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## Complete Genome Sequence of *Methanothermobacter marburgensis*, a Methanoarchaeon Model Organism<sup>∇</sup>

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The circular genome sequence of the chemolithoautotrophic euryarchaeon *Methanothermobacter marburgensis*, with 1,639,135 bp, was determined and compared with that of *Methanothermobacter thermautotrophicus*. The genomes of the two model methanogens differ substantially in protein coding sequences, in insertion sequence (IS)-like elements, and in clustered regularly interspaced short palindromic repeats (CRISPR) loci.

Methanothermobacter marburgensis (DSM 2133) (formerly Methanobacterium thermoautotrophicum strain Marburg), a member of the Methanobacteriales (2), was isolated in 1978 from anaerobic sewage sludge in Marburg, Germany (5). The hydrogenotrophic methanogen grows even faster (2 h versus 3 h doubling time) and to higher cell concentrations (3 g versus 1.5 g dry mass per liter) than Methanothermobacter thermautotrophicus (DSM 1053) (formerly Methanobacterium thermoautotrophicum strain  $\Delta$ H) (20) (for other differences, see references 3 and 19). Both methanogens were used in the last 35 years for the elucidation of the enzymes and coenzymes involved in CO<sub>2</sub> reduction to methane with H<sub>2</sub> (4, 16–18). The genome sequence of M. thermautotrophicus was reported in 1997 (15); that of M. marburgensis is announced here.

The genome size of M. marburgensis is 1,639,135 bp (that of M. thermautotrophicus is 1,751,377 bp), the genome G+C content is 48.64% (49.54% for M. thermautotrophicus), and the part coding is 90.94% (91.02% for M. thermautotrophicus). Comparison of the sequences (13) revealed that the two genomes have 1,607 protein coding sequences (CDS) in common and 411 CDS not in common (145 CDS are found only in M. marburgensis and 266 CDS only in M. thermautotrophicus) and show a high degree of synteny. The CDS not in common could be traced back to gene splitting (15%), gene deletion (30%), gene duplication (30%), and lateral gene transfer (24%) events (percentages given are for M. marburgensis). Of the 1,607 CDS in common, approximately 40% show BLAST search expectation values of  $>10^{-100}$  at the protein level, reflecting large differences in sequence divergence. Almost 470 CDS encode conserved hypothetical proteins.

The genome of *M. marburgensis* harbors 15 insertion sequence (IS)-like elements, whereas there is no evidence for a classically organized IS-like element in *M. thermautotrophicus*. Consistently, a CDS for a transposase is found only in *M. marburgensis*.

In the genome of *M. marburgensis* there is only one clustered regularly interspaced short palindromic repeat (CRISPR) locus with 36 repeats and only one CRISPR-associated (cas) gene (csa3), indicating that the organism is not protected from invasion by phage and plasmid DNA (7, 8, 10, 12). By comparison, in the genome of M. thermautotrophicus there are three CRISPR loci with 124, 4, and 47 repeats and 18 cas genes that encode proteins involved in adaptation and interference (http://genoweb1.irisa.fr/Serveur-GPO/outils/repeatsAnalysis /CRISPR/). The spacer sequences from locus 2 match DNA sequences found in phage  $\Psi M1$  of M. marburgensis (6, 11) and ΨM100 of M. wolfei (9), which supports the observation that M. thermautotrophicus is not lysed by those two phages. Unfortunately, there is no DNA sequence available for phage  $\Phi$ F1, which is able to lyse *M. thermautotrophicus* (14), to compare it with the spacer sequences of the CRISPR regions. In the plasmid pM2001 (= pMTBMA4) (4,439-bp circular multicopy plasmid found only in M. marburgensis) (1, 19), no sequence identities for CRISPR spacer sequences of M. thermautotrophicus were found (14).

Approximately 200 CDS were identified that are required for the synthesis of the enzymes, coenzymes, and prosthetic groups involved in  $\mathrm{CO}_2$  reduction to methane and in the coupling of this process with energy conservation. Some of the genes have been found only recently; others, such as those for coenzyme  $\mathrm{F}_{430}$  biosynthesis, still remain to be discovered.

**Nucleotide sequence accession number.** The complete genome sequence of *M. marburgensis* was deposited in GenBank under accession numbers CP001710 (chromosome) and CP001711 (pMTBMA4).

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