Correspondence Suzanna Bräuer

brauersl@appstate.edu

Genome of *Methanoregula boonei* 6A8 reveals adaptations to oligotrophic peatland environments

Suzanna Bräuer,¹ Hinsby Cadillo-Quiroz,² Nikos Kyrpides,³ Tanja Woyke,³ Lynne Goodwin,⁴ Chris Detter,⁴ Sheila Podell,⁵ Joseph B. Yavitt,⁶ and Stephen H. Zinder⁷

¹Department of Biology, Appalachian State University, Boone, NC 28608, USA
²Swette Center for Environmental Biotechnology at the Biodesign Institute, Arizona State University, Tempe, AZ 85287-4501, USA
³Department of Energy, Joint Genome Institute, Walnut Creek, CA 94598, USA
⁴Los Alamos National Laboratory, Los Alamos, NM 87545, USA
⁵Scripps Institution of Oceanography, La Jolla, CA 92093, USA
⁶Department of Natural Resources, Cornell University, Ithaca, NY 14853, USA
⁷Department of Microbiology, Cornell University, Ithaca, NY 14853, USA
Analysis of the genome sequence of *Methanoregula boonei* strain 6A8, an acidophilic methanogen isolated from an ombrotrophic (rain-fed) peat bog, has revealed unique features that likely allow it to available in acidia. putrient pear and iting. Eint *M* boons(in prodicted to a strain for the pear for the pear for the pear for the pear formation of the pear

that likely allow it to survive in acidic, nutrient-poor conditions. First, *M. boonei* is predicted to generate ATP using protons that are abundant in peat, rather than sodium ions that are scarce, and the sequence of a membrane-bound methyltransferase, believed to pump Na⁺ in all methanogens, shows differences in key amino acid residues. Further, perhaps reflecting the hypokalemic status of many peat bogs, *M. boonei* demonstrates redundancy in the predicted potassium uptake genes *trk*, *kdp* and *kup*, some of which may have been horizontally transferred to methanogens from bacteria, possibly *Geobacter* spp. Overall, the putative functions of the potassium uptake, ATPase and methyltransferase genes may, at least in part, explain the cosmopolitan success of group E1/E2 and related methanogenic archaea in acidic peat bogs.

Received 24 March 2015 Accepted 20 May 2015

INTRODUCTION

Methanoregula boonei is an acidophilic methanogen isolated from an ombrotrophic peat bog (McLean Bog) in New York State, USA (Bräuer *et al.*, 2006a). A member of the Euryarcheal order *Methanomicrobiales*, this archaeon demonstrates physiological evidence of adaptation to nutrient-poor low ionic strength environments, such as ability to grow at 0.4 mM Na⁺ and sensitivity to >50 mM sodium (Bräuer *et al.*, 2011) in contrast to methanogens described elsewhere (Jarrell & Kalmokoff, 1988). As *M. boonei* has been described previously (Bräuer *et al.*, 2011), this paper will focus on a summary of genomic evidence revealing the presence of putative genes specific for protonrich and sodium- and potassium-poor environments.

Abbreviations: COG, cluster of orthologous groups; JGI, Joint Genome Institute.

The GenBank/EMBL/DDBJ accession number for the complete genome sequence of *Methanoregula boonei* strain 6A8 is NC_009712.

Two supplementary tables and a supplementary figure are available with the online Supplementary Material.

M. boonei is the type strain (DSMZ=21154^T, JCM=14090^T) within the type genus of the family *Methanoregulaceae* (Sakai *et al.*, 2012). Cultures are dimorphic, containing thin rods (0.2–0.3 µm in diameter and 0.8–3.0 µm long) and irregular cocci (0.2–0.8 µm in diameter). In PM1 medium, *M. boonei* appears to be an obligate hydrogenotroph and is unable to utilize formate, acetate, methanol, ethanol, 2-propanol, butanol or trimethylamine (Bräuer *et al.*, 2011). Optimal growth conditions are near 35–37 °C and pH 5.1, with growth occurring at pH values as low as 3.8.

METHODS

Preparation of DNA and genome sequencing. *M. boonei* was cultured as described previously (Bräuer *et al.*, 2006b). An exponentially growing culture (11) was harvested by cold centrifugation and DNA was extracted using a GNOME DNA isolation kit (MP Biomedicals), following the manufacturer's protocols except that a final concentration of 0.1% SDS was added in addition to the cell lysis/denaturing solution to increase cell lysis. Genomic DNA was then evaluated for quality and concentration prior to sequencing.

Table	1. Genome	features	of N	1. boonei	6A8
-------	-----------	----------	------	-----------	-----

Feature	Genome (total)			
	Value	% of total*		
Size (bp)	2542943			
No. of $G + C$ bases	1386250	54.5		
Coding sequence (bp)	2201702	86.6		
Mean ORF length (bp)	893			
5S rRNA	1			
16S rRNA	1			
23S rRNA	1			
tRNA genes	48			
Other RNA genes	3			
Total no. of genes	2518			
Proteins with function prediction	1617	64.2		
Proteins without function prediction	847	33.6		
GenBank accession no.	NC_009712			

*The total is based on either the size of the genome in bp or the total number of protein encoding genes in the annotated genome.

The genome of M. boonei 6A8 was sequenced at the Joint Genome Institute (JGI) using a combination of 3, 8 and 40 kb (fosmid) DNA libraries. All general aspects of library construction and sequencing performed at the JGI can be found at http://jgi.doe.gov/. Draft assemblies were based on 37 430 total reads. All three libraries provided 13 × coverage of the genome. The Phred/Phrap/Consed software package (www.phrap.com) was used for sequence assembly and quality assessment (Ewing & Green, 1998; Ewing et al., 1998; Gordon et al., 1998). After the shotgun stage, reads were assembled with parallel Phrap (High Performance Software). Possible misassemblies were corrected with Dupfinisher (Han & Chain, 2006). Gaps between contigs were closed by editing in Consed or custom primer walk. A total of 921 additional reactions were necessary to close gaps and to raise the quality of the finished sequence. The completed genome sequences of M. boonei 6A8 contains 37 526 reads, achieving a mean of 13-fold sequence coverage per base with an error rate of less than 1 in 100 000.

