralogues). Alignments for phylogenetic analysis were performed by TCoffee (Notredame et al. 2000) and accessed at the Centre Nationale de la Recherche Scientifique website (<http://igs-server.cnrs-mrs.fr/Tcoffee/tcoffee.cgi/index.cgi>) and ClustalW alignments (Thompson et al. 1994) were performed at the European Bioinformatics Institute (<http://www.ebi.ac.uk/clustalw/>). Representatives from three bacterial phyla were included in the alignments (chosen by having the best match to one of the archaeal ORFs, either as an orthologue or by BLASTP at IMG). Phylogenetic analysis by neighbor-joining (Bootstrap 250) was performed using MEGA version 3.0 (Kumar et al. 2004) and by maximum-likelihood (Felsenstein 1996) using Molphy, accessed at the Institut Pasteur, biological software website (<http://bioweb.pasteur.fr/intro-uk.html#phylo>).

Closely linked two-component genes and probable operons that contain two or more two-component genes were constructed from the chromosome map images at IMG (<http://img.jgi.doe.gov/pub/main.cgi>), TIGR (<http://www.tigr.org/tdb/>) and the biology of extremophiles website (<http://www-archbac.u-psud.fr/homepage.html>)

Results and discussion

The structural classification of potential two-component proteins is shown in Table 2. No two-component encoding gene could be found in the Crenarchaeota or Nanoarchaeota (data not shown). Sensor domains are drawn as ellipses and two-component (HisKA, HATPase_c and response regulator) and output domains are drawn as rectangles. Parentheses followed by figures indicate the number of similar domains that may be found in the proteins listed in each subclass.

Histidine kinases

Different types of histidine kinases (HK) are listed in Table 2A. Histidine kinases contain two domains; a dimerization and a phosphoacceptor domain (HisKA or HisKA_2) and a HATPase_c domain (Grebe and Stock 1999). HisKA and HisKA_2 are part of a His kinase A phosphoacceptor domain superfamily that also includes HWE_HK and HisKA_3 (Karniol and Vierstra 2004; Pfam accession CL0025).

HKI Histidine kinase Is are HKs containing HisKA and HATPase domains. There may be other domains in some of these examples which are not currently recognized. The HKI ORFs vary greatly in size, ranging from 175 to 592 amino acids in length.

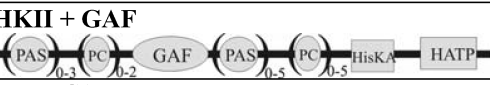
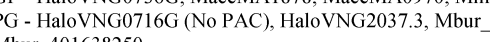
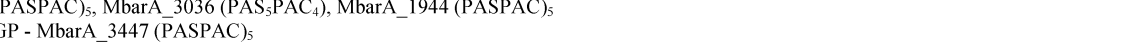
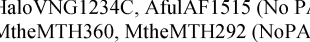
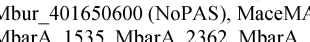
HKII Histidine kinase IIs are HKs containing sensor GAF and PAS/PAC domains. The GAF domains (cGMP phosphodiesterase, adenylyl cyclases, bacterial transcription factors FhlA) are associated with small molecule binding, in particular cAMP and cGMP (Aravind and Ponting 1997, Ho et al. 2000, Anantharaman et al. 2001). The GAF domain is usually found in combination with PAS (*Drosophila* period clock protein, vertebrate aryl hydrocarbon receptor nuclear translocator and *Drosophila* single-minded protein) or PAC (PAS-associated C-terminal motif) domain, or both. One class of PAS domains is known to bind cofactors such as heme and FAD (Bibikov et al 2000, Sardiwal et al. 2005). Sensing of light, oxygen or redox potential by PAS domains requires cofactors, whereas sensing signals such as voltage, xenobiotics and nitrogen availability does not (Ponting and Aravind 1997, Gilles-Gonzalez and Gonzalez 2004). The PAC domains are proposed to contribute to the PAS domain fold. The shared feature of GAF and PAS/PAC domains is the binding of a diverse set of regulatory small molecules that often remain unidentified; all three domains are common signal transduction system components (Anantharaman and Aravind 2001, Zhulin et al. 2003). There is one example containing Cache and one containing SBP_bac_3 (bacterial extracellular solute-binding proteins,

family 3). The Cache domain is a signalling domain found in animal calcium channel subunits and it is thought to form an extracellular or periplasmic ligand sensor (Anantharaman and Aravind 2001). SBP_bac_3 is involved in active transport of

solute across the cytoplasmic membrane and in the initiation of signal transduction pathways (Tam and Saier 1994). This is by far the largest subgroup of ORFs, containing 161 out of the total of 489 (33% of the total).

Table 2. Compilation and cartoon diagrams of all putative archaeal open reading frames encoding two-component proteins. Abbreviations; Cach = Cache; ConA = ConA-like glucanase; Gly = Glycos_transf_2; HKA = His_KA (and KA2); HATP = HATPase_c; Hkd = H-kinase_dim; H10 = HTH_10; PC = PAC; and SBPBac3 = SBP_bac_3. See Table 1 for gene name prefixes.

A. Histidine kinases

HKI	
HaloVNG2180.3, AfulAF1467, AfulAF0208, MtheMTH459, MtheMTH1124, MtheMTH123, MtheMTH444, MtheMTH356, Mburi_401648670, Mburi_401649920, Mburi_401657500, Mburi_401643290, MaceMA3962, MbarA_0679, MmazMM0948, HmarrnAC0477, HmarrnAC1693, HmarrnAC2945, HmarrnAC2997, HmarrnAC3370, HmarrnAC3440, Mhun_401789380, Mhun_401789570 (HisKA_2), Mhun_401795600	
HKII	
HKII + GAF	
HmarrnAC0247, HmarrnAC0964, NphaNP0730A	
HKII + GAF1PASPAC	
GP - HaloVNG0736G, MaceMA1878, MaceMA0970, MmazMM2886, MbarA_2594, HmarrnAC0413, HmarrnB0295. PG - HaloVNG0716G (No PAC), HaloVNG2037.3, Mburi_401647970 (No PAS), Mburi_401650390 (No PAS), Mburi_401653070, Mburi_401638250.	
HKII + GAF2PASPAC	
GP - Mburi_401656610 (No PAC), MaceMA2890, MaceMA2294, MaceMA2266 (PAC1), MbarA_3538, MbarA_0815 (PAS1) PG - HaloVNG1175G (No PAC), HmarrnAC0410 PGP - HaloVNG0916G (PAC), Mburi_401642670 (no PAC), NphaNP1882A (PAC1)	
HKII + GAF3PASPAC	
GP - MaceMA1628, MmazMM0518, NphaNP0090A (PAC1) PGP - AfulAF1483 (PAC), GPGP - NphaNP1912A	
HKII + GAF4PASPAC	
GP - MaceMA3370 PGP - Mburi_2935, MmazMM3295, NphaNP1622A (PAC3)	
HKII + mPASPAC - PGP	
MaceMA1270 (PACPAS), MaceMA1630 (PAC5PAC2), MaceMA0551 (PAS6PAC5), MaceMA1646 (PAS5PAC4), MaceMA0203 (PASPAC)5, MbarA_3036 (PAS5PAC4), MbarA_1944 (PASPAC)5 GP - Mburi_3447 (PASPAC)5	
HKII + mGAFPASPAC	
(PAS)2-G-PASPAC-G-(PASPAC)4 - MmazMM0168 G-(PASPAC)2-G-(PAS)-(PAC)8 - HmarrnB0180 PASPAC-G-(PASPC)4-G - Mburi_3037 (PAS)2-PAC-G-(PAS)7-(PAC)6-(GAF)2 - HmarrnB0299	
HKII + PASPAC	
HaloVNG1234C, AfulAF1515 (No PAC), AfulAF0450, AfulAF1639, AfulAF0021 (No PAC), MtheMTH1260 (NoPAC), MtheMTH360, MtheMTH292 (NoPAC), Mburi_401644730, Mburi_401643030, Mburi_401657970, Mburi_401641290, Mburi_401650600 (NoPAS), MaceMA2757, MaceMA1957, MaceMA0619, MaceMA1704, MaceMA0620 (NoPAS), MaceMA3481, Mburi_1535, Mburi_2362, Mburi_1687 (No PAC), Mburi_1668 (No PAC), MmazMM1781, MmazMM1915, MmazMM1777 (No PAC), MmazMM2990, HmarrnAC0789 (no PAC), HmarrnAC1114 (No PAC), HmarrnAC1391, HmarrnAC1486 (No PAC), HmarrnAC2219 (No PAS), HmarrnAC2461, HmarrnAC2477, HmarrnAC2789 (No PAC), HmarrnAC3347, HmarrnAC3349 (No PAC), HmarrnAC3482, HmarrnB0133(No PAC), NphaNP1506A, NphaNP1640A (No PAC), NphaNP1804A, NphaNP1850A (No PAC), NphaNP3536A (No PAC), NphaNP3752A.	
HKII + 1 PASPAC	
AfulAF0770, AfulAF1184, AfulAF1452, MtheMTH619 (No PAC), MtheMTH823, MtheMTH468 (No PAC), Mburi_401638920 (PAS), Mburi_401653820, MaceMA4026, MaceMA2294, MaceMA2082, MaceMA2732, Mburi_2184, Mburi_2907, Mburi_1483 (1PAS), HmarrnAC0991, HmarrnAC1096, NphaNP0144A, NphaNP1154A, NphaNP1914A, Mhun_401786790 (1PAS)	
HKII + 2 PASPAC	
HaloVNG1374G (PAS)2, AF2109, MtheMTH174 (1PAC), Mburi_401643080 (PAC)2, Mburi_401655300, MaceMA1627 (2PAC), MaceMA0758 (2PAS), MaceMA3346, MaceMA1149, MaceMA0490, MaceMA3543, Mburi_2691 (1PAC), MmazMM1671, MmazMM2748, MmazMM2178 (2PAC), MmazMM2646 (2PAS), HmarrnAC0086, HmarrnAC1501 (2PAS), NphaNP0862A, NphaNP6080A (PAC)2.	
HKII + 3 PASPAC	

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Table 2. Cont'd. Compilation of all putative archaeal open reading frames encoding two-component proteins.

