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Search for Past Life on Mars: Possible Relict Biogenic Activity in Martian Meteorite ALH84001

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Fresh fracture surfaces of the martian meteorite ALH84001 contain abundant polycyclic aromatic hydrocarbons (PAHs). These fresh fracture surfaces also display carbonate globules. Contamination studies suggest the PAHs are indigenous to the meteorite. High resolution scanning and transmission electron microscopy study of surface textures and internal structures of selected carbonate globules show that the globules contain finegrained, secondary phases of single-domain magnetite and Fe-monosulfides. The carbonate globules are similar in texture and size to some terrestrial bacterially induced carbonate precipitates. Although inorganic formation is possible, formation of the globules ules by biogenic processes could explain many of the observed features including the PAHs. The PAHs, the carbonate globules, and their associated secondary mineral phases of a past martian biota.

A long-standing debate over the possibility of present-day life on Mars was addressed by the Viking lander experiments in 1976. Although the results were generally interpreted to be negative for life in the tested surface soils, the possibility of life at other locations on Mars could not be ruled out (1). The Viking lander's mass spectrometry experiments failed to contirm the existence of organics for the martian surface samples analyzed. Furthermore, the Viking results contained no information on possible fossils. Another source of information about possible ancient martian life is the Shergotty-Nakhla-Chassigny (SNC) class of meteorites, which appear to have come to the Earth by impact events on Mars (2, 3). We have examined ALH84001 collected in Antarctica and recently recognized as a meteorite from Mars (4). Our objective was to look for signs of past (fossil) life within the pore space or secondary minerals of the martian meteorites. Our task is difficult because we only have a small piece of rock from Mars and we are searching for martian

biomarkers on the basis of what we know about life on Earth. Therefore, if there is a martian biomarker, we may not be able to recognize it, unless it is similar to an earthly biomarker. Additionally, no information is available on the geologic context of the rock on Mars.

ALH84001 is an igneous orthopyroxenite consisting of coarse-grained orthopyroxene {(Mg,Fe)SiO1) and minor maskelynite (NaAlSi,O.), olivine [(Mg,Fe)SiO.], chromite (FeCr.O.), pyrite (FeS.), and apatite [Ca, (PO4)2] (4-6). Its crystallized 4.5 billion years ago (Ga) (6). It records at least two shock events separated by a period of annealing. The age of the first shock event has been estimated to be 4.0 Ga (7). Unlike the other SNC meteorites, ALH84001 contains secondary carbonate minerals that form globules from 1 to \sim 250 μ m across (4, 6, 8, 9). These carbonate globules have been estimated to have formed 3.6 Ga (10). Petrographic and electron microprobe results (4, 11) indicate that the carbonates formed at relatively high temperatures (~700°C), however, the stable oxygen isotope data indicate that the carbonates formed between 0° and 80°C (12). The carbonate globules are tound along tractures and in pore spaces. Some of the carbonate globules were shock-faulted (4, 5). This shock event occurred on Mars or in space. and thus rules out a terrestrial origin for the globules (3, 8, 13). The isotopic composition of the carbon and oxygen associated with the carbonate giobules also indicates that they are indigenous to the meteorite

and were not formed during its 13.000-year residence in the Antarctic environment (13).

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The δ^{13} C values of the carbonate in ALH84001 range up to 42 per ml for the large carbonate spheroids (12) and are higher than values for carbonates in other SNC meteorites. The source of the carbon is the martian atmospheric CO2, which has been recycled through water into the carbonate (12). The carbon isotopic compositions of ALH84001 are similar to those measured in CM2 carbonaceous chondrites (14). Consequently, the carbonates in ALH84001 and the CM2 meteorites are believed to have been formed by aqueous processes on parent bodies. The $\delta^{13}C$ in martian meteorite carbonates ranges from -17 to +42‰ (12. 15. 16). This range of ¹³C valves exceeds the range of ¹³C generated by most terrestrial inorganic processes (17, 18). Alternatively, biogenic processes are known to produce wide ranges in ¹³C fractionations on Earth (19, 20).