Additional gene functional annotation and comparative analyses were performed within the Integrated Microbial Genomes (IMG/ER) platform (Markowitz *et al.*, 2006). Alignments of functional genes were conducted in BioEdit using CLUSTAL w (Larkin *et al.*, 2007). Phylogenetic trees were reconstructed using the PHYLIP software

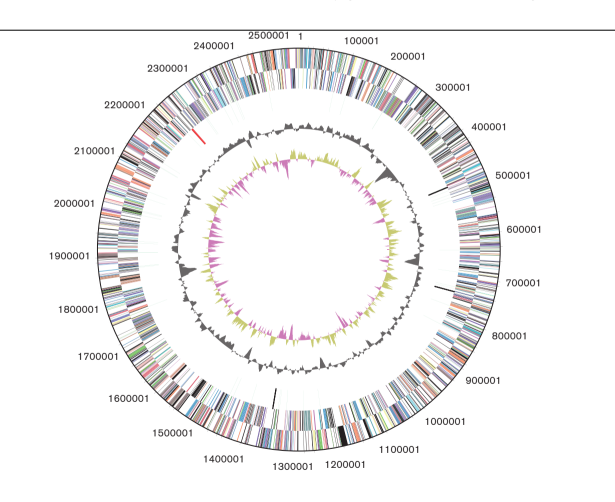


Fig. 1. Circular map of the *M. boonei* 6A8 genome. From the outside ring to the centre: (i) genes on the forward strand, coloured by COG category (Tatusov *et al.*, 2000); (ii) genes on the reverse strand, coloured by COG category; (iii) RNA genes with tRNA genes coloured green, rRNA genes red and other RNAs black; (iv) GC content; (v) GC skew. This figure is available on the IMG website (https://img.jgi.doe.gov/) (Markowitz *et al.*, 2006).

package (Felsenstein, 2004) by conducting both neighbour-joining and maximum-likelihood analysis.

Nucleotide accession number. The complete genome sequence of *M. boonei* strain 6A8 is available in the National Center for Biotechnology Information database (Wheeler *et al.*, 2007) under GenBank/EMBL/DDBJ accession number NC_009712. Additionally, the genome is available in the IMG system (Markowitz *et al.*, 2006) and the JGI genome portal (Grigoriev *et al.*, 2012).

RESULTS AND DISCUSSION

Genome sequencing and annotation information

M. boonei was selected for sequencing due to its potential energy production (methane), biogeochemical importance in global carbon cycling and occurrence in habitats that are unique for cultured methanogens, i.e. proton rich and nutrient element poor. Sequencing, assembly and annotation were conducted by the Department of Energy JGI. A summary of the genome sequencing information can be found in Table 1.

Genome properties

The genome consists of one single circular chromosome of approximately 2.5 million bp with 2518 genes identified, including one rRNA gene operon (Table 1, Fig. 1). Highlighting our dearth of knowledge of methanogenic archaea, only 36% of the genes were associated with one of the welldefined cluster of orthologous groups (COG) categories (Tatusov *et al.*, 2000), with the remaining 64% either not associated with a COG (39%), or associated only by general (14%) or unknown (12%) function (Table 2). The majority of the COG genes in *M. boonei* were predicted to be involved in energy production and conversion (9%), translation (8%), and transport and metabolism of amino-acids (8%), coenzymes (6%) and ions (6%).

Adaptation to high proton concentrations

In McLean Bog, the pH is near 4 and the H⁺ concentration is approximately 10^{-4} M (100 μ M), three orders of magnitude greater than at pH 7. Moreover, sphagnum moss impedes the flow of mineral-rich groundwater into the bog so that the only water source is rain, essentially distilled water. Consequently, Na⁺ concentrations are typically low. For example, Na⁺ concentrations were measured as only 2 μ M in the McLean Bog porewater (Bräuer *et al.*, 2004). These exceedingly low external Na⁺ concentrations make developing and conserving a sodium motive force challenging; however, sodium motive force is considered essential to energy conservation by methanogens (Schlegel & Müller, 2013).

All cultured members of the groups *Methanobacteriales*/*Methanobacteri*

1574

(Mtr) (Schlegel & Müller, 2013; Thauer et al., 2008). Further, the A1A0 ATPase/synthases studied among the members of the groups Methanobacteriales/Methanococcales have clearly been shown to pump Na⁺ (McMillan et al., 2011; Mulkidjanian et al., 2008). In contrast, the cvtochrome-containing methanogens in the Methanosarcinales have Mtr complexes, but also have steps that pump protons (Schlegel & Müller, 2013; Thauer et al., 2008), and evidence has favoured H⁺ pumping by the ATPases in these organisms (Müller et al., 1999; Pisa et al., 2007). More recently, it was demonstrated that the A_1A_0 archaeal ATPase/synthase in Methanosarcina acetivorans is 'promiscuous', pumping either Na⁺ or H⁺ (Schlegel et al., 2012) with both ions possible at neutral pH, especially at seawater salinity of 0.4 M Na⁺, whereas only protons were pumped at pH 5. M. boonei belongs to the Methanomicrobiales, which lack cytochromes like the Methanobacteriales/Methanococcales cluster, but are more closely related to the Methanosarcinales, and it is not clear which patterns of bioenergetics this group follows.

It is the AtpC/K subunit of the membrane bound A_0 subunit of the A_1A_0 ATPase/synthase complex that is responsible for

Table 2. Genes associated with COG functional categories

COG category	No. of genes	% of total*
Energy production and conversion	157	9.35
Translation, ribosomal structure and biogenesis	134	7.98
Amino acid transport and metabolism	133	7.92
Coenzyme transport and metabolism	108	6.43
Inorganic ion transport and metabolism	96	5.71
Signal transduction mechanisms	95	5.65
Transcription	83	4.94
Post-translational modification, protein turnover, chaperones	69	4.11
Replication, recombination and repair	69	4.11
Carbohydrate transport and metabolism	61	3.63
Cell wall/membrane/envelope biogenesis	55	3.27
Nucleotide transport and metabolism	53	3.15
Cell motility	28	1.67
Defence mechanisms	27	1.61
Lipid transport and metabolism	24	1.43
Intracellular trafficking, secretion and vesicular transport	22	1.31
Cell cycle control, cell division, chromosome partitioning	18	1.07
Secondary metabolites biosynthesis, transport and catabolism	15	0.89
Chromatin structure and dynamics	2	0.12
General function prediction only	227	13.51
Function unknown	204	12.14
Not in a COG category	971	38.56

*The total is based on the total number of protein encoding genes in the annotated genome.

pumping either H^+ or Na^+ , with the ion typically binding to a conserved aspartate or a glutamate located within a transmembrane helix. In Table 3, partial sequences of the AtpC/K are aligned, and residues critical to Na^+ binding in the *Methanobacteriales/Methanococcales* (Grüber *et al.*, 2014; McMillan *et al.*, 2011) are indicated in bold. Also shown in bold are residues in the *M. acetivorans* sequence that are considered crucial to being able to bind either Na^+ or H^+ . The *M. boonei* sequence shares these residues as well as many others with *M. acetivorans*, and it is likely that its ATPase pumps protons near pH 4.