HKII + 4 PASPAC	AfulAF0410 (PAC) ₃ , MaceMA1991 (3PAS), MaceMA1645, MbarA_1666, MbarA_2680, MbarA_2876 (PAS ₃), MbarA_2552, MmazMM2276, MmazMM2773, MmazMM1093, MmazMM0169 (no PAS), MmazMM2435 (3PAC), MmazMM2275 (3PAS).
HKII + 5 PASPAC	MaceMA2784 (PAS) ₄ , MaceMA0759 (PAS) ₄ , MaceMA0552, MbarA_0404 (PAS) ₃ , MmazMM2515, MmazMM2277, MbarA_3250, NphaNP4606A (PAC) ₄ .
HKII + mPASPAC	MaceMA1844 (7PAC, 8PAS), MaceMA1274 (6PAS, 7PAC), MaceMA1322 (6PAS, 7PAC), MaceMA1470 (5PAC, 6PAS), MaceMA3368 (PACPAS) ₇ ,
HKII + Cache Mbur_401646300	
HKII + SBP Mbur_401644740	
HKIII	
HKIII	MmarMMP1303, MmarMMP1120, HmarrnAC0988, NphaNP0142A
HKIII + PASPAC ConA-like glucanase Mbur_401641830	
HKIII + GAFPASPAC HmarrnB0156	
HKIII + CHASE4 AfulAF1721, MaceMA2553, MmazMM3099.	
HKIII + CHASE4PASPAC MaceMA1739 (No PAC), MaceMA2555, MaceMA2348 (No PAC), MmazMM2629, MmazMM3101.	
HKIII + Cache Mbur_401655290, Mbur_401649130, MmazMM2955 (PASPAC), Mhun_401789550	
HKVI	
HKVI	
No P2 - MmazMM1325., MaceMA0014, HmarrnAC2205, MbarA_0984, Mhun_401776240, Mhun_401784470 P2 HaloVNG0971G, AfulAF1040, MaceMA3066, MmazMM0328, Mhun_401793120, NphaNP2172A (P2)₂ - Mbur_401647520, MmarMMP0927, PAB1332, PH0484, Tkod_610170420/30	
HATPase_c	
HATPase_c	Mbur_401643110, Mbur_401662170, Mbur_401641680, Mbur_401642700, Mbur_401657760, MbarA_1241, HmarrnAC0457, HmarrnAC2337.
HATPase_c + PASPAC	AfulAF0893, HmarrnAC0130, NphaNP2516A (PASPAC) ₂ , NphaNP6064A (PAS) ₃ (PAC) ₂ .
HATPase_c + (GAF)₂(PASPAC)_n	MaceMA4561 (PAS) ₂ (PAC) ₃ , MaceMA0863 PAS(PAC) ₂
HATPase_c + PAS	MaceMA0777, MmazMM1931, HmarrnAC1346, HmarrnAC2616 (PAS) ₂ , HmarrnAC2694, NphaNP3356A, Mhun_401806820.
HATPase_c + HAMP	HaloVNG1375.3
His_KA	
His_KA Mhun_401787170	
His_KA (PAC)₁₋₂(PAS)₂₋₃(GAF)₁₋₂ AfulAF0277 (PAC) ₂ (PAS) ₃ -GAF, AfulAF2420_332 (PASPAC) ₂ GAF. AfulAF0448 GAF(PAS)-PACGAF.	
His_KA + (PAS)₀₋₄(PAC)₀₋₃ AfulAF1035 (PAS) ₂ , AfulAF2032 (PAS) ₄ (PAC) ₃ .	

B. Response regulators

RRI-CheY	
HaloVNG0735G, HaloVNG2036G, HaloVNG0974G, HaloVNG0917G (213aa), AfulAF1063, AfulAF1898, AfulAF0449, AfulAF2249, AfulAF1256, AfulAF2419, AfulAF1384, AfulAF1473, AfulAF1042, MtheMTH549, MtheMTH445, MtheMTH447 (277aa), MmarMMP1304, MmarMMP0933, Mbur_401656780 (361aa), Mbur_401641730, Mbur_401640890, Mbur_401660760,	

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Table 2. Cont'd. Compilation of all putative archaeal open reading frames encoding two-component proteins.

Mbur_401647540, Mbur_401657490, Mbur_401644620, Mbur_401658280 (287aa), Mbur_401641840, MaceMA4671, MaceMA1268, MaceMA1269, MaceMA2861, MaceMA3068, MaceMA2445 (292aa), MaceMA1468, MaceMA1469, MaceMA0016, MaceMA0018, MaceMA2012, MaceMA4376, MaceMA1366, MbarA_3448, MbarA_2388, MbarA_3321, MbarA_2896, MbarA_0988, MbarA_0986, MbarA_3109 (260aa), MbarA_1051, MbarA_3248, MbarA_2510, MmazMM0330, MmazMM0049, MmazMM007 (292aa), MmazMM3206, MmazMM1068, MmazMM2351, MmazMM2516, MmazMM2880, MmazMM2953, MmazMM2954, MmazMM1327, MmazMM1328, PAB1330, PH0482, HmarpNG4019, HmarrnAC0361, HmarrnAC0411, HmarrnAC0536 (201aa), HmarrnAC1118, HmarrnAC1494, HmarrnAC2168, HmarrnAC2194, HmarrnAC3308, HmarrnB0297, Tkod_610170400, NphaNP0028A, NphaNP0140A, NphaNP0516A, NphaNP0682A, NphaNP2102A, NphaNP2906A, Mhun_401793100, Mhun_401778650, Mhun_401785170, Mhun_401776210 (284aa), Mhun_401789540, Mhun_401778900, Mhun_401780830, Mhun_401781980, Mhun_401783330, Mhun_401783500, Mhun_401783520, Mhun_401785150, Mhun_401786800, Mhun_401789100, Mhun_401789600, Mhun_401791380, Mhun_401792540, Mhun_401793270, Mhun_401793820 (220aa), Mhun_401794450, Mhun_401795290 (216aa), Mhun_401795560, Mhun_401796600, Mhun_401799230 (214aa), Mhun_401799550, Mhun_401799610 (211aa), Mhun_401799660 (208), Mhun_401799870, Mhun_401800570 (352aa), Mhun_401801850, Mhun_401801900, Mhun_401801910, Mhun_401803600, Mhun_401803780, Mhun_401804010, Mhun_401806750 (238aa).	RRIII-(PACPAS)GAFHTH_10 HmapNG7159 – (PASPAC) ₂ (GAF) ₂ , HmapNG7223 – (GAF) ₂	
RRIII-DUF24 HmarrnB0301		
RRIV-CheB 		
HaloVNG0973G, AfuLF1041, MmarMMP0926, Mbur_401647530, MaceMA3067, MaceMA0015, MbarA_0985, MmazMM0329, MmazMM1326, PAB1331, PH0483, HmarrnAC2204, Tkod_610170410, NphaNP2174A, Mhun_401793110, Mhun_401776230		
RRIV-Glycos_transf_2 MtheMTH548		
RRIV-(PAC)₀₋₅(PAS)₀₋₄(GAF)₀₋₁ MtheMTH1607, HmarrnAC3271, NphaNP1846A, NphaNP2716A –RR-GAF MtheMTH1764 – RR-PAC-GAF HmarrnAC2211 – RR-GAF-PAS Mhun_401786010 – RR-GAF-PAS ₂ MtheMTH440, HmarrnAC0339, HmarrnAC0674, HmarrnAC2109, Mhun_401793830, Mhun_401773960, Mhun_401775740, Mhun_401792240 – RR-PAS Mhun_401789530, Mhun_401795570, Mhun_401803130 - RR-PASPAC MtheMTH457 – RR-PAS-PAC-(PAS) ₂ HmarrnAC0356 – RR-(PASPAC) ₂ -GAF HmarrnAC1142 – RR-PAC-PAS-PAC Mhun_401776490 – RR-(PAS) ₃ (PAC) ₂ Mhun_401796390 RR-(PAS) ₅ (PAC) ₄ .		
C. Hybrid kinases		
HY1 		
MtheMTH901, HmarrnAC0301, HmarrnAC0412, HmarrnAC0475, HmarrnB0296, NphaNP3458A, Mhun_401776220, Mhun_401784480, Mhun_401792550, Mhun_401801890, Mhun_401799920 (No HisKA)		
HY1+PASPAC 		
MtheMTH902 (No PAS), Mbur_401644630, Mbur_401651890, Mbur_401641180, HmarrnAC0487 (No PAC), NphaNP0138A (No PAS), Mhun_401789580 (No HATPase), Mhun_401796780, Mhun_401800550 (HisKA_2), Mhun_401806740 (HisKA_2)		
HY1+(PASPAC)₂ MtheMTH446 (No PAC), MaceMA1267, Mbur_401642690, HmarrnAC0574, HmarrnAC0848 (PAC) ₁ , HmarrnAC1806 (PAS) ₁ , NphaNP0188A, NphaNP0622A, Mhun_401803140 (PAC ₁ , No HATPase), Mhun_401795550 (No HATPase), Mhun_401785160, Mhun_401795880, Mhun_401803320 (No PAC, HATPase), Mhun_401803800 (No PAC), Mhun_401803980		
HY1 + mPASPAC HaloVNG5037G – (PAS) ₅ (PAC) ₃ , MbarA_3247 - (PAS) ₂ (PAC) ₃ , MmazMM2518, HmarrnAC2044, HmarrnAC3050, Mhun_401801140, Mhun_401805200 - (PAS) ₃ (PAC) ₂ , HmarrnAC0075, NphaNP6028A, NphaNP6202A, Mhun_401774520, Mhun_401777310, Mhun_401801130, Mhun_401801880 (HisKA_2) – (PACPAS) ₃ , Mhun_401777350, - (PAS) ₄ (PAC) ₂ , HmarrnAC1626, NphaNP2742A – (PASPAC) ₄ , Mhun_401774510 (PASPAC) ₅ , Mbur_401644900 (PAS) ₅ (PAC) ₄ (No HATPase), Mhun_401795210 (PASPAC) ₆ , Mhun_401774210 (6PAS, 4PAC), NphaNP2656A (7PAC, 8PAS), Mhun_401790080 (PAS) ₄ ,		
HY1 + GAF HmarrnAC0794		

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Table 2. Cont'd. Compilation of all putative archaeal open reading frames encoding two-component proteins.

