Three-Dimensional Raman Tomographic Microspectroscopy: A Novel Imaging Technique

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Abstract Chemical constituents and their spatial distributions in terrestrial and extraterrestrial samples may provide clues to their formation histories and secondary processes they were subjected to. Most analytical tools are either limited in imaging the interior of samples or they are destructive such that the sample is chemically modified. In this work, we have conducted three-dimensional Raman tomographic imaging experiments on both terrestrial and extraterrestrial samples. The in situ nondestructive analytical technique presented here provides spatial distribution of various chemical constituents from the interior of the samples with high spatial and depth resolution. Raman data were first collected from two-dimensional layers at the surface and at various depths inside the samples by means of confocal Raman imaging. Individual chemical distribution maps were generated for each layer, which were then stacked to form and visualize the three-dimensional distributions. In addition to three-dimensional distributions, Raman spectral information was also collected from each pixel within the measured areas, thus allowing us to also perform detailed spectral investigations of the samples.

1. Introduction

Raman spectroscopy is an inelastic scattering analytical technique for molecular identification and characterization, and it has been used for the investigation of variety of samples in different areas of research. In meteoritics and planetary science, Raman spectroscopy is a widely used analytical technique. For instance, Ling et al. (2011) used Raman spectroscopy to investigate lunar soils in the laboratory. Interfaces between chondrules and the matrix of a carbonaceous chondrite were studied by Raman spectral mapping in order to understand the physical processes during the planetesimal formation (Ghimire et al., 2012). Insoluble organic matter extracted from various meteorites has been investigated using Raman spectroscopy in order to assess the degree of primitiveness of organics in meteorites (Busemann et al., 2007). Jaret et al. (2017) used Raman spectroscopy and imaging to investigate the occurrence and distribution of coesite in a shocked silica amygdule in Deccan basalts from Lonar Crater, India. Similarly, Raman spectroscopy has been widely used to identify high-pressure, high-temperature polymorphs in meteorites (Ohtani, 2007).

Although less common in terrestrial geological sciences, Raman spectroscopy is growing in use in these fields as well, particularly for the identification of polymorphs and fluid inclusions (Dubessy et al., 2012, and references therein). Additionally, Raman spectroscopy plays a critical role in diamond anvil cell studies of minerals at high pressures as it is used for a well-established pressure calibration of experiments (Bassett, 2009; Li et al., 2013; Schmidt et al., 2013).

Until recently, Raman spectroscopy has mostly been used in one- or two-dimensional space such that either a Raman spectrum of a point on a sample is collected or intensity distribution maps of an area on the surface of a sample are acquired. By collecting a single Raman spectrum from powdered grains of the Sutter’s Mill meteorite, Yesiltas et al. (2014) presented the unusual presence of two distinct carbonaceous phases in this meteorite with different thermal histories. Wang et al. (2015) utilized Raman imaging spectroscopy and reported Raman spectral features of major, minor, and trace species in extraterrestrial as well as terrestrial materials and also presented their spatial correlations to reveal genetic relationships and the geological processes that these materials have experienced. Whereas these studies provide important information and clues, they are limited to the surface of the samples that are under investigation, because both Raman spectra and Raman images are collected only from the surface within the focal plane of the microscope. In these cases, therefore, organic and inorganic compounds below the surface are completely missed and not
observed. However, it is highly important in some fields, especially in meteoritics and planetary science, to extract as much information from a sample as possible due to the limited amount and precious state of the samples. Additionally, context of small phases can often provide insight into formation mechanism or process that would otherwise be lost during the investigation of powders or through the analyses of point spectra alone (e.g., Jaret et al., 2017). This goal can be achieved via three-dimensional tomographic investigation. Using three-dimensional Raman tomographic microspectroscopy, the sample of interest can be investigated in situ in three-dimensional space both spatially and spectrally with high resolution at the same time without minimal to no sample preparation.

In this paper, we report the application of three-dimensional Raman tomographic microspectroscopy to study various extraterrestrial and terrestrial samples. Namely, we report Raman tomographic imaging microspectroscopy of an unequilibrated ordinary chondrite Meteorite Hills 00526 (MET 00526), coesite occurrence inside SiO2 glass from the Lonar Impact Crater in India, pyrite within calcite carbonate matrix from the Shullsburg formation in Wisconsin, rutile occurrence inside quartz from the Pikwitonei granulite of the Superior Province, and a banded iron formation (BIF) that is composed of alternating layers of iron-rich minerals, to cover a range of materials that are of importance in the field of meteoritics and geology. We remark that the extraterrestrial samples that will be returned to Earth in the near future from the primitive asteroids Ryugu and Bennu through Hayabusa2 (Tsuda et al., 2013) and OSIRIS-REx missions (Lauretta et al., 2015), respectively, can also benefit from this technique for the investigation of their chemical compositions.

### Table 1

<table>
<thead>
<tr>
<th>Experimental parameters</th>
<th>MET 00526</th>
<th>Coesite/glass</th>
<th>Pyrite/calcite</th>
<th>Rutile/quartz</th>
<th>BIF</th>
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<tr>
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<td>80 × 80</td>
<td>80 × 80</td>
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<td>1.92 × 10⁵</td>
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Figure 1. (left) Distribution maps of a chemical component (olivine in MET 00526 meteorite) collected at different depths (Z) layer by layer starting from the surface. (right) TIFF image of olivine in the MET 00526 meteorite stacked together using ImageJ software package (Movie S1).