Another question of interest is whether the Mtr complex in *M. boonei* pumps Na^+ . In *Methanosarcina mazei* and *Methanothermobacter marburgensis*, Na^+ pumping was demonstrated for the Mtr complex, thereby leading to a sodium motive force. From these results, it was extrapolated

that all Mtr complexes pump Na⁺ (Schlegel & Müller, 2013). The pumping is attributed to the membrane-bound MtrE subunit, and a specific aspartate (indicated by the bold D) predicted to be within a transmembrane helix, part of the motif 168-IWGITIGAIGSSTGDVHYGAER-191 that is conserved between the two organisms (Gottschalk & Thauer, 2001). M. boonei and other members of the Methanomicrobiales have an asparagine instead of aspartate at that position (position 190 in Fig. S1, available in the online Supplementary Material) in their MtrE sequences, which renders that residue unable to pump cations. There is a glutamate at position 253 in Fig. S1 within a region (predicted to be a transmembrane alpha helix by the IMG website) that is conserved amongst Methanomicrobiales as well as some other methanogens, but is not present in the M. marburgensis or Methanosarcina barkeri sequences. This residue may play a role in pumping but, as of now, it is

Table 3. Amino acid alignment of AtpCK demonstrating the two conserved glutamine (Q) and tyrosine (Y) residues (shown in bold) (identified by: McMillan *et al.*, 2011; Mulkidjanian *et al.*, 2008; Sakai *et al.*, 2011) that appear to be unique for methanogens predicted to have sodium-driven ATPases versus those predicted to have proton-driven or sodium-proton driven ATPases

M. boonei is shaded in grey. Organisms predicted to have proton-driven ATPases are shown in red font and those predicted to have sodium-driven ATPases are shown in blue font. Organisms with experimental evidence supporting sodium-driven ATPases are shown in bold blue and include: *Methanococcus jannaschii* (Morsomme *et al.*, 2002), *Methanococcus voltae* (Dybas & Konisky, 1992), *Methanobacterium thermoautotrophicum* (Schönheit & Perski, 1983) and *Methanobrevibacter ruminatum* (McMillan *et al.*, 2011). Organisms with experimental evidence supporting proton-driven (or sodium/proton-driven) ATPases are shown in bold red and include: *M. mazei* Gö1 (Becher & Müller, 1994; Pisa *et al.*, 2007), *Methanosaeta thermophila* (Inatomi *et al.*, 1993) and *M. barkeri* (Blaut & Gottschalk, 1984; Müller *et al.*, 1999). *M. acetivorans* C2A was recently shown to have a sodium-proton driven ATPase (Grüber *et al.*, 2014; Schlegel & Müller, 2013) and the residues considered crucial to being allowing binding of both ions are indicated in bold, as are the corresponding amino acids in the *M. boonei* sequence directly above. IMG gene numbers follow the colon. Blue shading has been added to indicate sequences that align with those of organisms predicted to have sodium-driven ATPases.

Organism	Partial amino aci	id alignment					
Methanospirillum hungatei JF-1:637896821	KAVGAGLAVG	LAGVGSGLGE	MGIGAAAMGA	VAENKDMFGL	ALLFTVLPET	IVIFGLVVAL	LL
Methanospirillum hungatei JF-1:637897383	VPIGAAIAFA	GGAIATGIAQ	SKIGAAGAGT	VAERPESAGT	VIVLEAIPET L	VILGFVVAA	MI
Methanosarcina mazei strain Gö1:638165281	KALGAAIAIA	VTGLASAIAE	KDIGTAAIGA	MAENEGLFGK	GLILTVIPET	IVIFGLVVAL	LI
Methanoregula boonei 6A8:640869605	KAIGAGLAVG	LTGVGTGVAE	MGIGAAAVGA	IAENKDFFGL	GLLFTVIPET	IVIFGLVIAL	LL
Methanosarcina acetivorans C2A: 638179041	KALGAALAIT	VTGLASAWAE	KEIGTAAIGA	MAENEGLFGK	GLILTVIPET	IVIFGLVVAL	LI
Methanosarcina barkeri str. fusaro: 637699281	KAIGASIAIA	LTGIASAIAE	KDIGTAAIGA	MAENEGLFGK	GLILTVIPET	IVIFGLVVAL	LI
Methanoculleus marisnigri JR1:640114955	SAVGAGLAVG	LTGVGTGLAE	MGIGAAAVGA	TAENRDMFGL	ALLFTVIPET	IVIFGLVVAL	LL
Methanosphaerula palustris E1-9c: 643571272	KAVGAGLAVG	LAGIGTGLGE	MGIGAAAMGA	TAENKDMFGL	ALLFTVIPET	IVIFGLVVSL	LL
Methanocella paludicola SANAE: 646465407	VAIGAGLAVG	LAGIGSGIAE	KDIGAAAVGA	IAEDRSFFGQ	GLIFTVIPET	IVIFGLVIAI	LL
Methanosaeta concilii GP6:650798510	IAVGAGLATG	LAGIGAGVGE	QGIGAAVVGV	VAEEPGFLGK	GLFLMLLPET	LIIFGLAVSL	IL
Methanocella paludicola SANAE: 646467386	IPLGAAIAFG	AGAISTGFAQ	ARIGSAGAGA	LSERPELSGL	IIILEAIPET	LAILGFVVAA	MI
Methanohalophilus mahii DSM 5219:646707499	KAIGAGLAVG	LTGLASGIAE	KDIGAAAIGA	MAENEGLFGK	GLIMTVIPET	IVIFGLVVAL	LI
Methanobacterium sp. AL-21:650750551	AAIGAGLAVG	LAGLGSGIGQ	GIAAAGSVGA	VAEDPDMFAR	GIIFTALPET	QAIYGFLIAI	LL
Methanobacterium sp. SWAN-1:650872332	AAIGAGVAVG	FAALGSGIGQ	GIASAGAVGA	VAEDKSMFAQ	GMVFTAIPET	QAIYGFLISI	LL
Methanopyrus kandleri AV19:638169043	AAIGAGLAAG	VAGVGSGIGQ	GIAAAAGAGA	VAEDEATFGK	AIVFSVLPET	QAIYGLLTAI	LI
Methanocaldococcus jannaschii DSM	GAVGAGLAVG	IAGLGSGIGA	GITGASGAGV	VAEDPNKFGT	AIVFQALPQT	Q GL Y GFLVAI	LI
2661:638201515							
Methanospirillum hungatei JF-1:637897394	MAIGAGIAVG	CSAIGSGIGV	GIVGSAASGV	ISERSEKFGM	ALVFTAIPQT	QAIYGLLIAI	LI
Methanothermobacter thermoautotrophicus:	AAIGAGVAVG	FAGLGSGLGQ	GIAAAESVGA	VAENSDMFAR	GIIFSTLPET	QAIYGFLIAI	LL
638155490							
Methanococcus maripaludis C5:640166057	GAIGAGLAVG	IAGLGSGIGA	GITGASGAGV	VAEDPNKFGT	AIVFQALPQT	Q GL Y GFLVAI	LI
Methanobrevibacter smithii DSM	AAIGAGVAIG	FAGLGSGLGQ	GMAAAGSVGA	VAEDNDMFAR	GIIFSALPET	QAIYGFLIAI	LL
2375:644143574							
Methanosphaera stadtmanae DSM	AAIGAGVAVG	FAALGSGIGQ	GIASSASVGA	VAEDSSMFAQ	GLVFTAIPET	QAIYGFLIAI	LL
3091:637847029							
Methanobrevibacter smithii ATCC	AAIGAGVAIG	FAGLGSGLGQ	GMAAAGSVGA	VAEDNDMFAR	GIIFSALPET	QAIYGFLIAI	LL
35061:640592228							
Methanobrevibacter ruminantium	AAIGAGVAIG	FAGLGSGLGQ	GMAAAGSVGA	VAEDNDMFAR	GIIFSALPET	QAIYGFLIAI	LL
M1:646531773							
Methanococcus voltae A3:646858602	GAIGAGLAVG	IAGLGSGIGA	GITGASGAGV	LAEDPKQFSK	VIVFQALPQT	Q GL Y GFLVAI	LI
Methanotorris igneus Kol 5:650856311	GAIGAGLAVG	IAGLGSGIGA	GITGASGAGV	VAEDPNKFGT	AIVFQALPQT	Q GL Y GFLVAI	LI
Methanothermococcus okinawensis	GAVGAGLAVG	IAGLGSGIGA	GITGASGAGV	VAEDPNKFGT	AIVFQALPQT	Q GL Y GFLVAI	LI
IH1:650918276						-	