HYI + PASPACGAF	
	
HmarrnAC2372, HmarrnAC3379, NphaNP5274A - PAS-GAF NphaNP5120A - PASPAC-GAF HmapNG7155, HmarrnAC2533, HmarrnAC2692 - GAF-PAS-PAC HmapNG7156 - (PAS) ₂ -PAC-GAF-(PACPAS) ₂ NphaNP0626A - PASPAC-GAF-(PACPAS) ₂ -GAF HmarrnAC2416 - PASPAC-GAF-(PAS) ₂ (PAC) ₃ HmarrnAC3361, Mhun_401786010 - PASPAC-GAF-(PASPAC) ₃ HmarrnAC1495 - (PAS) ₂ -PAC-GAF-PASPAC-GAF-(PASPAC) ₂ HaloVNG2334 - (PAS) ₂ -PAC-GAF-(PAC) ₃ (PAS) ₂ -GAF Mhun_401793840 - GAF-(PASPAC) ₃ Mhun_401775750 - GAF-PASPAC-HisKA-(PAS) ₄ (PAC) ₂ Mhun_401779810 - (PAS) ₃ (PAC) ₂ -HisKA-(PAS) ₂ PAC-GAF-(PAS) ₄ (PAC) ₃ NphaNP4696A - RR-PASPAC-HisKA-(PAS) ₄ (PAC) ₃	
HYII+Hpt	
MaceMA2013	
HYIII	
PAC(PAS) ₂ - MaceMA2256. (PACPAS) ₂ - MmazMM3205, MmazMM2881	
HYIII+CHASE4HAMP	
MaceMA4377, MbarA_1052	
HYIHisKA	
AfulAF1472	

HKIII Histidine kinase IIIIs are HKs that possess a HAMP “linker” (histidine kinase, adenylyl cyclase, methyl-accepting chemotaxis protein and phosphatase) domain. The HAMP domain is usually associated with the transmission of a signal across a membrane from periplasmic ligand-binding domains (Aravind and Ponting 1999, Appleman and Stewart 2003, Zhu and Inouye 2004). Eight examples of HKIIIs have an N-terminal putative periplasmic signalling CHASE4 domain and four have an N-terminal periplasmic signalling Cache domain (Anantharaman and Aravind 2001, Zhulin et al. 2003). These domains are positioned next to the HAMP domain, presumably for efficient transfer of the signal.

HKVI Histidine kinase IVs are the CheA-like chemotaxis signalling proteins that contain an N-terminal Hpt (histidine phosphotransfer) and Hkd (histidine kinase dimerization) domain and a C-terminal CheW domain. Some contain one or two P2 domains between the Hpt and Hkd. The Hpt domain is involved in mediating phosphotransfer from one receiver domain to another (Hoch 2000). Hkd (H-kinase-dim) is the dimerization domain of CheA and CheW that interacts with methyl-accepting chemotaxis proteins (MCPs), relaying signals to CheY, and thereby affecting flagellar rotation (West et al. 1995). The P2 domain is involved in enhancing the interaction of CheY with the HK (Jahreis et al. 2004, Stewart and van Bruggen 2004). *Thermococcus kodakaraensis* has two open reading frames with a frame shift mutation that probably encodes for a CheA-like protein. All of the HKVI genes discussed are located close to other genes that could be involved with signal transduction and are probably transcribed as single operons (see Table A2).

HATPase_c These contain no dimerization or phosphoac-

ceptor domains currently recognized at INTERPRO.

His_KA There are five groupings that contain His_KA without a discernable HATPase_c domain.

Response regulators

Response regulators (RR) are listed in Table 2B. These contain a characteristic receiver (RR/T_{reg}) domain, which is about 120 amino acids long and contains a conserved aspartate residue about halfway along the molecule that accepts a phosphate group from an HK.

RR I Response regulator Is are simple orphan (no other domain detected) RRs, representing the second largest group of two-component ORFs (24% of the total).

RR III Response regulator IIs contain an RR fused to a potential DNA binding domain. Such regulators are found only in *H. marismortui*. Of these, there are only three examples that contain either the HTH_10 or DUF24 domain (PF04967 and PF01638). These are the only RRs that are possibly transcriptional regulators, but there may be other currently unidentified DNA-binding domains in other RRs or hybrid kinases.

RR IV Response regulator IVs contain an N-terminal RR fused to output or signal domains. There are 16 examples of CheB fused to the RR. The CheB domain is related to methyl-esterase and is likely to be concerned with chemotaxis (West et al. 1995). There is one example of two RRs fused to a glycosyl transferase domain in *M. thermoautotrophicus* (Pfam Accession number: PF00353). The glycosyl transferase domain is involved in transferring sugar moieties from a donor to recipient molecules. There are a lot of *Methanospirillum hungatei* ORFs fused with PAS/PAC or GAF domains, or both, however, as the

annotation is incomplete, some of these ORFs may turn out to be part of hybrid kinases.

Hybrid kinases

Hybrid kinases (HY) are shown in Table 2C. They are defined as containing both HK and RR domains. The nomenclature is based on the position and number of RR with respect to the HK. There is an incomplete HYI in *A. fulgidus* that has a PAC/PAS and GAF sensor domain, but no discernable HATPase domain.

HYI Hybrid kinase Is have a single RR N-terminal to the HK.

HYII Hybrid kinase IIs have a single RR C-terminal to the HK. There is only one example in *M. acetivorans*.

HYIII Hybrid kinase IIIs have two RRs either N or C-terminal to the HK.

Distribution of putative two-component ORFs

The total number of ORFs within each class of two-component proteins, for each species of Euryarchaeota, is shown in Table 3. No two-component ORFs were found in the four Crenarchaeota species or *N. equitans* (data not shown). No two-component ORF was found in *M. jannaschii*, *M. kandleri*, *P. furiosus* or in any of the members of the Thermoplasmatales. The three other *Pyrococcus* species each have only three two-component ORFs (*Thermococcus kodakaraensis* HKVI that has a frame shift Tkod_61070420/30, that could be a sequencing error has been counted as one), representing 0.17% of the protein-coding capacity of the genome. *Methanococcus maripaludis* and *Halobacterium* also have a small number, six (0.34%) and 16 (0.64%), respectively. *Archaeoglobus fulgidus*, *M. thermoautotrophicus* and the four Methanosarcinales groups have a comparatively large number of two-com-

ponent ORFs, from 23 to 67. This represents from 1.03 to 1.48% of the coding capacity of the four complete genome assignments. *Haloarcula marismortui* has the largest number of two-component encoding genes, of the complete annotations, which represents 1.93% of the total protein coding capacity of the genome. *Methanospirillum hungatei* appears to have the largest number of two-component genes at 87, though the annotation of the genome is incomplete (so no percentage is given in Table 3). The HKs form half to two-thirds of the two-component ORFs for each species (except *Pyrococcus* sp. and *Methanospirillum hungatei*). The DNA-binding domains (putative) were only detected as part of the RRs in *H. marismortui*.

The PAS/PAC and GAF sensory domains are found in 293 of the 489 putative proteins surveyed. These sensory domains are absent in the *Pyrococcus* sp. A total of 18 ORFs were found that contain the HAMP domain that would in most cases be involved in transferring signals from sensor domains detecting information outside the cell.

Orthologous groups

Potential orthologous groups are shown in Table 4. These results are based on the bidirectional best hits from BLASTPs at IMG. The identification of orthologous groups at IMG may not be correct in all cases as some groupings may include ORFs that are due to gene duplication, hence a parologue (in a different organism) rather than an orthologue. It is, nevertheless, a useful tool for assigning putative orthologous groups when no functional information is available. The groups have been named with a three letter acronym for ease of reference (see Table 4). In *arr18/19*, RRIV-CheB, the grouping was modified from the information at IMG based on the phylogenetic analysis presented in Figure 1 (see below). There are many orthologous groups that contain two or three members, partic-

Table 3. Distribution of euryarchaeal two-component open reading frames. The total number of identified two-component genes are shown (Total 2-C) and are given as a percentage of the total protein coding capacity of each genome (% 2-C). Abbreviations: HK = histidine kinase; RR = response regulator; HY = hybrid kinase; and incom = incomplete.