ALH84001 arrived on Earth 13.000 years ago and appears to be essentially free of terrestrial weathering (8). ALH84001 does not have the carbon isotopic compositions typically associated with weathered meteorites (12, 15) and detailed mineralogical studies (8) show that ALH84001 has not been significantly affected by terrestrial weathering processes.

ALH84001 is somewhat friable and breaks relatively easily along preexisting fractures. It is these fracture surfaces that display the carbonate globules. We analyzed freshly broken fracture surfaces on small chips of ALH84001 for polycyclic aromatic hydrocarbons (PAHs) using a microprobe two-step laser mass spectrometer (μ L²MS) (22, 23).

Polycyclic aromatic hydrocarbons. Spatial distribution maps of individual PAHs on interior fracture surfaces of ALH84001 demonstrate that both total PAH abundance and the relative intensities of individual species have a heterogeneous distribution at the 50- μ m scale. This distribution appears to be consistent with partial geochromatographic mobilization of the PAHs (24). The average PAH concentration in the interior fracture surfaces is estimated to be in excess of 1 part per million (25). The PAHs were found in highest concentration in regions rich in carbonates.

From averaged spectra we identified two groupings of PAHs by mass (Fig. 1A). A middle-mass envelope of 178 to 276 atomic mass units (amu), dominates and is composed mostly of simple 3- to 6-ring PAH skeletons with alkylated homoiogs accounting for less than 10% of the total integrated signal intensity. Principal peaks at 178, 202. 228, 252, and 278 amu are assigned to

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phenanthrene $(C_{14}H_{10})$, pyrene $(C_{16}H_{10})$, chrysene $(C_{18}H_{12})$, perylene or benzopyrene $(C_{20}H_{12})$, and anthanthracene $(C_{22}H_{12})$ (26). A second weak, diffuse high-mass envelope extends from about 300 to beyond 450 amu. The peak density is high and shows a periodicity at 14 and 2 amu. This distribution implies that there is a complex mixture of PAHs whose parent skeletons have alkylated side chains with varying degrees of dehydrogenation; specific assignments are ambiguous.

Contamination checks and control experiments indicate that the observed organic material is indigenous to ALH84001. The accumulation of PAHs on the Greenland ice sheet over the past 400 years has been studied in ice cores (27). The total concentration of PAHs in the cores varies from 10 parts per trillion for preindustrial times to 1 part per billion for recent snow deposition. Because Antarctica is in the less industrialized Southern Hemisphere we may expect concentrations of PAHs in Antarctic ice to lie between these two limits. The primary source of PAHs is anthropogenic emissions, which are characterized by extensive alkylation (~10 fold greater than that of the parent PAHs) (28) and by the presence of abundant aromatic heteroprimarily dibenzothiophene cycles. (C12H8S; 184 amu). In contrast, the PAHs in ALH84001 are present at the part per million level ($\sim 10^3$ to 10^5 times higher concentration), show little alkylation, and dibenzothiophene was not observed in any of the samples we studied.

Analysis of Antarctic salt deposits on a heavily weathered meteorite (LEW 85320) by μL^2MS did not show the presence of

terrestrial PAHs within detection limits. which suggests an upper limit for terrestrial contamination of ALH84001 of 1%. Measurements of four interior fragments of two Antarctic ordinary chondrites (ALH83013 and ALH83101) of petrologic classes H6 and L6 showed no evidence of indigenous PAHs. These represent equivalent desorption matrix blanks; previous studies have shown that no indigenous organic material is present in meteorites of petrologic class six (29).

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Spatially resolved studies of exterior fragments of ALH84001, with intact fusion crust show that no PAHs are present within the fusion crust or a zone extending into the interior of the meteorite to a depth of ~500 µm (Fig. 1, B-E). The PAH signal increases with increasing depth, leveling off at ~ 1200 µm within the interior, well away from the fusion crust. This concentration profile is consistent with volatilization and pyrolysis of indigenous PAHs during atmospheric entry of the meteorite and formation of a fusion crust (30) but inconsistent with terrestrial introduction of organic material into the interior of ALH84001 along cracks and pore spaces during burial in the Antarctic ice sheet. These results indicate that the PAHs are indigenous to ALH84001.

