FOCUSED ION BEAM MICROSCOPY OF ALH84001 CARBONATE DISKS

Kathee L. Thomas-Kepra1,2, Simon J. Clemetti1, Dennis A. Bazykin1, Joseph L. Kirschvink1, David S. McKay2, Hojatollah Vali2, Everett K. Gibson, Jr.3, and Christopher S. Romanek1; 1Lockheed Martin Space Operations, 2400 NASA Road 1, Mail Code C23, Houston, TX 77058 (kthomas@ems.jsc.nasa.gov), 2Texas Southern University, 3100 Cleburne Ave., Houston, TX 77004, 3Iowa State University, Department of Microbiology, 207 Science I, Ames, IA 50011, California Institute of Technology, Division of Geological and Planetary Sciences, 1200 East California Boulevard, Pasadena, CA 91125, NASA Johnson Space Center, Mail Code SN, Houston, TX 77058, 4McGill University, Department of Earth and Planetary Sciences, 3450 University Street, Montreal, PQ H3A 2A7, Canada, and 5Savannah River Ecology Laboratory, Drawer E, University of Georgia, Aiken, SC 29802.

Introduction: Our aim is to understand the mechanism(s) of formation of carbonate assemblages in ALH84001. A prerequisite is that a detailed characterization of the chemical and physical properties of the carbonate be established. We present here analyses by transmission electron microscopy (TEM) of carbonate thin sections produced by both focused ion beam (FIB) sectioning and ultramicrotomy. Our results suggest that the formation of ALH84001 carbonate assemblages were produced by considerably more complex process(es) than simple aqueous precipitation followed by partial thermal decomposition as proposed by other investigators [e.g., 1-3].

Method: All the carbonate assemblages studied appeared as pseudo-spherical disks embedded within the orthopyroxene (Opx) matrix and were flush with the fracture surface (Fig 1). These were removed intact by careful application of pressure to the carbonate/Opx interface (and a certain amount of luck). A typical extracted assemblage was ~200 µm in diameter and ~20 µm thick. Electron transparent sections necessary for TEM analysis were prepared by two methods: (1) ultramicrotomy of epoxy embedded samples; and, (2) FIB sectioning of Pt-coated samples. Ultramicrotomy: 30-40 µm size fragments from the intact carbonate disk were embedded in epoxy and sectioned with a 45° diamond knife onto distilled water from which they were transferred to continuous C film TEM grid. Typical section thickness was 90-100 nm. FIB Sectioning: A complete carbonate disk was attached to a scanning electron microscope (SEM) mount by carbon adhesive tape and coated with ~40 nm of Pt. Regions of interest identified by SEM were subsequently extracted using a FEI dual beam Strata 237 Ga FIB1 (for a more complete discussion of this technique see [4]); a typical FIB slab was ~ 20 x 7 x 0.1 µm. Ultramicrotome and FIB sections were analyzed with a JEOL 2000 FX TEM equipped with a Thermo Noran System Six energy dispersive x-ray spectrometer (EDS).

Results and Discussion: In ultramicrotomy the ‘knife’ that sectioned the sample is (in our case) actually a 45° cleaved diamond wedge. This requires that as a section is cut it must be flexed to an equivalent angle at the point the knife intersects the sample. To obtain a contiguous, uniform thin section this requires the sample has a relatively low Young’s modulus, be physically homogeneous, and structurally free of pre-existing fractures; unfortunately ALH84001 carbonate assemblages meet none of these criteria. Consequently carbonate microtome sections are typically fragmented and fail to preserve µm-scale structural and textural relationships (Fig. 2). Furthermore, once a section has been cut by the diamond wedge, it is pushed into a distilled water bath and suspended by surface tension. Capillary action likely penetrates the sample resulting in possible contamination and/or relocalization of indigenous water soluble salts.

FIB sectioning uses a focused Ga+ beam to sputter away selected regions of the sample to render an isolated slab that can subsequently be picked out of the sample using a micromanipulator and transferred directly to a continuous C film TEM mount. Since there is little or no structural flexing, the physical size of the section is not constrained by the same factors as microtomy (Fig. 2). Consequently, large slabs of electron transparent carbonate with a length-to-thickness ratio > 300 can be removed intact from the sample (Fig. 3). This allows the spatial location of the sample to both the carbonate and the Opx host matrix to be fully preserved (Fig. 3). One potential disadvantage of the FIB technique is perturbation of the carbonate matrix surface by embedded Ga+ ions.

Sectioning by FIB reveals the carbonate to have a low porosity (~20%), with parallel strain fractures and cross-cut with S-rich, grain filled veins. The chemical zonation of the carbonates occurs only along two axes and is invariant on the third, that is zoning propagates radially outward from the carbonate center but is constant through the thickness of the carbonate at a given radius (Fig. 4).

Void space appears throughout the carbonate, sometimes the presence of a void is accompanied by magnetite crystal(s), however we also observed fully encapsulated voids within the section containing no magnetite. Other investigators have suggested that void spaces with associated magnetite(s) were formed by thermal decomposition of Fe-carbonate to magnetite. We note that in many cases the void size associated with magnetite grains is significantly larger than could be explained solely by the density difference between magnetite and carbonate.

A, as yet undetermined, Si-rich phase(s) is present homogeneously throughout the carbonate at ~ 0.1-1.0 wt.% level. We cannot discount the presence of phyllosilicates and/or a Si-gel/glass.

Single crystal magnetite exist throughout the carbonate assemblages from core to rim but have the highest concentration in the optically dark outer rims. All magnetites characterized in the FIB sections were chemically pure, even when embedded within a carbonate matrix both Mg- and Mn-rich (Fig. 5). Both of these elements are capable of forming solid solutions of Mg- and Mn-ferrite, in particular incorporation of small amounts of Mn into magnetite may be thermodynamically favored due to the resulting lowering of the Curie temperature [5].

Implications: The use of FIB sectioning provides a hitherto unique view of the carbonate assemblages at the nm to µm scales in the absence of any embedding medium and under anhydrous conditions. The presence of carbonates inset into the Opx and flush with the fracture surface displaying cylindrically symmetrical zoning seems inconsistent with a simple aqueous precipitation process. Furthermore, the presence of the embedded magnetites within the sections studied here arguably requires one or more processes, other than thermal decomposition, to explain their origin.


We gratefully acknowledge and thank Giles Graham (Lawrence Livermore) and M. Bernas (FEI) for the milling and extraction of ALH84001 carbonate FIB sections.
Figure 1. ALH84001 carbonate disk embedded within a form-fitting pit and flush with the Opx matrix surface.

Figure 2. Images of a typical ALH84001 carbonate microtome section (top) and FIB section (bottom). Note the lack of sample continuity in the microtome section.

Figure 3. False color SEM image (left) of two FIB pits (orange) in ALH84001 carbonate (gray) and corresponding TEM views of extracted FIB sections (right).

Figure 4. Cartoon of ALH84001 carbonate disk depicting X, Y, and Z directions. Cylindrical symmetrical chemical zoning occurs in the X-Y plane. Carbonate composition is homogeneous through the thickness of the disk (Z value) at any specific X-Y point. ALH84001 carbonate disks appear to be composed of vertical, rather than horizontal, bands of zoned composition carbonate.

Figure 5. EDS spectra of two magnetites (bottom) embedded in carbonate matrix (top) in FIB sections. Carbonate contains detectable Mn while magnetite does not. Identical analysis time (1500 sec) and spot size were used.