Species	HKI	HKII	HKIII	HKVI	HK other	HK total	HK	RRI	RRIII	RRIV total	RR	HY	Total 2-C	% 2-C
<i>Archaeoglobus fulgidus</i>	2	10	1	1	6	20	9	0	1	10	1	31	31	1.26
<i>Haloarcula marismortui</i>	6	23	2	2	6	39	10	3	8	21	22	82	82	1.93
<i>Halobacterium</i> sp. NRC-1	1	7	0	1	1	10	4	0	1	5	2	17	17	0.64
<i>M. thermoautotrophicus</i>	5	7	0	0	0	12	3	0	5	8	3	23	23	1.2
<i>Methanococcus maripaludis</i>	0	0	2	1	0	3	2	0	1	3	0	6	6	0.34
<i>Methanococcoides burtonii</i>	4	17	3	1	5	30	9	0	1	10	5	45	45	incomp
<i>Methanosarcina acetivorans</i>	1	38	4	2	3	48	13	0	2	15	4	67	67	1.48
<i>Methanosarcina barkeri</i>	1	21	0	1	1	24	10	0	1	11	2	35	35	1.03
<i>Methanosarcina mazei</i>	1	20	4	2	1	28	12	0	2	14	3	45	45	1.33
<i>Methanospirillum hungatei</i>	3	1	1	3	1	9	36	0	12	48	30	87	87	incomp
<i>Natronomonas pharaonis</i>	0	17	1	1	3	22	6	0	3	9	11	42	42	incomp
<i>Pyrococcus abyssi</i>	0	0	0	1	0	1	1	0	1	2	0	3	3	0.17
<i>Pyrococcus horikoshii</i>	0	0	0	1	0	1	1	0	1	2	0	3	3	0.14
<i>Thermococcus kodakarensis</i>	0	0	0	1	0	0	1	0	1	2	0	3	3	0.13
												489		

Table 4. Potential orthologous two-component open reading frames.

Gene name	Classification	Orthologue number
MaceMA3962, MbarA_0679, MMAZMM0948	HKI	<i>ahk1</i>
AfulAF208	HKI	<i>ahk2</i>
Mbur_401643110	HATPase	
MaceMA1878, MbarA_3037	HKII	<i>ahk3</i>
MaceMA1628, MMAZMM0518	HKII	<i>ahk4</i>
MaceMA2890, MbarA_2935, MmazMM3295, MtheMTH174	HKII	<i>ahk5</i>
Mhun_401800550	HYI PASPAC	
MaceMA1646, MbarA_3036	HKII	<i>ahk6</i>
MaceMA0203, MbarA_1944	HKII	<i>ahk7</i>
MmazMM2515, MbarA_3447, MaceMA1470	HKII	<i>ahk8</i>
MaceMA0552, MmazMM0168	HKII	<i>ahk9</i>
MaceMA2294, Mbur_401657970	HKII	<i>ahk10</i>
MbarA_2362, MmazMM2990	HKII	<i>ahk11</i>
MaceMA0619, MMAZMM1777	HKII	<i>ahk12</i>
MaceMA1704, MmazMM2990	HKII	<i>ahk13</i>
MaceMA0620, MmazMM1781, MbarA_1535	HKII	<i>ahk14</i>
MaceMA1149, MmazMM2178	HKII	<i>ahk15</i>
MaceMA0490, MmazMM1671, MbarA_2876	HKII	<i>ahk16</i>
MaceMA1645, MbarA_2680, MmazMM2748, Mhun_401774840	HKII	<i>ahk17</i>
MbarA_1687, MmazMM1931, Mhun_401789380, HmarrnAC0789	HKII	<i>ahk18</i>
HaloVNG1234C, HmarrnAC3482	HKII	<i>ahk19</i>
AfulAF0770, MmazMM2518, MaceMA3405	HKII	<i>ahk20</i>
MaceMA4026, MmazMM0889	HKII	<i>ahk21</i>
MaceMA2294, Mbur_401657970, HaloVNG1374G, Mhun_401803800, AfulAF0450, NphaNP1154A	HKII	<i>ahk22</i>
HmarrnB0156	HKIII	
AfulAF2109, Mhun_401800710, HmarrnAC1626	HKII	<i>ahk23</i>
MaceMA3368, Mbur_401643080, MaceMM2773, MbarA_3250	HKII	<i>ahk24</i>
MaceMA1627, MmazMM0172, MbarA_0404	HKII	<i>ahk25</i>
MaceMA1991, MtheMTH468, MmazMM1915	HKII	<i>ahk26</i>
MbarA_1666, Mhun_401774840	HKII	<i>ahk27</i>
MmazMM2276, MaceMA1274	HKII	<i>ahk28</i>
MmazMM1093, MbarA_3538	HKII	<i>ahk29</i>
MaceMA0759, Mhun_401779810	HKII	<i>ahk30</i>
MaceMA1844, MtheMTH823	HYI	
HmarMMP1303,	HKII	<i>ahk31</i>
Mbur_401650390	HKII	<i>ahk32</i>
(NphaNP1640A)	HKIII	
HmarrnAC2616	HKII	<i>ahk33</i>
(NphaNP1804A)	HATPasePAS	
HmarrnAC0130	HKII	<i>ahk34</i>
HmarrnB0133, NphaNP3536A	HATPasePAS	
MaceMA2553, MmazMM3099	HKII	<i>ahk35</i>
MaceMA2555, MMAZMM3101	HKIII	<i>ahk36</i>
AfulAF1721, Mhun_401794250	HKIII CHASE	<i>ahk37</i>
MaceMA1739, MmazMM2629	HKIII CHASE	<i>ahk38</i>
MaceMA3066, MmazMM0328, AfulAF1040, Mbur_401647520	HKVI	<i>ahk39</i>
MaceMA0014, MmazMM1325, MbarA_0984	HKVI	<i>ahk40</i>
PAB1332, PH0484, Tkod_610170420/30, MmarMMP0927	HKVI	<i>ahk41</i>
HaloVNG0971G, HmarrnAC2205, NphaNP2172A, Mhun_401793120	HKVI	<i>ahk42</i>
MaceMA0777, MmazMM1931	HATPase+	<i>ahk43</i>
Mhun_401789380	HKI	
HmarrnAC0789	HKII	
MaceMA0863, Mbur_401650390	HATPaseGAF	<i>ahk44</i>
	HKII	

continued overleaf

Table 4. Cont'd. Potential orthologous two-component open reading frames.

Gene name	Classification	Orthologue number
HaloVNG2036G, HmarrnAC1118 (NphaNP2906A 50%)	RRI	<i>arr1</i>
MmarMMP1304, Mbur_401640890, Mhun_401785170	RRI	<i>arr2</i>
MaceMA4671, MmazMM2880	RRI	<i>arr3</i>
MbarA_2510, MmazMM2953	RRI	<i>arr4</i>
MmazMM2954, Mbur_401641840	RRI	<i>arr5</i>
MtheMTH447, Mbur_401660760, Mhun_401806750	RRI	<i>arr6</i>
HaloVNG0974G, AfuAF1042, MmarMMP0933, Mbur_401647540, MaceMA3068, MmazMM0330, PAB1330, PH0482, HmarrnAC2194, Tkod_610170400, Mhun_401793100, NphaNP2102A	RRI	<i>arr7</i>
Mbur_401660760, MaceMA1469, MbarA_3448, MMAZMM2516, MtheMTH1764 (PACGAF), Mhun_401793830 (PAS)	RRI	<i>arr8</i>
MaceMA1268, MbarA_3248, HmarrnAC0411, AfuAF1473, MtheMTH445, Mhun_401778650, HmarrnAC0536, (NphaNP0140A)	RRI	<i>arr9</i>
MaceMA0016, MbarA_0986, MmazMM1327	RRI	<i>arr10</i>
MaceMA2861, MbarA_2388, MmazMM0049, HmarrnAC0339 (PAS)	RRI	<i>arr11</i>
MaceMA4376, MbarA_1051, MmazMM1068, AfuAF2419, Mbur_401644620, Mhun_401789540	RRI	<i>arr12</i>
MaceMA1366, MbarA_2896, MMAZMM2351, Mbur_401657490, Mhun_401803780	RRI	<i>arr13</i>
MaceMA2445, MbarA_3109, MMAZMM3007, Mbur_401658280, Mhun_401776210	RRI > 200aa	<i>arr14</i>
HmarrnAC3308, Mhun_401796600	RRI > 200aa	<i>arr15</i>
MaceMA0018, MbarA_0988, MMAZMM1328	RRI	<i>arr16</i>
HmapNG7223, Mhun_401799920	RRIII-HTH	<i>arr17</i>
MaceMA0015, MbarA_0985, MmazMM1326	HYI-HATPase	
PAB1331, PH0483, Tkod_610170410, HaloVNG0973G, HmarrnAC2204, MaceMA3067, MmazMM0329, Mbur_401647530, AfuAF1041, MmarMMP0926, NphaNP2174A, Mhun_401793110	RR CheB	<i>arr18</i>
MtheMTH1607	RR IV	<i>arr20</i>
AfuAF0448	His_KAPACPASGAF	
MtheMTH440,	RRIVPAS	<i>arr21</i>
Mbur_401642690, Mhun_401789580	HYI PASPAC	
HmarrnAC1142	RRIVPAS	<i>arr22</i>
Mhun_401803140	HYI PASPAC	
Mhun_401784480, Mbur_401644630	HYI	<i>ahy1</i>
MaceMA1267, MbarA_3247, MtheMTH446, Mhun_401795550	HYI PASPAC	<i>ahy2</i>
AfuAF1472	HYIHisKA	
HmarrnAC3379,	HYI PASGAF	<i>ahy3</i>
HaloVNG1175G	HKIIPASGAF	
HmarrnAC2533	HYI PASGAF	<i>ahy4</i>
HaloVNG0916G, (NphaNP1912A)	HKIIPASGAF	
HmapNG7156, HaloVNG2334	HYI PASGAF	<i>ahy5</i>
MaceMA2256, MmazMM2881	HYIII	<i>ahy6</i>
MaceMA4377, MbarA_1052	HYIII	<i>ahy7</i>
HmarrnAC0475	HYI	<i>ahy8</i>
Mhun_401787170	HisKA	
Mhun_401801890,	HYI	<i>ahy9</i>
MtheMTH444	HKI	
HmarrnAC2044, Mhun_401801140	HYIPASPAC	<i>ahy10</i>
Mhun_401774520, HaloVNG5037G, Mbur_401644900	HYIPASPAC	<i>ahy11</i>
HmarrnAC3050, Mhun_401777310	HYIPASPAC	<i>ahy12</i>

ularly within the Methanosaecinales. The more interesting groups are those that have more members or that have members in different genera. These will be discussed below, particularly those that are part of taxis operons (*ahk40–43*, *arr7*, *arr10*, *arr18* and *arr19*).