No evidence can be found for laboratorybased contamination introduced during processing. Samples for analysis were prepared at the meteorite clean labs at NASA Johnson Space Center and sealed in containers before they were transported to Stanford University. A contamination study conducted prior to analysis of these samples showed no evidence for any PAH contamination (31). We also conducted experiments in

which chips of ALH84001 were cultured in nutrient measure under aerobic and anerobic conditions; we found the chips to be sterile.

PAHs have been found in a wide range of extraterrestial materials using the μL^2MS technique, is cluding carbonaceous and ordinary chor stites (29), interplanetary dust 32) and interstellar graphite particles (2 grains (33). Each material is characterized ev differing PAH distributions reflecting the differer environments in which the PAHs form and their subsequent evolution (for example, as a result of aqueous alteration and thermal metamorphism). Comparison of the mass distribution of PAHs observed in ALH84001 with that of PAHs in other extraterrestrial materials indicates that the closest match is with the CM2 carbonaceous chondrites (34). The PAHs in ALH84001, however. differ in several respects from the CM2 chondrites: low-mass PAHs such as naphthalene (C10Hs: 123 amu) and acenaphthalene (C12H2; 152 amu) are absent in ALH84001, the middle-mass envelope shows no alkylation, and the relative intensity of the 5- and 6-ring PAHs and the relative intensity and complexity of the extended high-mass distribution are different.

On Each, PAHs are abundant as fossil molecules in ancient sedimentary rocks, coal, and petroleum where they are derived from chemical aromatization of biological precursors such as marine plankton and early plant life (35). In such samples, PAHs are typically present as thousands, if not hundreds of thousands, of homologous and isomeric series: in contrast, the PAHs we observed in ALH84001 appear to be relatively

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Fig. 1. (A) Averaged mass spectrum of an intenor, carbonate-rich, fracture surface of ALH84001. The spectrum represents the average of 1280 individual spectra defining an analyzed surface region of 750 by 750 μm mapped at a spatial resolution of 50 by 50 μm. (B-E) PAH Signal intensity as a function

of distance from the AL+84001 fusion crust for the four binmary PAHs shown in (A). The fusion crust fragment, we can showed no preexisting fractures, was cleaved immediately prior to analysis using a stainless steel scapel and introduced in <2 minutes into the μ L*1.0 Each plot represents a section perpendicular to the fusion crust surface which starts at the extension and extension distance of 1200 μ m inward. The sub-threshold to μ the analysis shot being the summed average of 5 time-of-flight spectra.

Fig. 2. Faise-color backscatter electron (BSE) image of fractured surface of a chip from ALH84001 meteorite showing distribution of the carbonate ! giobules. Orthopyroxene s green and the carponate ciopules are orange. Surrounding the i Mg-carbonate are a i black rim (magnesite) and a white, Fe-rich rim. Scale bar is 1 mm. (False color produced by C. Schwanot).



simple. The *in situ* chemical aromatization of naturally occurring biological cyclic compounds in early diagenesis can produce a restricted number of PAHs (36). Hence, we would expect that diagenesis of microorganisms on ALH84001 could produce what we observed—a few specific PAHs—rather than a complex mixture involving alkylated homologs.

Chemistry and mineralogy of the carbonates. The freshly broken, but preexisting fracture surfaces rich in PAHs also typically display carbonate globules. The globules tend to be discoid rather than spherical, and are flattened parallel to the fracture surface. Intact carbonate globules appear orange in visible light, have a rounded appearance, and many display alternating black and white rims. Under high magnification stereo light microscopy or SEM stereo imaging, some of the globules appear to be quite thin and pancake-like, suggesting that the carbonates formed in the restricted width of a thin fracture. This geometry limited their growth perpendicular to but not parallel to the fracture.