unclear whether the Mtr complex in *M. boonei* or other *Methanomicrobiales* pumps Na^+ or H^+ , or perhaps is not a pump at all. Because of these fundamental differences in the MtrE sequences between *Methanomicrobiales* and other methanogens, the role of Mtr in their bioenergetics warrants examination, especially since it is considered the only site for energy conservation in these organisms.

Adaptation to low potassium concentrations

Similar to the case for Na⁺, the K⁺ concentrations in McLean Bog porewater are extremely low, less than 25 µM (Bräuer et al., 2004), and cells typically accumulate K⁺ (Epstein, 2003) as well as expel Na⁺. *M. boonei* is predicted to carry genes for three different K⁺ uptake mechanisms including the low-affinity trk genes that many methanogens carry, in addition to the medium-affinity kup genes and the ATP-driven high-affinity kdp genes, both of which are more rarely found among methanogenic archaea (Table S1). Only one other methanogen (sequenced to date) carries all three predicted K⁺ uptake systems, Methanosphaerula palustris E1-9c, and it was also isolated from a peatland ecosystem (Cadillo-Quiroz et al., 2009; Cadillo-Quiroz et al., 2008), a fen in which the pH was neutral but the K⁺ porewater concentrations were only 3-8 µM (Dettling et al., 2007).

In Escherichia coli, the kdp uptake system shows both high specificity and high affinity for potassium, and is required for growth during extreme potassium limitation (Altendorf & Epstein, 1996; Epstein, 2003; Epstein et al., 1990). E. coli cultures with a mutation in the kdp genes have shown growth deficiencies at K⁺ concentrations below 300 µM (Rhoads et al., 1976). Highlighting its importance in M. boonei, the kdpCAB operon has been duplicated and can be identified in two locations in the genome (Fig. 2). Compared to E. coli, both predicted KdpA proteins in M. boonei (Mboo 0443 and 0894) have all four regions (I, 112-NTNWQ-116; II, 230-TNGGG-234; III, 343-SCGAV-347; IV, 468-NNGSA-472; E. coli numbering) demonstrated experimentally (Bertrand et al., 2004; Buurman et al., 1995; Dorus et al., 2001; Schrader et al., 2000; van der Laan et al., 2002) and in 3D structural models (Greie, 2011; Hu et al., 2008) to be responsible for K⁺ binding. Originally identified by the HGT-detection program, DarkHorse (Podell & Gaasterland, 2007), the predicted KdpA proteins in M. boonei and in other methanogens cluster phylogenetically within the Proteobacteria, perhaps most closely resembling those of Geobacter spp. (Fig. 3). Since there are apparently three closely related clades of methanogen KdpA protein sequences, it is unclear how many transfer events have occurred. Moreover, the KdpC protein is predicted to be fused to the N-terminal of KdpA (Fig. 2) in both sets of genes, an arrangement shared with Methanomassiliicoccus luminyensis, a methanogen isolated from human faeces (Dridi et al., 2012) belonging to a new phylum related to Thermoplasma and only able to use H₂ and methanol for methanogenesis. All other methanoarchaea, including the closely related Methanosphaerula, have the canonical

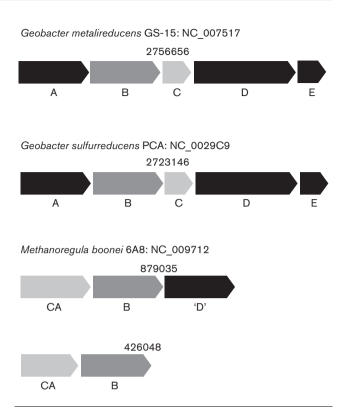


Fig. 2. Diagram of high affinity, ATP-driven potassium uptake (kdp) gene arrangement in *Geobacter* spp., which shows the canonical arrangement, compared to the two kdp operons [Mboo 0894-6 (top) and Mboo 0443-4 (bottom)] in *M. boonei* 6A8. This figure was modified from an image on the IMG website (https://img.jgi.doe.gov/) (Markowitz *et al.*, 2006). Genes encoding for KdpA, KdpB, KdpC, KdpD or KdpE subunits are indicated by an A, B, C, D or E, respectively. The fused kdpC/A gene is indicated by a CA and the predicted pseudogene kpdD is indicated by a 'D'.

kdpABC gene order. Thus, the arrangement and close phylogenetic relationship of their *kdpCA* genes relative to that of other organisms suggests that a gene transfer event occurred between ancestors of *M. boonei* and *M. luminyensis*.