Phylogenetic analysis

Figures 1 to 5 show neighbor-joining and the supplementary Figures A1 to A5 contain the maximum likelihood phylogenetic analyses of alignments from TCoffee analysis. Phylogenetic analysis of the same ORFs was also performed on align-

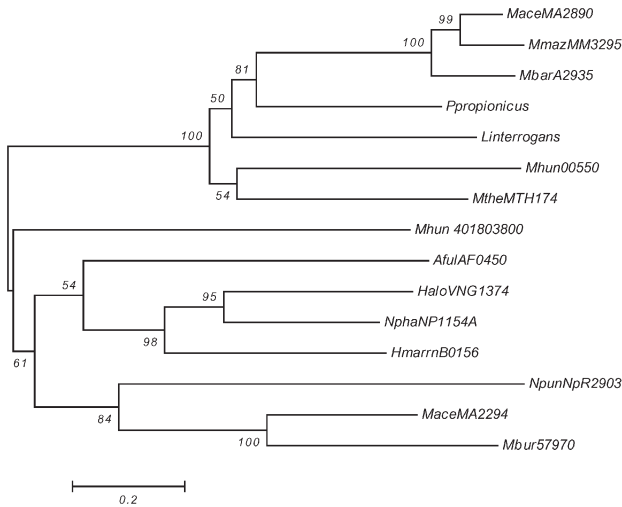


Figure 1. Phylogenetic analysis by neighbor-joining of putative two-component open reading frames (ORFs) from histidine kinase II groups *ahk5* and *ahk22*. Group *ahk5* = MaceMA2890, MbarA_2935, MmazMM3295, MtheMTH174 and Mhun_400550; and group *ahk22* = MaceMA2294, Mburi57970, HaloVNG1374G, Mhun_401803800, AfuAF0450, NphaNP1154A and HmarrnB0156. The bacterial ORFs are Ppropionicus, NpuncNpR2903 and Linterrogans. Abbreviations: Ppropionicus = *P. propionicus*_500413890; NpuncNpR2903 = *N. punctiforme*; and Linterrogans = *L. interrogans*-LA2540

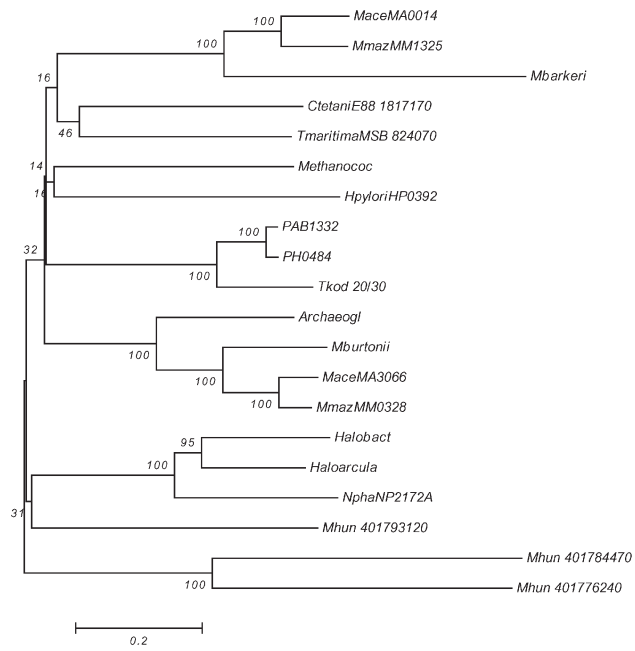


Figure 2. Phylogenetic analysis by neighbor-joining of putative two-component open reading frames (ORFs) from the histidine kinase IV groups *ahk40-43*. Group *ahk40* = MaceMA3066, MmazMM0328, AfuAF1040 and Mburi58280; group *ahk41* = MaceMA0014, MmazMM1325 and MbarA_0984; group *ahk42* = PAB1332, PH0484, Tkod_610170420/30 and MmarMMP0927; and group *ahk43* = HaloVNG0971G, HmarrnAC2205, NphaNP2172A and Mhun_401793120. The bacterial ORFs are CtetaniE17170, TmaritimMSB8_824070 and HpyloriHP0392. Abbreviations: CtetaniE17170 = *C. tetani*E88_1817170; TmaritimMSB8_824070 = *T. maritima*; and HpyloriHP0392 = *H. pylori*.

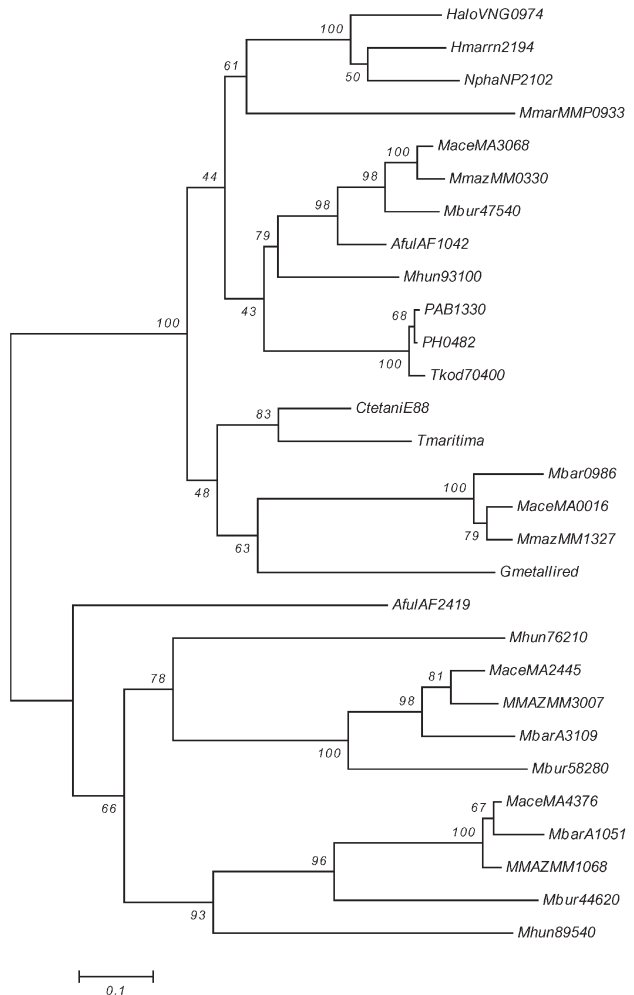


Figure 3. Phylogenetic analysis by neighbor-joining of putative two-component open reading frames (ORFs) from the response regulator I orphans *arr7*, *arr10*, *arr12* and *arr14*. Group *arr7* = HaloVNG0974G, AfuAF1042, MmarMMP0933, Mburi47540, MaceMA3068, MmazMM0330, PAB1330, PH0482, HmarrnAC2194, Tkod_610170400, Mhun_401793100 and NphaNP2102A; group *arr10* = MaceMA0016, MbarA_0986 and MmazMM1327; group *arr12* = MaceMA4376, MburiA_1051, MmazMM1068, AfuAF2419, Mburi44620 and Mhun_401789540; and group *arr14* = MaceMA2445, MburiA_3109, MMAZMM3007, Mburi401658280 and Mhun_401776210. The bacterial ORFs are CtetaniE88, Tmaritim and Gmetallired. Abbreviations: CtetaniE88 = *C. tetani*E88_1817-150; Tmaritim = *T. maritima*MSB8_24050; and Gmetallired = *G. metallireducens*_401349760.

ments made by ClustalW (data not shown), but the results were not found to differ significantly. Figure 1 contains the ORFs from the two HKII groups, *ahk5* and *ahk22*, with the three closest bacterial ORFs (to MaceMA2890) from Cyanobacteria, Firmicutes and Proteobacteria. Figure 2 is composed of ORFs from the four HKVI ‘CheA like’ groups, *ahk40-43*, and the three closest bacterial ORFs (to MaceMA0014) from Thematogae, Firmicutes and Proteobacteria. Figure 3 is the analysis of a number of RRI orphans, *arr7* and *arr10* (from putative taxis operons), *arr12* and *arr14* (> 200 amino acids)

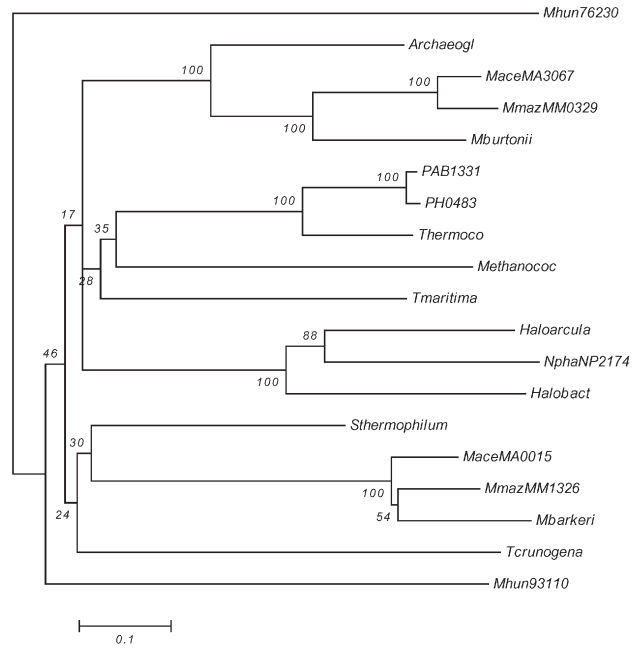


Figure 4. Phylogenetic analysis by neighbor-joining of putative two-component open reading frames (ORFs) from response regulator IV group *arr18* and *arr19*. Group *arr18* = MaceMA0015, MbarA_0985 and MmazMM1326; and group *arr19* = PAB1331, PH0483, Tkod_610170410, HaloVNG0973G, HmarrnAC2204, MaceMA3067, MmazMM0329, Mburr_401647530, AfuAF1041, MmarMMP0926, NphaNP2174A and Mhun_401793110. The bacterial ORFs are Sthermophilum, Tmaritima and Tcrunogena. Abbreviations: Thermoco = *T. kodakaraensis*; Sthermophilum = *S. thermophilum*_3768360; Tmaritima = *T. maritima*MSB8_21060; and Tcrunogena = *T. crunogena* Tcr0758

with bacterial ORFs from Thematogae, Firmicutes and Proteobacteria (closest to MaceMA3068). Figure 4 shows the results for the two RRIV CheB orthologous groups from taxis operons, *arr18* and *arr19* with bacterial representatives from Thematogae, Proteobacteria and Actinobacteria (closest to MaceMA0015). Figure 5 shows results for *ahy2* and bacterial representatives from Cyanobacteria, Actinobacteria and Proteobacteria.