We selected a typical globule, $\sim 50 \,\mu\text{m}$ in diameter, for analysis by TEM and electron microprobe (37). On the basis of backscatter electron (BSE) images (Fig. 2), the

Fig. 3. BSE image and microprobe electron maps showing the concentration of five elements in a carbonate from ALH84001. The element maps show that the carchemically bonate is zoned. Colors range through red, green, light blue, and deep blue. reflecting the highest to lowest element concentrations. Scale bars for all images are 20 µm. (A) BSE image snowing location of orthopyroxene (OPX), clinopyroxene (CPX), apatite (A), and carbonate (MgC, C). Fe-nch nms (R) separate the center of the carbonate (C) from a Monon carbonate (MgC) rim. Region in the box is described in Figs. 5 and 6. (B) Felis most abundant in



the barallelinms. \sim 3 μ m across, and in a region of the carbonate \sim 20 μ m in size. (C) Highest S is associated with an Fe-rich nim; it is not homogeneously distributed, but rather located in discrete 3 regions or hot spots in the nm. A lower S abundance is present throughout the globule in batchy areas. (D) Higher concentrations of Mg are shown in the Fe-rich nums. (E) Ca-rich region are associated with the abatite, the Fe-rich core of the carbonate and the clinopyroxene (F). P-rich regions are associated with the abatite.

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larger globules (>10 μ m) have Ca-ric cores (which also contain the highest M abundances) surrounded by alternating Fe and Mg-rich bands (Fig. 3). Near the edge of the globule, several sharp thin bands are present. The first band is rich in Fe and S the second is rich in Mg with no Fe, ththird is rich in Fe and S again (Fig. 3 Detectable S is also present in patchy area throughout the globule.

In situ TEM analyses of the globule in Fig. 4 revealed that Fe- and Mg-rich car bonates located nearer to the rim range u composition from ferroan magnesite to pure magnesite. The Fe-rich rims are composed mainly of fine-grained magnetite ranging ir size from ~10 to 100 nm and minor amounts of pyrrhotite (~5 vol%) (Figs. 3 region i and 4A). Magnetite crystals are cuboid, teardrop, and irregular in shape. Individual crystals have well-preserved structures with no lattice defects. The mag netite and Fe-sulfide are in a fine-grained carbonate matrix (Figs. 4A-4C). Composition of the fine-grained carbonate matrix matches that of coarse-grained carbonates located adjacent to the rim (Fig. 4A).

High-resolution transmission electron microscopy (HRTEM) and energy dispersive spectroscopy (EDS) showed that the Fe sulfide chase associated with the Fe-rich rims is pyrchotite (Fig. 4C). Pyrchotite par ticles are composed of S and Fe only, no oxygen was observed in the spectra; particle: have atomic Fe/S ratios ranging from ~0.92 to 0.97. The size and shape of the FeS particles varies. Single euhedral crystals of pyrrhoute range up to ~100 nm across polycrystalline particles have more rounded shapes ranging from ~20 to 60 nm acros. (Fig. 4C). HRTEM of these particles showed that their basal spacing is 0.57 nm, which corresponds to the {111} reflection of the pyrthoute in a 4C monoclinic system. The magacute is distributed uniformly in the rim, whereas the pyrrhotite seems to be dis tributed randomly in distinct domains ~5 to 10 pair long (Fig. 3C). Magnetite grains it AL184001 did not contain detectable amounts of minor elements. In addition these magnetite grains are single-domain create having no structural defects.

distinct region, located toward the connected the carbonate spheroid but computers separate from the magnetite-rich re-deterabed above, also shows accumula ton or magnetite and an Fe-rich sulfide (also Sector types of textures: The first on the sector types of textures: The first on the sector types and electron-dense unde to the massive and electron-dense unde to the massive and is porous. The porouteration occurs mainly in crosscuttin builts and rarely in isolated patches. We interest this porous texture as a region a which the massive carbonate has been partially dissolved. The nanometer-size magnetite and Fe-sulfide phases are everywhere associated with the fine-grained Mg-Fe-rich carbonate. In the regions containing high concentrations of magnetite, dissolution of carbonate is evident (Fig. 5A). In contrast to the magnetite-rich rim, the core area contains few magnetite particles. The Fesulfide phases in this magnetite-poor region have chemical compositions similar to that of the pyrrhotite. However, unlike pyrrhotite grains that have a large variety of morphologies, most of these Fe-sulfide particles have elongated shapes (Fig. 5B). We could not obtain a diffraction pattern of these Fe-monosulfide particles because they were unstable in the electron beam. Possible candidates for these Fe-monosulfide minerals include mackinawite (FeS_{1-x}), greigite (Fe₃S₄), and smythite (Fe₉S₁₁). Because of the morphological similarity to terrestrial greigite (Fig. 5C), we suggest these Fe sulfide minerals are probably greigite (38).