In *E. coli* and many other bacteria, KdpD is a membranebound osmosensitive K^+ -sensing histidine kinase component and KdpE is the response regulator of a two-component transcriptional regulatory system that induces *kdp* genes when K^+ is low (Nakashima *et al.*, 1992; Poolman & Glaasker, 1998) (Fig. 2). A number of methanoarchaea with *kdp* genes possess a *kdpD* gene (Table S1). In *M. boonei*, the *kdpD* gene is predicted to encode a truncated protein lacking the histidine kinase domain (Table S2) and to also lack *kdpE* compared to bacteria. Thus, it is unlikely that KdpD is a transcriptional regulator and it may play some other role in regulating activity of the Kdp or other proteins, since it still maintains membrane-bound sensing domains. Some *Bacteria*, including *Cyanobacteria* and *Deinococcus radiodurans*,

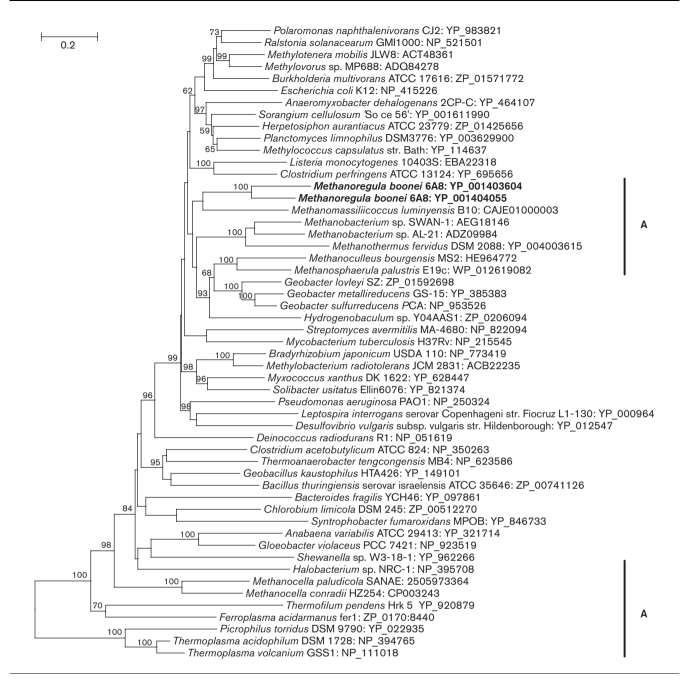


Fig. 3. Neighbour-joining dendrogram of KdpA. The tree was rooted using deep-branching members of the *Thermoplasmatales*. Archaeal sequences are indicated with an A. Accession numbers or IMG gene numbers follow the colon. Bootstrap values greater than 60 are shown for nodes that were supported by maximum-likelihood analysis. *M. boonei* is shown in bold. The scale bar indicates the number of protein changes per site.

also contain a truncated *kdpD* and lack *kdpE* (Ballal *et al.*, 2007). Although *kdpF* is also absent, this gene is non-essential for potassium transport *in vivo*, according to studies in *E. coli* (Gaßel *et al.*, 1999).

Similarly, the *kup* genes may have also been horizontally transferred between *Geobacteraceae* and *Methanosarcina* spp. and several members of the *Methanomicrobiales*

(Fig. 4). However, the data are less robust since the *Geobacter* spp. KupA protein sequences do not cluster with other *Proteobacteria*, so the direction of transfer is unclear. Still, it's most likely that the methanoarchaeal KupA sequences were derived from *Bacteria*, according to the phylogenetic clustering (Fig. 4). Essentially all of the methanoarchaea possess low-affinity potassium uptake genes (*trk*; Table S1); thus, these genes will not be discussed.

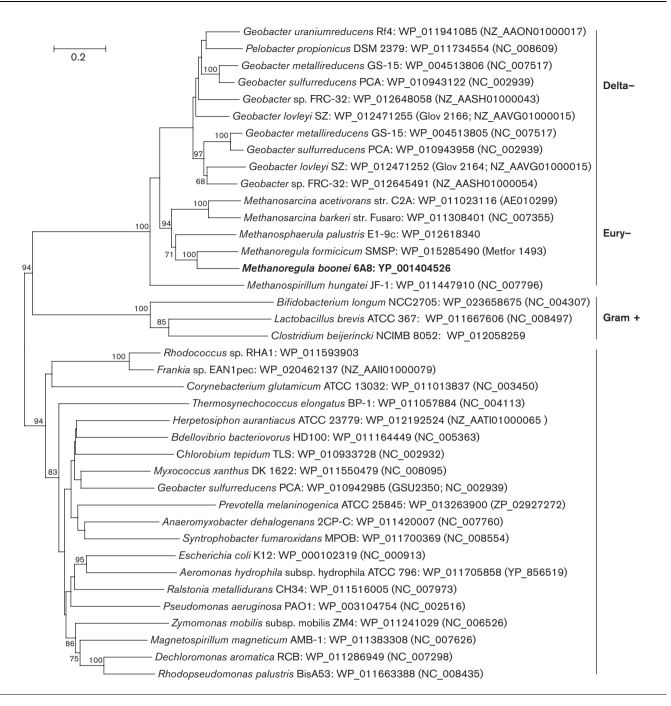


Fig. 4. Unrooted neighbour-joining dendrogram of the predicted KupA protein in *M. boonei* compared to that of other organisms. Several distinct clusters are shown including: one that contains members of the phyla *Cyanobacteria*, *Choroflexi*, *Chlorobi*, *Proteobacteria* and *Actinobacteria* (unlabelled); one that includes Gram-positive *Actionobacteria* and *Firmicutes* (Gram+); and one that includes *Proteobacteria* of the genus *Geobacter* (Delta-), as well as members of the *Methanomicorbia* class of the *Euryarchaeota* (Eury-). Bootstrap values greater than 60 are shown for nodes that were supported by maximum-likelihood analysis. *M. boonei* is shown in bold. The scale bar indicates the number of protein changes per site.