Linked genes

Genes that are located close to each other on the genome and transcribed in the same orientation are shown in Table A2. Most of these are likely to be part of operons. This provides clues to some cognate pairs of HKs, HYs and RRs. All putative HKVI encoding genes are located with other “chemotaxis” genes in “chemotaxis operons,” including two such operons for *M. acetivorans* and *M. mazei*. Included in these “chemotaxis operons” are the orthologous groups, *ahk40–43*, *arr7*, *arr10* and *arr18* and *arr19* (see Table 4).

Conclusions

Distribution of two-component ORFs

Ten species, representing four genera have at least 17 putative

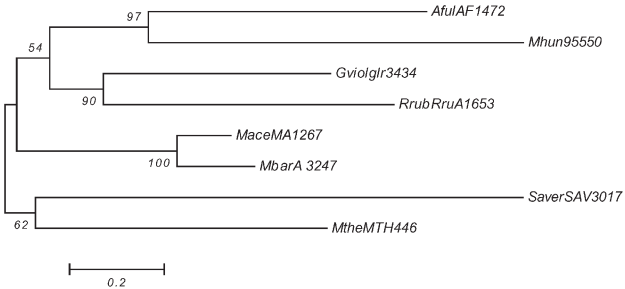


Figure 5. Phylogenetic analysis by neighbor-joining of putative two-component open reading frames (ORFs) from hybrid kinase 1 group *ahy2*. The bacterial ORFs are Gviolglr3434, RrubRruA1653 and SaverSAV3017. Abbreviations: Gviolglr3434 = *G. violaceus*; RrubRruA1653 = *R. rubrum*; and SaverSAV3017 = *S. avermitilis*.

two-component ORFs. Some of these two-component ORFs are quite sophisticated in structure, including the multiple sensor HKIIs, CheA-like HKVIs and the hybrid kinases. The results presented here show that a number of euryarchaeal species have an extensive array of two-component sensory ORFs. These proteins may sense a number of different internal signals by means of PAS/PAC domains and their associated co-factors (Ponting and Aravind 1997, Gilles-Gonzalez and Gonzalez 2004). In addition, the potential to sense other small molecules (particularl cNMPs) via the GAF domains (Aravind and Ponting 1997, Ho et al. 2000, Anantharaman et al. 2001) and extracellular signals, by the CHASE4 and Cache putative sensory domains, via HAMP domains (Anantharaman and Aravind 2001, Zhulin et al. 2003) shows that these organisms (in particular *H. marismortui*, *Natronomonas pharaonis*, *Methanospirillum hungatei* and the Methanosarcinales) possess sophisticated and complex sensory networks. As yet, none of these putative two-component genes have a functional name, so functions can be assigned only by similarity. The DNA-binding RRs are common in bacteria that regulate gene expression (Ashby 2004, Galperin 2005). However only three RRs have been identified with putative DNA binding domains, all in *H. marismortui*. If regular indiscriminate HGT were taking place, one would expect to see more DNA-binding RRs in archaeal sequences. Presumably the large number of orphan RRs are involved in regulation of cellular activity by interacting directly with other proteins. Transcriptional control is probably maintained by the many DNA-binding domains that have been identified as part of one-component systems in archaea (Ulrich et al. 2005). In these systems the DNA-binding output domain is linked directly to a sensor domain without any phosphotransfer.

Of the species that have the most two-component genes, *H. marismortui* and *Natronomonas pharaonis* are halophilic and the Methanosarcinales and *Methanospirillum hungatei* are mesophiles. The mesophiles coexist with a large and diverse population of bacteria, giving ample opportunity for HGT, whereas the opportunity for HGT in the halophilic organisms would be more restricted. This begs the question of how the distribution of two-component genes that can be seen in the

Euryarchaeota arose. Was it through HGT exclusively or by vertical transfer from a common ancestral euryarchaeal organism coupled with gene duplications?

Phylogeny and inheritance of two-component ORFs

The phylogenetic analysis of five different sets of orthologous ORFs, chosen because they are found in most of the species that contain two-component ORFs (Figures 1–5), were found to closely match the published phylogenies for these organisms (Matte-Tailliez et al. 2002, Brochier et al. 2004, Baptiste et al. 2005).

For *ahk5* and *ahk22*, shown in Figure 1, the phylogeny of each group agrees with the current phylogeny of these organisms and the position of the three bacterial examples indicates that the two groups may have arisen through a separate HGT event in an ancestral euryarchaeal species for *ahk5* and possibly, into an ancestral methanogen for *ahk22*.

The results for the CheA-like HKVI ORFs are shown in Figure 2. *Ahk40*, *ahk42* and *ahk43* (except Mhun_401793120) cluster together and probably represent vertical inheritance from a single HGT event into an ancestral Euryarchaeota species (one bacterial ORF from *T. maritima* giving the best match). *Ahk41* appears to be a separate group, found in the Methanosarcinales, that clusters on its own and seems to be more closely associated with the Firmicutes and Proteobacterial examples, presumably representing a separate HGT event. The three *Methanospirillum hungatei* ORFs seem to be due to separate (Mhun_401793120 probably should not be in *ahk43*) HGT events and Mhun_401784470 and Mhun_401776240 are probably true paralogues.

Figure 3 shows the results for four orphan RR groups. The two groups, associated with putative taxis operons *ahk7* and *ahk10*, group closely together, however, *ahk10*, which is found only in the Methanosarcinales is probably due to an HGT event into a direct ancestor of this group. The other two orthologous groups, *arr12* and *arr14* are quite separate from the first two mentioned groups (*arr7* and *arr10*) and probably arose from separate HGT events into the ancestors of methanogens (*AfulAF2419* appears to be a distant member of *ahk12*).

Figure 4 shows the results for the two RRIV-CheB orthologous groups associated with taxis. The *arr18* orthologous group found in Methanosarcinales groups separately from *arr19*, being closer to two of the bacterial ORFs. Therefore *arr18* appears to be the result of a separate HGT event in an ancestor of the Methanosarcinales, whereas *arr19* appears to be the result of an HGT event into an ancestor of Euryarchaeota.

The phylogeny for *ahy2* (the biggest hybrid kinase orthologous group), shows that these members probably arose from more than one HGT event. The combined results for the orthologous groups found in potential taxis operons are shown in Table A2.

The operon that contains HKVI (*ahk40/42/43*), RRI (*arr7*) and RRIV-CheB (*arr19*) appears to have arisen as an HGT event that transferred the whole operon into an ancestor of the Euryarchaeota. In contrast, the taxis operon containing HKVI (*ahk41*), RRI (*arr10*) and RRIV-CheB (*arr18*) appears to have

arisen from a separate HGT event of the whole operon into a direct ancestor of the Methanosarcinales.

The results presented here suggest that HGT has taken place from bacterial species both into ancestral Euryarchaeota and more recently into the methanogens. However the large numbers of two-component genes in the mesophilic methanogens and the Halobacterales probably reflect their well known metabolic flexibility (Baptiste et al. 2005, Falb et al. 2005). This in turn, necessitates an increased requirement for regulation of cellular activity in a changing environment rather than the increased potential for HGT from bacteria. Most of the two-component ORFs that can be observed in these groups of organisms are probably derived from paralogous gene duplication events, the number of two-component ORFs observed would be driven by the requirement to control cellular activity as the organisms evolve. A limited number of HGT events could be sufficient to account for the diversity of phosphotransfer and sensory domains.

Any function of two-component ORFs is inferred by homology to known bacterial genes (e.g. HKVI and chemotaxis) and awaits *in situ* or *in vitro* studies, or both. This highlights the importance of interfacing between bioinformaticians and biochemists to plan experiments in an informed way, particularly where orthologues are identified and found in more than one genus and hence may play central roles in cellular regulation.

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Appendix

Table A1 shows the two-component protein domains used for BLASTP. Figures A1 to A5 show maximum likelihood phylogenetic analyses. Table A2 shows closely linked genes that may be part of operons.

Table A1. Two-component protein domains from *M. acetivorans* (*M. ace*) and *Escherichia coli* K12 (*E. coli*) used for BLASTP searches, showing the online accession numbers and the amino acid range that was used. Abbreviation: aa = amino acids.