Formation of the magnetite and iron sulfides. The occurrence of the fine-grained carbonate. Fe-monosulfide, and magnetite phases could be explained by either inorganic or biogenic processes. Single-domain magnetite can precipitate inorganically under ambient temperature and neutral pH conditions by partial oxidation of ferrous

solutions (39). This synthetic magnetite ranges in size from about 1 to more than 100 nm and is chemically very pure (39). Simultaneous inorganic precipitation of magnetite and pyrthotite requires strongly reducing conditions at high pH (40). However, carbonate is normally stable at high pH, and the observed dissolution of carbonate would normally require low pH acidic conditions. It is possible that the Fe-sulfides, magnetite, and carbonates all formed under high pH conditions, and the acidity changed at some point to low pH causing the partial dissolution of the carbonates. But the Fe-sulfide and magnetite do not appear to have undergone any corrosion or dissolution (41), which would have likely occurred under acidic conditions. Moreover, as previously mentioned, the dissolution of carbonate is always intimately associated with the presence of Fe-sulfides and magnetite. Consequently, neither simultaneous precipitation of Fe-sulfides and magnetite along with dissolution of carbonates nor sequential dissolution of carbonate at a later time without concurrent dissolution of Fe-sulfides and magnetite seems plausible in simple inorganic models, although more complex models could be proposed.

In contrast, the coexistence of magnetite and Fe-sulfide phases within partially dissolved carbonate could be explained by bio-





Fig. 4. TEM images of a thin section obtained from part of the same fragment shown in Fig. 3A (from the region of arrow I, Fig. 3A). (A) Image at low magnification showing the Ferrich nm containing finegrain magnetite and Fe-sulfide phases and their association with the surrounding carbonate (C) and orthopyroxene (CPX). (B) High magnification of a magnetite-rich area in (A) showing the distribution of individual magnetite crystals inigh contrast) within the fine-grain carbonate (low contrast). (C) High magnification of a symmotite-rich region showing the distribution of individual pyrmotite particles itwo black arrows in the center) together with magnetite (other arrows) within the the-grained carbonate (low contrast).

genic processes, which are known to operate under extreme disequilibrium conditions. Intercellular coprecipitation of Fesulfides and magnetite within individual bacteria has been reported (42). In addition, extracellular biomediated precipitation of Fe-sulfides and magnetite can take place under anaerobic conditions (43, 44).

Magnetite particles in ALH84001 are similar (chemically, structurally, and morphologically) to terrestrial magnetite particles known as magnetofossils (45), which are tossil remains of bacterial magnetosomes (46) found in a variety of sediments and soils (41, 47, 48) and classified as singledomain (~20 to 100 nm) or superparamagnetic (<20 nm).(49). Single-domain magnetite has been reported in ancient limestones and interpreted as biogenic (48). Some of the magnetite crystals in the ALH84001 carbonates resemble extracellular precipitated superparamagnetic magnetite particles produced by the growth of anaerobic bacterium strain GS-15 (43).

Surface features and origin of the carbonates. We examined carbonate surfaces on a number of small chips of ALH84001 using high-resolution SEM (50). The Ferich rim of globules typically consist of an aggregate of tiny ovoids intermixed with small irregular to angular objects (Fig. 6A). Ovoids in the example are about 100 nm in longest dimension, and the irregular objects range from 20 to 80 nm across. These features are typical of those on the Fe-rich rims of many carbonate globules. These objects are similar in size and shape to features in the Fe-rich rims identified as magnetite and pyrrhotite (Figs. 4B-4C). These objects are too small to obtain compositional analysis under the SEM.