CONCLUSION

Organisms related *to M. boonei* in the E1/E2 cluster, R10, or fen cluster are widespread throughout acidic to moderately acidic peatlands in Germany (Hamberger *et al.*, 2008; Wüst et al., 2009), England (Edwards et al., 1998; Hales et al., 1996), Russia (Kotsyurbenko et al., 2007), Scandinavia (Galand et al., 2005; Høj et al., 2005), the United States (Basiliko et al., 2003; Cadillo-Quiroz et al., 2006; Hawkins et al., 2014) and Canada (Godin et al., 2012; Yavitt et al.,

2006). Further, this group tends to dominate in ombrotrophic bogs and is often outcompeted in minerotrophic fens, where the methanogenic community becomes more diverse (Galand *et al.*, 2005; Kotsyurbenko, 2010; Kotsyurbenko *et al.*, 2007). For example, microbial diversity was shown to increase along a gradient from pH 4.2 in an ombrotrophic bog to 5.1 in a mesotrophic fen in Finland (Juottonen *et al.*, 2005). Similarly, a fen in Minnesota was found to have higher diversity than that of a nearby bog (Lin *et al.*, 2012). The genome of *M. boonei* harbours evidence of adaptation to a proton-rich, sodium-poor and potassium-poor environment, which may, in part, explain the cosmopolitan success of this and related organisms in acidic peat bogs.

ACKNOWLEDGEMENTS

The authors thank the following employees of the US Department of Energy Joint Genome Institute - Marcel Huntemann, Alex Copeland, Amy Chen, Victor Markowitz, Krishnaveni Palaniappan, Natalia Ivanova, Natalia Mikhailova, Galina Ovchinnikova, Evan Andersen, Amrita Pati, Dimitrios Stamatis, T. B. K. Reddy, Chew Yee Ngan, Mansi Chovatia, Chris Daum, Nicole Shapiro and Michael N. Cantor - as well as the following employees of the Los Alamos National Lab - Hazuki Teshima, Olga Chertkov, Hajnalka Daligault, Karen Davenport, Wei Gu, Christine Munk, Xiaojing Zhang, David Bruce, Yan Xu, Beverly Quintana, Krista Reitenga, Yulia Kunde, Lance Green, Tracy Erkkila, Cliff Han and Patrick Chain - for assistance with genome sequencing. The work conducted by the US Department of Energy Joint Genome Institute, a Department of Energy Office of Science User Facility, is supported by the Office of Science of the US Department of Energy under contract no. DE-AC02-05CH11231.

REFERENCES

Altendorf, K. & Epstein, W. (1996). The Kdp-ATPase of *Escherichia coli*. In *Biomembranes: a Multi-Volume Treatise*, 5, pp. 403–420. Edited by A. G. Lee. Greenwich, CT: JAI Press.

Ballal, A., Basu, B. & Apte, S. K. (2007). The Kdp-ATPase system and its regulation. J Biosci 32, 559–568.

Basiliko, N., Yavitt, J. B., Dees, P. M. & Merkel, S. M. (2003). Methane biogeochemistry and methanogen communities in two northern peatland ecosystems. New York State. *Geomicrobiol J* **20**, 563–577.

Becher, B. & Müller, V. (1994). Delta mu Na⁺ drives the synthesis of ATP via an delta mu Na⁺-translocating F_1F_0 -ATP synthase in membrane vesicles of the archaeon *Methanosarcina mazei* Gö1. *J Bacteriol* **176**, 2543–2550.

Bertrand, J., Altendorf, K. & Bramkamp, M. (2004). Amino acid substitutions in putative selectivity filter regions III and IV in KdpA alter ion selectivity of the KdpFABC complex from *Escherichia coli*. *J Bacteriol* 186, 5519–5522.

Blaut, M. & Gottschalk, G. (1984). Coupling of ATP synthesis and methane formation from methanol and molecular hydrogen in *Methanosarcina barkeri. Eur J Biochem* 141, 217–222.

Bräuer, S. L., Yavitt, J. B. & Zinder, S. H. (2004). Methanogenesis in McLean Bog, an acidic peat bog in upstate New York: stimulation by H_2/CO_2 in the presence of rifampicin, or by low concentrations of acetate. *Geomicrobiol J* **21**, 433–443.

Bräuer, S. L., Cadillo-Quiroz, H., Yashiro, E., Yavitt, J. B. & Zinder, S. H. (2006a). Isolation of a novel acidiphilic methanogen from an acidic peat bog. *Nature* 442, 192–194.

Bräuer, S. L., Yashiro, E., Ueno, N. G., Yavitt, J. B. & Zinder, S. H. (2006b). Characterization of acid-tolerant H₂/CO₂-utilizing methanogenic enrichment cultures from an acidic peat bog in New York State. *FEMS Microbiol Ecol* **57**, 206–216.

Bräuer, S. L., Cadillo-Quiroz, H., Ward, R. J., Yavitt, J. B. & Zinder, S. H. (2011). *Methanoregula boonei* gen. nov., sp. nov., an acidiphilic methanogen isolated from an acidic peat bog. *Int J Syst Evol Microbiol* 61, 45–52.

Buurman, E. T., Kim, K.-T. & Epstein, W. (1995). Genetic evidence for two sequentially occupied K^+ binding sites in the Kdp transport ATPase. *J Biol Chem* **270**, 6678–6685.

Cadillo-Quiroz, H., Bräuer, S., Yashiro, E., Sun, C., Yavitt, J. & Zinder, S. (2006). Vertical profiles of methanogenesis and methanogens in two contrasting acidic peatlands in central New York State, USA. *Environ Microbiol* **8**, 1428–1440.

Cadillo-Quiroz, H., Yashiro, E., Yavitt, J. B. & Zinder, S. H. (2008). Characterization of the archaeal community in a minerotrophic fen and terminal restriction fragment length polymorphism-directed isolation of a novel hydrogenotrophic methanogen. *Appl Environ Microbiol* **74**, 2059–2068.

Cadillo-Quiroz, H., Yavitt, J. B. & Zinder, S. H. (2009). *Methanosphaerula palustris* gen. nov., sp. nov., a hydrogenotrophic methanogen isolated from a minerotrophic fen peatland. *Int J Syst Evol Microbiol* 59, 928–935.

Dettling, M. D., Yavitt, J. B., Cadillo-Quiroz, H., Sun, C. & Zinder, S. H. (2007). Soil–methanogen interactions in two peatlands (bog, fen) in Central New York State. *Geomicrobiol J* 24, 247–259.

Dorus, S., Mimura, H. & Epstein, W. (2001). Substrate-binding clusters of the K⁺-transporting Kdp ATPase of *Escherichia coli* investigated by amber suppression scanning mutagenesis. *J Biol Chem* **276**, 9590–9598.