Domain	Species	Gene	Amino acid region
CheY Receiver	<i>E. coli</i>	NP_416396.1	
	<i>M. ace</i>	MA0016	
	<i>M. ace</i>	MA3068	
OmpR Receiver	<i>E. coli</i>	NP_417864.1	aa 6–124
	<i>E. coli</i>	NP_417863.1	aa 234–439
Histidine kinase	<i>M. ace</i>	MA0490 (HisKA_2)	aa 639–847
	<i>E. coli</i>	NP_415513.1	aa 815–896
Hpt	<i>M. ace</i>	NP_614988	aa 5–106

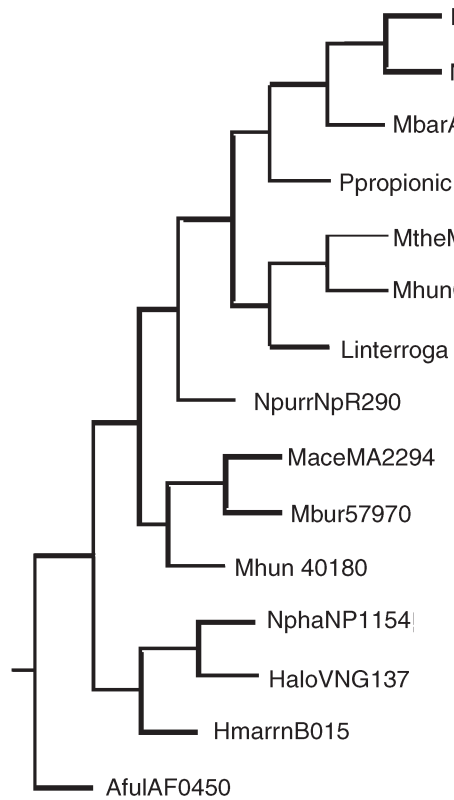


Figure A1. Phylogenetic analysis by neighbor-joining of putative two-component open reading frames (ORFs) from histidine kinase II groups *ahk5* and *ahk22*. Group *ahk5* = MaceMA2890, MbarA_2935, MmazMM3295, MtheMTH174 and Mhun_401800550; and group *ahk22* = MaceMA2294, Mbur_401657970, HaloVNG1374G, Mhun_401803800, AfulAF0450, NphaNP1154A and HmarrnB0156. The bacterial ORFs are Ppropionicus, NpurrNpR2903 and Linterrogans. Abbreviations: Ppropionicus = *P. propionicus*_500413890; NpurrNpR2903 = *N. punctiforme*; and Linterrogans = *L. interrogans*-LA2540.

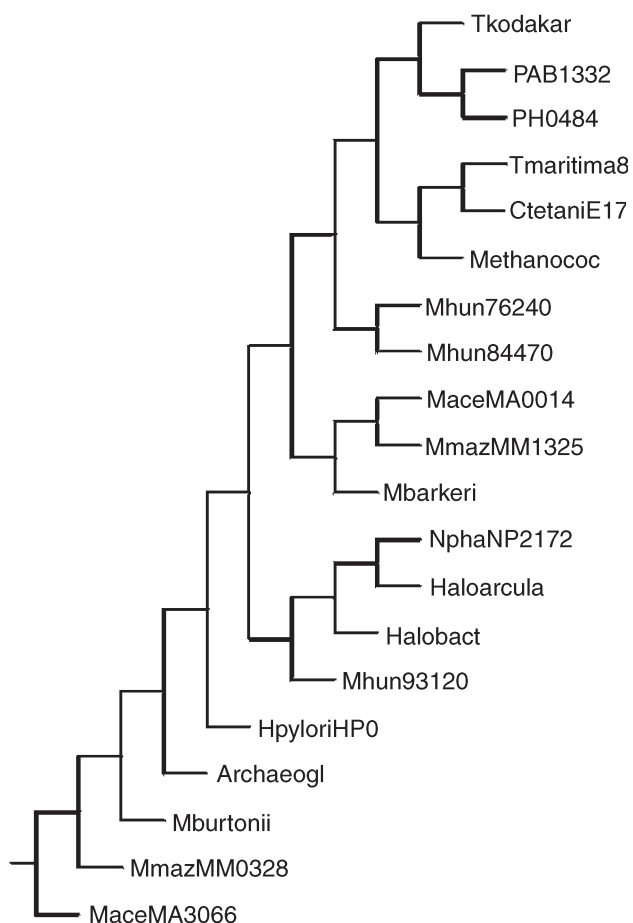


Figure A2. Phylogenetic analysis by neighbor-joining of putative two-component open reading frames (ORFs) from histidine kinase VI groups *ahk40-43*. Group *ahk40* = MaceMA3066, MmazMM0328, AfulAF1040 and Mbur_401647520; group *ahk41* = MaceMA0014, MmazMM1325 and MbarA_0984; group *ahk42* = PAB1332, PH-0484, Tkod_610170420/30 and MmarMMP0927; and group *ahk43* = HaloVNG0971G, HmarrnAC2205, NphaNP2172A and Mhun_401793120. The bacterial ORFs are CtetaniE17170, TmaritimaMSB_824070 and H pyloriHP0392. Abbreviations: CtetaniE17170 = *C. tetani*E88_1817170; *T. maritima* = TmaritimaMSB_824070; and *H. pylori* = H pyloriHP0392.

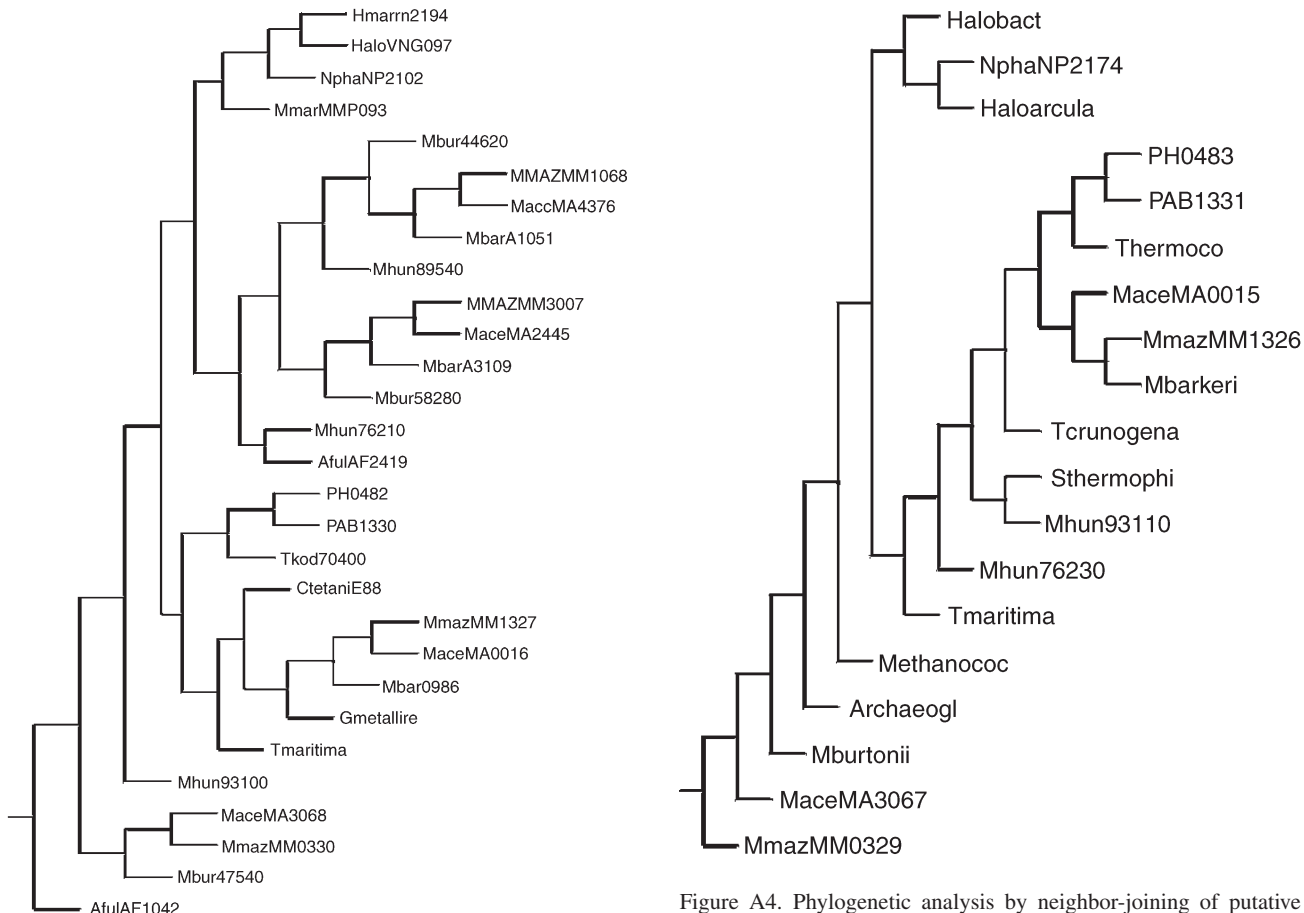


Figure A3. Phylogenetic analysis by neighbor-joining of putative two-component open reading frames (ORFs) from response regulator I groups *arr7*, *arr10*, *arr12* and *arr14*. Group *arr7* = HaloVNG0974G, AfulAF1042, MmarMMP0933, Mbur_401647540, MaceMA3068, MmazMM0330, PAB1330, PH0482, HmarrnAC2194, Tkod_610170400, Mhun_401793100 and NphaNP2102A; group *arr10* = MaceMA0016, MbarA_0986 and MmazMM1327; group *arr12* = MaceMA4376, MbarA_1051, MmazMM1068, AfulAF2419, Mbur_401644620 and Mhun_401789540; and group *arr14* = MaceMA2445, MbarA_3109, MMAZMM3007, Mbur_401658280 and Mhun_401776210. The bacterial ORFs are CtetaniE88, Tmaritima and Gmetallire. Abbreviations: CtetaniE88 = *C. tetanii*E88_1817150; Tmaritima = *T. maritima*MSB8_24050; and Gmetallire = *G. metallireducens*_401349760.