In the center of some of the globules (Fig. 2), the surface of the carbonate shows an irregular, grainy texture. This surface texture does not resemble either cieavage or a growth surface of synthetic and diagenetic carbonates (51). These surfaces also display small regularly shaped ovoid and elongated forms ranging from about 20 to 100 nm in longest dimension (Fig. 6B). Similar tex tures containing ovoids have been found or, the surface of calcite concretions growr from Pleistocene ground water in southerr Italy (52) where they are interpreted a nannobacteria that have assisted the calcite precipitation.

The origin of these textures on the sur face of the ALHS4001 carbonates (Fig-6A-6B) is unclear. One possible explantion is that the textures observed on the carbonate surface is a result of the partidissolution of the carbonate, that is they a erosional remnants of the carbonate the happen to be in the shape of ovoids a elongate forms perhaps because the carbonate to be in the shape of ovoids a elongate forms perhaps because the carbonate to be the the carbonate the

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ate has preferentially eroded along grain boundaries or dislocations. Shock effects may have enhanced such textures. However, as we know of no similar example from the terrestrial geologic record or from laboratory experiments, we cannot fully evaluate this possible explanation for the textures. A second possibility is that artifacts can be created during sample preparation or may result from laboratory contamination. For example, the application of a thick Au-conductive coating can produce textures resembling mud cracks, and even droplets or blobs of Au. Laboratory contamination can include dust grains, residue from sample cleaning, and organic contamination from epoxy. For comparison, we examined several control samples treated identically to the meteorite chips. We conclude that the complex textures (Fig. 6) did not result from procedures used in our laboratory. Only interior or freshiv broken surfaces of chips were used (52). We did observe an artifact texture from our Au-Pd conductive coating that consists of a mud crack-like texture visible only at 50,000× magnification or greater. None of the controls display concentrations or blobs of coating material. A lunar rock chip carried through the same procedures and examined at high magnification showed none of the features seen in Fig. 6.

An alternative explanation is that these textures as well as the nanosite magnetite



Fig. 5. TEM images of a thin section showing the morphology of the Fe-sulfide chases present in ALH84001 and a terrestrial soil sample. Iron sulfide phase (greigite?-Fe3Sa) is located in a magnetitepoor region separate and distinct from the magnetite-rich nms (Fig. 3A, arrow II). (A) TEM of a thin section showing a cross section of a single carbonate crystal (large black regions: The apparent cleavage features are due to knife damage by ultramicrotomy.) A vein of fine-grained carbonate (light grey) is observed between the large carbonate crystal. Possibly greigite and secondary magnetite (fine, dark crystals) have been precipitated in this fine-grained matrix. There is a direct relation between the degree of carbonate dissolution and the concentration of the fine-grained magnetite and Fe-sulfide phases. This region shows less alteration of carbonate and fewer Fe-rich particles, while regions shown in Fig. 4 contain carbonate that has been highly altered and contains abundant Felnch particles. The cleavage surface of the carbonate crystal does not show any dissolution features (arrows); there is no evidence of structural selective dissolution of carbonate. (B) a representative elongated Fe-sulfide particle, located in the dissolution region of the carbonate described in (A), is most likely composed of greigite. The morphology and chemical composition of these particles are similar to the plogenic greigite described in (C). (C) High magnification of an individual microorganism within a root cell of a soil sample showing an elongated, multicrystalline core of greigite within an organic envelope.



Fig. 6. High-resolution SEM images showing ovoid and elongate features associated with ALH84001 carbonate globules. (A) Surface of Fe-rich rim area. Numerous ovoids, approximately 100 nm in diameter, are present (arrows). Tubular-shaped bodies are also apparent (arrows). Smaller angular grains may be the magnetite and cyrmotite found by TEM. (B) Close view of central region of carbonate (away from nm areas) showing textured surface and nanometer ovoids and elongated forms (arrows).