Dridi, B., Fardeau, M. L., Ollivier, B., Raoult, D. & Drancourt, M. (2012). *Methanomassiliicoccus luminyensis* gen. nov., sp. nov., a methanogenic archaeon isolated from human faeces. *Int J Syst Evol Microbiol* **62**, 1902–1907.

Dybas, M. & Konisky, J. (1992). Energy transduction in the methanogen *Methanococcus voltae* is based on a sodium current. *J Bacteriol* **174**, 5575–5583.

Edwards, C., Hales, B. A., Hall, G. H., McDonald, I. R., Murrell, J. C., Pickup, R., Ritchie, D. A., Saunders, J. R., Simon, B. M. & Upton, M. (1998). Microbiological processes in the terrestrial carbon cycle: methane cycling in peat. *Atmos Environ* **32**, 3247–3255.

Epstein, W. (2003). The roles and regulation of potassium in bacteria. *Prog Nucleic Acid Res Mol Biol* **75**, 293–320.

Epstein, W., Walderhaug, M. O., Polarek, J. W., Hesse, J. E., Dorus, E., Daniel, J. M., Green, N. M. & Broome-Smith, J. (1990). The bacterial Kdp K⁺-ATPase and its relation to other transport ATPases, such as the Na⁺/K⁺- and Ca²⁺-ATPases in higher organisms. *Philos Trans R Soc Lond B Biol Sci* 326, 479–487.

Ewing, B. & Green, P. (1998). Base-calling of automated sequencer traces using Phred. II. Error probabilities. *Genome Res* 8, 186–194.

Ewing, B., Hillier, L., Wendl, M. C. & Green, P. (1998). Base-calling of automated sequencer traces using Phred. I. Accuracy assessment. *Genome Res* 8, 175–185.

Felsenstein, J. (2004). PHYLIP (phylogeny inference package) version 3.68. Distributed by the author. Seattle, USA: Department of Genome Sciences, University of Washington.

Galand, P. E., Fritze, H., Conrad, R. & Yrjälä, K. (2005). Pathways for methanogenesis and diversity of methanogenic archaea in three boreal peatland ecosystems. *Appl Environ Microbiol* **71**, 2195–2198.

Gaßel, M., Möllenkamp, T., Puppe, W. & Altendorf, K. (1999). The KdpF subunit is part of the K⁺-translocating Kdp complex of *Escherichia coli* and is responsible for stabilization of the complex *in vitro. J Biol Chem* **274**, 37901–37907.

Godin, A., McLaughlin, J. W., Webster, K. L., Packalen, M. & Basiliko, N. (2012). Methane and methanogen community dynamics across a boreal peatland nutrient gradient. *Soil Biol Biochem* **48**, 96–105.

Gordon, D., Abajian, C. & Green, P. (1998). Consed: a graphical tool for sequence finishing. *Genome Res* **8**, 195–202.

Gottschalk, G. & Thauer, R. K. (2001). The Na⁺-translocating methyltransferase complex from methanogenic archaea. *Biochim Biophys Acta* **1505**, 28–36.

Greie, J.-C. (2011). The KdpFABC complex from *Escherichia coli*: a chimeric K^+ transporter merging ion pumps with ion channels. *Eur J Cell Biol* **90**, 705–710.

Grigoriev, I. V., Nordberg, H., Shabalov, I., Aerts, A., Cantor, M., Goodstein, D., Kuo, A., Minovitsky, S., Nikitin, R. & other authors (2012). The genome portal of the Department of Energy Joint Genome Institute. *Nucleic Acids Res* 40, D26–D32.

Grüber, G., Manimekalai, M. S., Mayer, F. & Müller, V. (2014). ATP synthases from archaea: the beauty of a molecular motor. *Biochim Biophys Acta* 1837, 940–952.

Hales, B. A., Edwards, C., Ritchie, D. A., Hall, G., Pickup, R. W. & Saunders, J. R. (1996). Isolation and identification of methanogenspecific DNA from blanket bog peat by PCR amplification and sequence analysis. *Appl Environ Microbiol* 62, 668–675.

Hamberger, A., Horn, M. A., Dumont, M. G., Murrell, J. C. & Drake, H. L. (2008). Anaerobic consumers of monosaccharides in a moderately acidic fen. *Appl Environ Microbiol* 74, 3112–3120.

Han, C. S. & Chain, P. (2006). Finishing repeat regions automatically with Dupfinisher. In *Proceedings of the 2006 International Conference on Bioinformatics and Computational Biology*, pp. 141–146. Edited by H. R. Arabnia & H. Valafar. CSREA Press.

Hawkins, A. N., Johnson, K. W. & Bräuer, S. L. (2014). Southern Appalachian peatlands support high archaeal diversity. *Microb Ecol* 67, 587–602.

Høj, L., Olsen, R. A. & Torsvik, V. L. (2005). Archaeal communities in High Arctic wetlands at Spitsbergen, Norway (78°N) as characterized by 16S rRNA gene fingerprinting. *FEMS Microbiol Ecol* 53, 89–101.

Hu, G.-B., Rice, W. J., Dröse, S., Altendorf, K. & Stokes, D. L. (2008). Three-dimensional structure of the KdpFABC complex of *Escherichia col*i by electron tomography of two-dimensional crystals. *J Struct Biol* 161, 411–418.

Inatomi, K., Kamagata, Y. & Nakamura, K. (1993). Membrane ATPase from the aceticlastic methanogen *Methanothrix thermophila*. *J Bacteriol* 175, 80–84.

Jarrell, K. F. & Kalmokoff, M. L. (1988). Nutritional requirements of the methanogenic rchaebacteria. *Can J Microbiol* 34, 557–576.

Juottonen, H., Galand, P. E., Tuittila, E. S., Laine, J., Fritze, H. & Yrjälä, K. (2005). Methanogen communities and bacteria along an ecohydrological gradient in a northern raised bog complex. *Environ Microbiol* 7, 1547–1557.

Kotsyurbenko, O. R. (2010). Soil, wetlands, peat. In *Handbook of Hydrocarbon and Lipid Microbiology*, pp. 626–634. Edited by K. N. Timmis. Berlin: Springer-Verlag.

Kotsyurbenko, O. R., Friedrich, M. W., Simankova, M. V., Nozhevnikova, A. N., Golyshin, P. N., Timmis, K. N. & Conrad, R. (2007). Shift from acetoclastic to H_2 -dependent methanogenesis in a West Siberian

peat bog at low pH values and isolation of an acidophilic *Methanobacterium* strain. *Appl Environ Microbiol* **73**, 2344–2348.