Figure A4. Phylogenetic analysis by neighbor-joining of putative two-component open reading frames (ORFs) from response regulator IV groups *arr18* and *arr19*. Group *arr18* = MaceMA0015, MbarA_0985 and MmazMM1326; and group *arr19* = PAB1331, PH0483, Tkod_610170410, HaloVNG0973G, HmarrnAC2204, MaceMA3067, MmazMM0329, Mbur_401647530, AfulAF1041, MmarMMP0926, NphaNP2174A and Mhun_401793110. The bacterial ORFs are Sthermophilum and Tmaritima, Tcrunogena. Abbreviations: Thermoco = *T. kodakaraensis*; Sthermophilum = *S. thermophilum*_3768360; Tmaritima = *T. maritima*MSB8_21060; and Tcrunogena = *T. crunogena* Tcr0758.

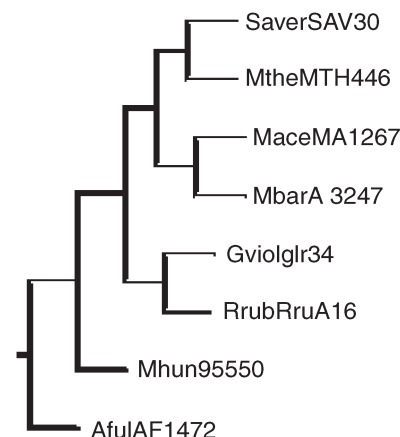


Figure A5. Phylogenetic analysis by neighbor-joining of putative two-component open reading frames (ORFs) from hybrid kinase 1 group *ahy2*. The bacterial ORFs are Gviolglr3434, RrubRruA1653 and SaverSAV3017. Abbreviations: Gviolglr3434 = *G. violaceus*; RrubRruA1653 = *R. rubrum*; and SaverSAV3017 = *S. avermitilis*.

Table A2. Closely linked genes that may be part of operons.

Assigned gene no.	Classification	Orthologue number	Additional information
<i>Archaeoglobus fulgidus</i>			
AF448	His_KA PACPASGAF		
AF449	RRI-CheY		
AF450	HKIII + 1 PASPAC		
AF1042	RRI-CheY	arr7	AF1055 mcp
AF1041	RRIV-CheB	arr18	AF1044CheW
AF1040	HKVI	arr40	AF1039 CheC
AF1472	HYI-HisKA	ahy2	
AF1473	RRI-CheY	arr9	
AF2419	RRI-CheY	arr12	
AF2420	His_KA PACPASGAF		
<i>Halobacterium</i>			
VNG0974G	RRI-CheY	arr7	VNG0976G CheW
VNG0973G	RRIV-CheB	arr19	VNG0970G CheC1
VNG0971G	HKVI	ahk43	VNG0967G CheD
VNG0966G CheR	HKIII	ahk22	
VNG1374G	HATPase HAMP		
VNG1375G			
VNG2036C	RRI-CheY	arr1	
VNG2037C	HKII		
<i>Haloarcula marismortui</i>			
rrnAC0410	HKII + 2PASPAC	arr9	
rrnAC0411	RRI-CheY		
rrnAC0412	HYI		
rrnAC0413	HKII + 1PASPAC		
rrnAC0456	HATPase_c+ gyra		Top6B
rrnAC0457	HATPase_c		Top6A
rrnAC2204	RRIV-CheB	arr19	rrnAC2206 CheR
rrnAC2205	HKVICheW-CheA	ahk43	
rrnAC2692	HYI + PASPACGAF		
rrnAC2694	HATPase_c + PAS		
<i>Methanobacter thermoautotrophicus</i>			
MTH444	HKI	ahy9	
MTH445	RRI-CheY	arr9	
MTH446	HYI + (PASPAC) ₂	ahy2	
MTH447	RRI-CheY	arr6	
MTH457	RRIV-PACPAS		
MTH459	HKI		
MTH548	RRIV-G_transf		
MTH549	RRI-CheY		
MTH901	HYI		
MTH902	HYI-PASPAC		
<i>Methanococcus maripaludis</i>			
MMP0926	RRI-CheB	arr19	MMP0925 CheW
MMP0927	HKVI	ahk42	MMP0928 CheD
MMP0933	RRI-CheY	arr7	MMP0929 mcp
MMP1303	HKIV	ahk32	
MMP1304	RRI-CheY	arr2	
<i>Methanosarcina acetivorans</i>			
MA0014	HKVI	ahk41	
MA0015	RRI-CheB	arr19	
MA0016	RRI-CheY	arr10	
MA0018	RRI-CheY	arr16	MA0019 mcp MA0020 CheW

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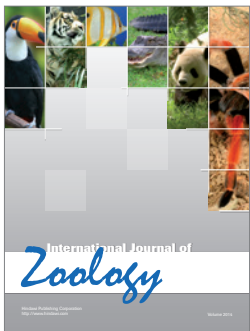
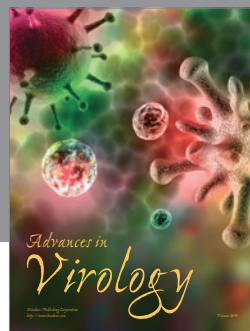
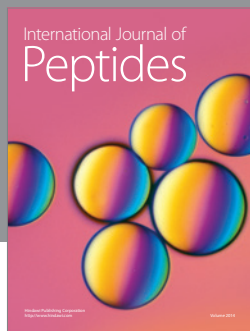
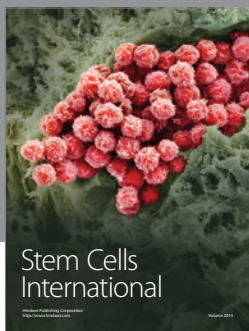
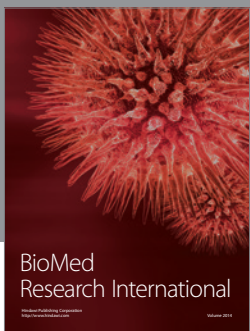
Table A2. Cont'd. Closely linked genes that may be part of operons.

Assigned gene no.	Classification	Orthologue number	Additional information
<i>Methanosa</i> cina acetivorans cont'd			
MA0551	HKII + PASPAC		
MA0552	HKIII	ahk9	
MA0619	HKIII	ahk12	
MA0620	HKIII	ahk14	
MA0758	HKIII		
MA0759	HKIII	ahk30	
MA1267	HYI + PASPAC	ahy2	
MA1268	RRI-CheY	arr9	
MA1269	RRI-CheY		
MA1270	HKII + PASPAC		
MA1468	RRI-CheY		
MA1469	RRI-CheY	arr8	
MA1470	HKIII	ahk8	
MA1627	HKIII	ahk25	
MA1628	HKII + PASPAC	ahk4	
MA1645	HKIII	ahk17	
MA1646	HKII + PASPAC	ahk6	
MA2012	RRI-CheY		
MA2013	HYII + Hpt		
MA3066	HKVI	ahk40	MA3063 CheR
MA3068	RRI-CheY	arr7	MA3064 CheD
MA3067	RRI-CheB	arr18	MA3065 CheC MA3070 CheW
MA3368	HKIII	ahk24	
MA3370	HKII + PASPAC		
MA4376	RRI-CheY	arr12	
MA4377	HYIII CHSE4HP	ahy7	
<i>Methanosa</i> cina barkeri			
MbarA_0984	HKVI	ahk41	MbarA_0983 CheR
MbarA_0985	RRIV-CheB	arr19	
MbarA_0986	RRI-CheY	arr10	
MbarA_0988	RRI-CheY		MbarA_0989 mcp MbarA_0990 CheW
Opp orientation to above			
MbarA_1051	RRI-CheY	arr12	
MbarA_1052	HYIII	ahy7	
MbarA_3036	HKII		
MbarA_3037	HKII		
MbarA_3247	HYIPASPAC		
MbarA_3248	RRI-CheY		
MbarA_3250	HKII		
MbarA_3447	HKII		
MbarA_3448			
<i>Methanosa</i> cina mazei			
MM0168	HKII	ahk9	
MM0169	HKIII		
MM0328	HKVI	ahk40	MM3025 CheR
MM0329	RRI-CheB	arr18	MM0326 CheD
MM0330	RRI-CheY	arr7	MM0327 CheC MM0332 CheW MM0333 mcp
MM1325	HKVI	ahk41	MM1323 CheC
MM1326	RRI-CheY	arr19	MM1324
MM1327		arr10	CheB
MM1328 opp orientation to above	RRI-CheY	arr16	MM1329 mcp MM1330 CheW

continued overleaf

Table A2. Cont'd. Closely linked genes that may be part of operons.

Assigned gene no.	Classification	Orthologue number	Additional information
<i>Methanosarcina mazei</i> cont'd.			
MM2275	HKIII	<i>ahk28</i>	
MM2276	HKIII		
MM2277	HKIII		
MM2515	HKIII	<i>ahk8</i>	
MM2516	RRI-CheY	<i>arr8</i>	
MM2880	RRI-CheY	<i>arr3</i>	
MM2881	HYIII	<i>ahy6</i>	
MM2953	RRI-CheY	<i>arr4</i>	
MM2954	RRI-CheY	<i>arr5</i>	
MM2955	HKVIPASPACCach		
MM3205	HYIII		
MM3206	RRI-CheY		
<i>Pyrococcus abyssi</i>			
PAB1330	RRI-CheY	<i>arr7</i>	PAB1329 CheR
PAB1331	RRVI-CheB	<i>arr19</i>	PAB1333 CheC
PAB1332	HKVI	<i>ahk42</i>	PAB1334 CheC PAB1335 CheW PAB1336 mcp
<i>Pyrococcus horikoshi</i>			
PH0482	RRI-CheY	<i>arr7</i>	PH0481 CheB
PH0483	RRIV-CheB	<i>arr19</i>	
PH0484	HKVI	<i>ahk42</i>	



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