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and Fe-sulfides, are the products of microl ological activity. It could be argued th these features in ALH84001 formed in Ara arctica by biogenic processes or inorgany weathering. It is unlikely that reduced phaes, such as iron sulfides, would form in Ari arctica during inorganic weathering; repor ed authigenic sulfur-bearing phases free Antarctic soils and meteorites are sulfates hydrated sulfates. In general, autnigenic s ondary minerals in Antarctica are oxidiz or hydrated (53). The lack of PAHs in the analyzed Antarctic meteorites, the sterilio of the sample, and the nearly unweather nature of ALH84001 argue against an Aarctic biogenic origin. As a control we amined three Antarctic ordinary chonduce (ALH78119, ALH76004, and ALH810// (which do not have indigenous PAHs) from the same ice field where ALH84001 wa collected, as well as a heavily weathereordinary chondrite that gave negative result for PAHs (LEW 85320). These meteories were chosen to cover the different degrees a weathering observed on Antarctic metricoites. Examination of grain surfaces ac at magnifications in weathered and unweath ered regions of these meteorites showed as sign of the ovoid and elongate forms seen ALH84001. However, none of these contra meteorites contained detectable carbonate

Ovoid features in Fig. 6 are similar in suand shape to nannobacteria in travectivand limestone (54). The elongate forms (Fig. 6B) resemble some forms of fossilizefilamentous bacteria in the terrestrial fossrecord. In general, the terrestrial bactermicrofossils (55) are more than an order of magnitude larger than the forms seen to the ALH84001 carbonates.

The carbonate globules in ALH3.87 are clearly a key element in the inter-or tion of this martian meteorite. The or. these globules is controversial; Harves McSween (11) and Mittlefehldt (4) as an on the basis of microprobe chemister equilibrium phase relationships, the globules were formed by high-tempermetamorphic or hydrothermal rea-Alternatively, Romanek et al. (12) on the basis of isotopic relationship the carbonates were formed under loss porature hydrothermal conditions (5) panophase magnetite and, iron me ficies present in these globules would net be detected in microprobe at which normally have a spatial resolution mout a micrometer. Our TEM obse und our S maps suggest that na proprietite and iron sulfides, while tranci in some zones, are present (can the globules. The effect of the to that exide and sulfide minerals serverse microprobe analyses m the interpretation of the micropr

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(4, 11) uncertain. Alternatively, if the globules are products of biologic activity, a low-temperature formation would be indicated. The textures of the carbonate globules are similar to bacterially induced carbonate crystal bundle precipitates produced in the laboratory and in a fresh water pond (57). Moreover, the observed sequence in the martian carbonate globules-Mn-containing carbonate production early (in the core) followed by Fe carbonate and finishing with the abundant production of re-Juced Fe-sulfides is a sequence that is common in terrestrial settings, as Mn is first reduced by biogenic action, followed by ferric iron and sulfate (57). Pure Mg-carbonate (magnesite) can also be produced by ciomineralization under alkaline conditions (59). On the basis of these observations, we interpret that the carbonate globules have a biogenic origin and were likely formed at low temperatures.

It is possible that all of the described features in ALH84001 can be explained by inorganic processes, but these explanations appear to require restricted conditions, for example sulfate reducing conditions in Antarctic ice sheets, which are not known to occur. Formation of the described features by organic activity in Antarctica is also possible but such activity is only poorly understood at present. However, many of the described features are closely associated with the carbonate globules which, based on textural and isotopic evidence, were likely formed on Mars before the meteorite came to Antarctica. Consequently, the formation of possible organic products imagnetite and iron-monosulfides) within the globules is difficult to understand. if the carbonates formed on Mars and the magnetite and iron monosulfides formed in Antarctica. Additionally, these products might require anerobic bacteria, and the Antarctic ice sheet environment appears to be oxvgen-rich: ferric oxide formed from metallic Fe is a common weathering product in Antarctic meteorites.