Larkin, M. A., Blackshields, G., Brown, N. P., Chenna, R., McGettigan, P. A., McWilliam, H., Valentin, F., Wallace, I. M., Wilm, A. & other authors (2007). Clustal W and Clustal X version 2.0. *Bioinformatics* 23, 2947–2948.

Lin, X., Kennedy, D., Fredrickson, J., Bjornstad, B. & Konopka, A. (2012). Vertical stratification of subsurface microbial community composition across geological formations at the Hanford Site. *Environ Microbiol* 14, 414–425.

Markowitz, V. M., Korzeniewski, F., Palaniappan, K., Szeto, E., Werner, G., Padki, A., Zhao, X., Dubchak, I., Hugenholtz, P. & other authors (2006). The integrated microbial genomes (IMG) system. *Nucleic Acids Res* 34, D344–D348.

McMillan, D. G. G., Ferguson, S. A., Dey, D., Schröder, K., Aung, H. L., Carbone, V., Attwood, G. T., Ronimus, R. S., Meier, T. & other authors (2011). A₁A₀-ATP synthase of *Methanobrevibacter ruminantium* couples sodium ions for ATP synthesis under physiological conditions. *J Biol Chem* 286, 39882–39892.

Morsomme, P., Chami, M., Marco, S., Nader, J., Ketchum, K. A., Goffeau, A. & Rigaud, J.-L. (2002). Characterization of a hyperthermophilic P-type ATPase from *Methanococcus jannaschii* expressed in yeast. J Biol Chem 277, 29608–29616.

Mulkidjanian, A. Y., Galperin, M. Y., Makarova, K. S., Wolf, Y. I. & Koonin, E. V. (2008). Evolutionary primacy of sodium bioenergetics. *Biol Direct* 3, 13.

Müller, V., Ruppert, C. & Lemker, T. (1999). Structure and function of the A₁A₀-ATPases from methanogenic Archaea. *J Bioenerg Biomembr* **31**, 15–27.

Nakashima, K., Sugiura, A., Momoi, H. & Mizuno, T. (1992). Phosphotransfer signal transduction between two regulatory factors involved in the osmoregulated *kdp* operon in *Escherichia coli*. *Mol Microbiol* **6**, 1777–1784.

Pisa, K. Y., Weidner, C., Maischak, H., Kavermann, H. & Müller, V. (2007). The coupling ion in the methanoarchaeal ATP synthases: H^+ vs. Na⁺ in the A₁A₀ ATP synthase from the archaeon *Methanosarcina mazei* Gö1. *FEMS Microbiol Lett* 277, 56–63.

Podell, S. & Gaasterland, T. (2007). DarkHorse: a method for genomewide prediction of horizontal gene transfer. *Genome Biol* 8, R16.

Poolman, B. & Glaasker, E. (1998). Regulation of compatible solute accumulation in bacteria. *Mol Microbiol* 29, 397–407.

Rhoads, D. B., Waters, F. B. & Epstein, W. (1976). Cation transport in *Escherichia coli*. VIII. Potassium transport mutants. *J Gen Physiol* 67, 325–341.

Sakai, S., Takaki, Y., Shimamura, S., Sekine, M., Tajima, T., Kosugi, H., Ichikawa, N., Tasumi, E., Hiraki, A. T. & other authors (2011). Genome sequence of a mesophilic hydrogenotrophic methanogen *Methanocella paludicola*, the first cultivated representative of the order *Methanocellales*. *PLoS One* **6**, e22898.

Sakai, S., Ehara, M., Tseng, I.-C., Yamaguchi, T., Bräuer, S. L., Cadillo-Quiroz, H., Zinder, S. H. & Imachi, H. (2012). *Methanolinea mesophila* sp. nov., a hydrogenotrophic methanogen isolated from rice field soil, and proposal of the archaeal family *Methanoregulaceae* fam. nov. within the order *Methanomicrobiales*. *Int J Syst Evol Microbiol* **62**, 1389–1395.

Schlegel, K. & Müller, V. (2013). Evolution of Na⁺ and H⁺ bioenergetics in methanogenic archaea. *Biochem Soc Trans* **41**, 421–426.

Schlegel, K., Leone, V., Faraldo-Gómez, J. D. & Müller, V. (2012). Promiscuous archaeal ATP synthase concurrently coupled to Na⁺ and H⁺ translocation. *Proc Natl Acad Sci U S A* **109**, 947–952. Schönheit, P. & Perski, H. J. (1983). ATP synthesis driven by a potassium diffusion potential in *Methanobacterium thermoautotrophicum* is stimulated by sodium. *FEMS Microbiol Lett* **20**, 263–267.

Schrader, M., Fendler, K., Bamberg, E., Gassel, M., Epstein, W., Altendorf, K. & Dröse, S. (2000). Replacement of glycine 232 by aspartic acid in the KdpA subunit broadens the ion specificity of the K⁺-translocating KdpFABC complex. *Biophys J* **79**, 802–813.

Tatusov, R. L., Galperin, M. Y., Natale, D. A. & Koonin, E. V. (2000). The COG database: a tool for genome-scale analysis of protein functions and evolution. *Nucleic Acids Res* 28, 33–36.

Thauer, R. K., Kaster, A. K., Seedorf, H., Buckel, W. & Hedderich, R. (2008). Methanogenic archaea: ecologically relevant differences in energy conservation. *Nat Rev Microbiol* 6, 579–591.

van der Laan, M., Gassel, M. & Altendorf, K. (2002). Characterization of amino acid substitutions in KdpA, the K⁺-binding and -translocating

subunit of the KdpFABC complex of *Escherichia coli*. J Bacteriol 184, 5491–5494.

Wheeler, D. L., Barrett, T., Benson, D. A., Bryant, S. H., Canese, K., Chetvernin, V., Church, D. M., DiCuccio, M., Edgar, R. & other authors (2007). Database resources of the National Center for Biotechnology Information. *Nucleic Acids Res* 35, D5–D12.

Wüst, P. K., Horn, M. A. & Drake, H. L. (2009). Trophic links between fermenters and methanogens in a moderately acidic fen soil. *Environ Microbiol* 11, 1395–1409.

Yavitt, J. B., Basiliko, N., Turetsky, M. R. & Hay, A. G. (2006). Methanogenesis and methanogen diversity in three peatland types of the discontinuous permafrost zone, boreal western continental Canada. *Geomicrobiol J* 23, 641–651.

Edited by: R. Parales