A Cultured Greigite-Producing Magnetotactic Bacterium in a Novel Group of Sulfate-Reducing Bacteria

Christopher T. Lefèvre, Nicolas Menguy, Fernanda Abreu, Ulysses Lins, Mihály Pósfai, Tanya Prozorov, David Pignol, Richard B. Frankel, Dennis A. Bazyliński

Magnetotactic bacteria contain magnetosomes—intracellular, membrane-bounded, magnetic nanocrystals of magnetite (Fe₃O₄) or greigite (Fe₃S₄)—that cause the bacteria to swim along geomagnetic field lines. We isolated a greigite-producing magnetotactic bacterium from a brackish spring in Death Valley National Park, California, USA, strain BW-1, that is able to biomineralize greigite and magnetite depending on culture conditions. A phylogenetic comparison of BW-1 and similar uncultured greigite- and/or magnetite-producing magnetotactic bacteria from freshwater to hypersaline habitats shows that these organisms represent a previously unknown group of sulfate-reducing bacteria in the Deltaproteobacteria. Genomic analysis of BW-1 reveals the presence of two different magnetosome gene clusters, suggesting that one may be responsible for greigite biomineralization and the other for magnetite.

Several different groups of magnetotactic bacteria biomineralize magnetic mineral nanocrystals of magnetite (Fe₃O₄) and greigite (Fe₃S₄) through fundamentally different mechanisms (1). A number of magnetite-producing magnetotactic bacteria have been isolated and their genomes sequenced, revealing details about magnetite biomineralization at the molecular level (2–4), but no greigite-producing bacteria have been isolated and grown in axenic culture. There are two known morphological types of greigite-producers: a group of multicellular prokaryotes [“many-celled magnetotactic prokaryotes” (MMPs)], and a group of large rod-shaped bacteria (5). Although environmental studies revealed a good deal of information regarding MMPs (6–11), little is known about the large rod-shaped group except that some appear to produce magnetite as well as greigite in the same cell (12–14). One report describes a putative large, greigite-producing rod belonging to the Gammaproteobacteria (8), although its affiliation with this group is uncertain (15).

Here, we report a comprehensive phylogenetic analysis of large, rod-shaped, greigite- and/or magnetite-producing magnetotactic bacteria and the isolation and characterization of one of these microorganisms, strain BW-1, from a brackish spring at Badwater Basin, Death Valley National Park, California, USA. We collected water and sediment samples from numerous freshwater, brackish, and hypersaline aquatic environments located in the southwestern United States (table S1) (16).

Although the magnetotactic bacterial populations in the samples were not homogeneous, the large rods were often the dominant magnetotactic bacterium present in the samples, as determined with microscopy by using the hanging drop technique (movie S1) (17). To obtain cells for analyses, magnetotactic bacteria were first magnetically enriched in sample bottles (18) and then purified using the magnetic racetrack procedure (19). After purification, cells were used to inoculate a variety of different growth media for electron microscopy, phylogenetic analysis, and fluorescence in situ hybridization.

We observed a wide range of large magnetotactic rods collected from different freshwater and saline environments (Fig. 1). In all cases, cells possessed a single polar flagellum (Fig. 1, A and D). Some cells contained mainly greigite (Fig. 1, A to C) or magnetite (Fig. 1, D to F), whereas others produced both minerals (Fig. 1, G to I, and table S1), as determined by means of selected-area electron diffraction (SAED) in the electron microscope. Although the morphologies of the greigite crystals were generally pleomorphic, in all cases the magnetite crystals were bullet- or arrowhead-shaped [elongated anisotropic (20)] (Fig. 1, E, F, H, and I). Using bacterial-specific primers for the 16S ribosomal RNA (rRNA)
Fig. 2. Fluorescent in situ hybridization (FISH) of large, greigite- and/or magnetite-producing, rod-shaped magnetotactic bacteria (LGRs) to the specific oligonucleotide rRNA probe LGRp. (A) FISH of LGR cells collected from Badwater Basin, Death Valley National Park, California, USA. (B) FISH of LGR cells collected from Rogers Spring at the Lake Mead National Recreation Area, Nevada, USA. (Left) Differential interference contrast (DIC) images of magnetically enriched cells from water and sediment samples. (Middle) Fluorescence microscope images of the same cells stained with 4',6-diamidino-2-phenylindole (DAPI). (Right) Fluorescence microscope images of the same cells hybridized with the LGR-specific Alexa594-labeled probe LGRp (5'-GCTGCCCTTGTACATAC-3'), complementary to nucleotides close to the 1260-base pair region of the 16S rRNA molecule of the different LGRs. Cells at arrows are not LGRs and serve here as negative controls because only LGRs cells fluoresced with this probe.

Fig. 3. Phylogenetic tree, based on 16S rRNA gene sequences, showing the phylogenetic position of LGRs in the Deltaproteobacteria class. Bootstrap values at nodes are percentages of 1000 replicates. 16S rRNA gene sequences of the uncultured magnetotactic bacteria Candidatus Thermomagnetovibrio puitensis (28) and Ca. Magnetovum mohavensis (29) of the Nitrospirae phylum were used to root the tree. GenBank accession nos. are given in parentheses. Bar represents 2% sequence divergence. Letters in sequence designations indicate sampling site listed in table S1, and numbers following represent clone number. BP, Bridgeport Reservoir; LK, Lake Mead; NML, freshwater pond near Mono Lake; ZZ, Lake Tuendae; ZP, freshwater pond near Zion National Park; ZBP, Zuma Beach; RG, Rogers Spring; GL, pools at Great Boiling Springs; PL, Pyramid Lake; SS, Salton Sea; SS-2c, lagoon at Salton Sea; BB, Badwater Basin; and WH, Salt Pond. Details and exact coordinates are provided in table S1. BW-1 refers to the cultured LGR described in the text.
to this bacterium (15). The recent discovery of magnetite-producing rods that belong to the **Gammaproteobacteria** and are found in the same habitats as rod-shaped greigite-producers (18) might also explain this discrepancy.