In examining the martian meteorite ALH84001 we have found that the following evidence is compatible with the existence of past life on Mars: (i) an igneous Mars rock (of unknown geologic context) that was penetrated by a fluid along fractures and pore spaces, which then became the sites of secondary mineral formation and possible biogenic activity, (ii) a formation age for the carbonate globules vounger than the age of the igneous rock, (iii) SEM and TEM images of carbonate globules and features resembling terrestrial microorganisms, terrestrial biogenic carbonate structures, or microfossils, (iv) magnetite and iron sulfide particles that could have resulted from oxidation and reduction reactions

known to be important in terrestrial microbial systems, (v) the presence of PAHs associated with surfaces rich in carbonate globules. None of these observations is in itself conclusive for the existence of past life. Although there are alternative explanations for each of these phenomena taken individually, when considered collectively, particularly in view of their spatial association, we conclude that they are evidence for primitive life on early Mars.

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- 21. When referring to the Martian meteorites, we use microorganism in a general sense-to signify a morphology that can be interpreted as fossilized microorganisms or small groupings of such organisms on the basis of our knowledge of terrestnal features.
- 22. μL²MS was used to analyze fresh fractured samples of ALH84001 for the presence of PAHs. The μL²MS instrument is capable of the simultaneous measurement of all PAHs present on a sample surface to a spatial resolution of 40 μm, and detection limits are

in the sub-attomole (>10⁷ molecules) range (* amol = 10^{-16} mol).

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- 26. At a single laser ionization wavelength, µLTMS is unable to distinguish between structural maners however, because different isomers have afferent photoionization cross sections, mass assignments are based on the most probable isomer. In the case of masses 178 and 202, the possible isomerizamonations are phenanthrene or anthracene andavrene or fluoranthene. At the photoionization waveength used in this study, 266 nm, the µL2MS instrument is ~19 times as sensitive to phenanthrene asto anthracene and ~23 times as sensitive to pyreleas to fluoranthene (R. Zenobi and R. N. Zare, in Astences in Multiphoton Processes and Spectroscopy, S. H Lin, Ed. (World Scientific, Singapore, 1991).vol. 7 pp. 1-167]. Hence, masses 178 and 202 are assigned to phenanthrene and pyrene. In thesase c* higher masses, more structural isomers event, and assignments are based on those PAHs known to have high cross sections from companyees with standards.
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- Sources of laboratory contamination fall me three 31. categories: sample handling, laboratory artand wtual leaks (that is, sources of PAHs inside mail_2MS vacuum chamber). Possible contamination.dunng sample preparation was minimized by peterming nearly all sample preparation at the NASA-SC meteorite curation facility. In cases where sussicuent sample handling was required at Stanford, almaniculation was performed in less than 15 minutes using only stamless steel tools previously nosed and uttrasonicated in methanol and acetone. Dust-fitteploves were worn at all times and work was performed on a clean aluminum foil surface. To quantifyattoome contamination from exposure to laboratoniar, two clean quartz discs were exposed to amplementationatory environments both at NASA-JSC and Bantorc Each disc received an exposure typical of the excerienced subsequently by samples of ALH84E1 outing sample preparation. No PAHs were observed cr either quartz disk at or above detection was. Because contamination can depend on thesenvsica. characteristics of the individual sample (for.eample) a porous material will likely give a larger collamination signal than a nonporous one) additionalization ination studies have been previously condicted at Stanford [see (28)]. Briefly, samples of the research acid residues of Banwell (L6) and Bishunder (L2." were exposed to laboratory air for 1 and# davs

respectively. Barweil (L6) is known to contain no incigenous PAHs and none were observed on the exposed sample. Bishunpur (L3.1), in contrast, contains a nch suite of PAHs, but no discernible differences in signal intensities were observed between exposed and unexposed samples. To test for contamination from virtual leaks, the µL2MS instrument was penodically checked using samples of Murchison (CM2) meteorite matrix whose PAH distribution has been previously well characterized. No variations n either signal intensity or distribution of PAHs were coserved for µL2MS instrument exposure times in excess of 3 days. No sample of ALH84001 was in the instrument for longer than 6 hours. The μL^2MS vacuum chamber is pumped by an oil-bree system: two turbomolecular pumps, and a liquid-hitrogencooled cyropump

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Distance from Fusion Crust Exterior (Micrometers)