Magnetically purified, large, rod-shaped, magnetotactic bacteria from several sites were inoculated into a variety of growth media. Growth of some of these organisms was observed in an anaerobic, complex, liquid growth medium for the enrichment of heterotrophic sulfate-reducing bacteria. Through three successive rounds of dilution to extinction in this medium, two pure cultures were obtained: strain BW-1 from Badwater Basin and strain SS-2 from the Salton Sea (California, USA). Both strains phylogenetically lie within the clade of the uncultured large rods in the **Deltaproteobacteria** class (Fig. 3). Strain SS-2 appeared to only biominalize magnetite under our culture conditions.

In culture, BW-1 produces one or two loose chains of magnetosomes that contain greigite and/or magnetite, as determined through a combination of elemental mapping by using energy-filtered imaging (Figs. S1 and S2) and electron diffraction (Fig. 4, A to C, and Figs. S2C and S3B). Magnetosome mineral composition correlated with the concentration of hydrogen sulfide formed during growth. For example, when sulfide was allowed to accumulate in the growth medium (>0.3 mM sulfide) most magnetosomes contained greigite (Fig. 4, and Figs. S1 and S2).

When the headspace of the cultures was purged every other day with oxygen-free argon gas so as to decrease the concentration of hydrogen sulfide in the cultures (<0.3 mM), cells contained chains of bullet-shaped, magnetite-containing magnetosomes (Fig. S3). Thus, whether greigite and/or magnetite are produced in the magnetosomes of these organisms is affected by external environmental conditions such as hydrogen sulfide concentration and/or redox potential. This finding is consistent with observations on uncultured rod-shaped, greigite-producing magnetotactic bacteria in a chemically stratified, oxygen/sulfide inverse gradient system (13).

Magnetite crystals produced by BW-1 were consistently bullet-shaped, whereas greigite crystals had irregular outlines and lacked a well-defined crystal habit (Fig. 4, B and C). Thin sections of greigite magnetosomes reveal an electron-dense layer surrounding the crystal that is consistent with the presence of a magnetosome membrane (Fig. 4, E and F), which has also been observed in greigite magnetosomes of the MMP **Candidatus Magnetoglobus multicellularis** (23).

Our sampling effort demonstrates that greigite-producing magnetotactic bacteria are not confined to marine habitats as previously thought (8). Although several of these organisms collected from different sites appeared to only contain magnetite, it seems likely that all can biominalize greigite under the appropriate conditions (such as when sulfite is present as the terminal electron acceptor and is reduced to sulfide), on the basis of evidence presented here and on the fact that environmental conditions are thought to have an effect on the magnetosome mineral produced in these organisms (13). Although they differ morphologically, there are several similarities between the large rods and the greigite-producing MMP group. These obligately multicellular organisms consist of 10 to 60 genetically identical cells and are also phylogenetically affiliated with the **Deltaproteobacteria** class of the **Proteobacteria** phylum (9–11, 24).

Some are known to biominalize magnetite as well as greigite (25), and based on phylogenetic and genetic evidence, the MMPs appear to be anaerobic, sulfate-reducing bacteria (9–11, 24).

Recently, magnetosome (mam) genes, similar to those in magnetite-producing magnetotactic bacteria, have been found in an MMP that produced only greigite, leading to the idea that the magnetotactic trait, whether based on iron oxide or iron sulfide production, is monophyletic (26). The fact that magnetite and/or greigite production can be controlled in cells of a single species by modifying the external chemistry and redox potential suggests that there might be specific genes involved in greigite production in some large rods and MMPs that are expressed in certain conditions. We identified mam genes in the genome of strain BW-1 that are mostly present as two copies (table S2) and appear to form two separate clusters (fig. S5). The first cluster contains genes encoding for magnetosome proteins most closely related to those of the magnetite-producer **Desulfovibrio magneticus** strain RS-1 (27). The second cluster contains genes encoding for proteins most closely related to those of the greigite-producer **Candidatus Magnetoglobus multicellularis** (26). These results point to the possibility that different forms of mam genes are responsible for magnetite (cluster one) and greigite (cluster two) biominalization (fig. S5) and that the two gene clusters are regulated differentially, resulting in different proportions of magnetite and greigite in cells.

**Fig. 4.** TEM images of strain BW-1. (A) Darkfield scanning TEM image of a magnetosome chain containing both greigite (labeled B) and magnetite (labeled C). (B) High-magnification TEM image of greigite crystal labeled in (A). (Inset) SAED of crystal viewed along the [0 1 1] zone axis. Pattern is consistent with greigite. (C) High-magnification TEM image of magnetite crystal labeled in (A). (Inset) SAED pattern of crystal viewed along the [1 1 0] zone axis. (D) TEM image of a stained thin-section of a cell of BW-1 showing several magnetosomes aligned in the cell. Dark, intracellular, electron-dense mass represents large cell inclusion that is also visible by use of light microscopy. (E) High-magnification TEM image of magnetosomes in (D) showing that an electron-dense layer surrounds the greigite crystals, suggesting the presence of a magnetosome membrane.

**References and Notes**

Materials and methods are available as supporting material online.

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Supporting Online Material
www.sciencemag.org/cgi/content/full/334/6063/1720/DC1
Materials and Methods
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Movies S1 and S2
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