### 2. Materials and Descriptions

#### 2.1. MET 00526

Meteorites provide invaluable information about the formation and evolution of planetary bodies (Day, 2015; Veronique et al., 2012). We gain significant insights from the organic and inorganic content of meteorites by studying their chemistry in the laboratory. Meteorites are divided into six groups based on their petrology (petrologic types). Meteorites of petrologic types from 1 to 2 are from aqueously altered parent bodies and include substantial secondary mineral components. On the other hand, petrologic types 4 to 6 indicate thermal metamorphism in the parent bodies. Most of the type 3 meteorites were subjected to some thermal metamorphism (much less than types 4 to 6), except type 3.0, which experienced minimum thermal metamorphism. Type 3 meteorites are perhaps the most primitive samples (or least altered) considering that aqueous alteration and thermal metamorphism modify the chemistry of parent bodies. The MET 00526 meteorite is an unequilibrated ordinary chondrite with petrologic type of 3.0 (Grossman & Brearley, 2005). Ordinary chondrites normally lack carbonaceous matter due to parent body processes they underwent.
Figure 2. (a) Polarized image of the MET 00526 thin section, roughly ~0.5 cm. Large chondrules of different mineralogy are shown with a dark matrix in the meteorite. (b) Expanded version of the measured region shown with a red square. (c) visible micrograph of the region obtained with a 20X objective. (d) Color-coded distributions of observed components overlayed on the visible image. The colors correspond to the individual components (yellow: pyroxene; green: olivine; blue: carbon). (e–h) Two-dimensional intensity distribution maps of pyroxene, carbonaceous phase, olivine, and high fluorescence region as obtained from the first layer (surface) of the meteorite sample. Among the possibilities for such fluorescence are Fe associated with the olivine or rare-earth element that may be present on the sample. Raman spectra of the components observed in the first layer (surface) of MET 00526 are given in the bottom. See Table 2 for peak positions.
and complex formation mechanisms for carbonaceous matter. However, unequilibrated chondrites do contain carbon and other primary chemical components.

We received the MET 00526 meteorite sample from the Antarctic meteorite collection of NASA Johnson Space Center in the form of a thin section. MET 00526 is an unequilibrated ordinary chondrite with carbon-rich matrix and silicate-rich mineralogy. The thin section cut from the meteorite (roughly ~30 μm thick) was glued on a glass slide using epoxy at the NASA’s Johnson Space Center. We studied this meteorite for the visualization of various chemical components present in the meteorite sample.

2.2. Coesite Occurrence inside SiO₂ Glass

Coesite is a high-pressure, high-temperature polymorph of SiO₂ that is found in ultrahigh pressure metamorphic rocks (e.g., eclogites, Chopin, 1984) and in impact metamorphosed samples (Chao et al., 1960; Jackson et al., 2016; Jaret et al., 2017; Kieffer, 1971). Coesite can form as a direct solid-state product from quartz (Dachille et al., 1963) or in rare cases can form as the crystallization product from a SiO₂ melt (Chen et al., 2010; Stahle et al., 2008).

The coesite sample is within shocked basalts from the Lonar Crater, India (Jaret et al., 2017; Newsom et al., 2013; Wright et al., 2011). The coesite is located within SiO₂ glass that is itself within basalts that were shocked to Class 2 (Kieffer et al., 1976) or ~25–40 GPa. Coesite and glass are confined only to small (1–3 mm) vesicle fillings, which are interpreted to represent secondary silica precipitates formed prior to the shock event (Jaret et al., 2017). Measurements were acquired on standard polished petrographic thin sections (approximately 30 μm thick).

2.3. Pyrite Within Calcite Matrix

The pyrite sample is from the Blackstone Mine in the Upper Mississippi Valley District (Shullsburg quadrangle) of Wisconsin, composed of Ordovician to Silurian sandstones and carbonates. These units have been heavily mined for Pb and Zn during the nineteenth century and are well-established regions of mineralization and fluidization (Mullens, 1964). This mine primarily produced sphalerite and galena, with lesser amounts of marcasite, calcite, barite, and pyrite. Rare pyrite occurs as thin layers of euhedral cubes within wall rocks.

2.4. Rutile Occurrence inside Glass

This sample is a large quartz vein in the Pikwitonei granulite domain of the Superior province in Manitoba (Hemming et al., 1994). The quartz contains numerous inclusions of rutile and zircon, with rutile occurring in two size populations: ~500 μm and thinner needles that appear to follow crystallographic orientations.

2.5. BIF

BIF is a term that refers to the sedimentary iron-rich rocks that include layers of iron oxides, silica, and more rarely, other phases. These sedimentary iron rocks are believed to have deposited during a period of

<table>
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<th>Table 2</th>
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<tr>
<td>Positions of the Observed Raman Peaks and Their Corresponding Assignments</td>
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<tr>
<td>Sample</td>
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<td>MET 00529</td>
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<tr>
<td>Coesite in silica glass</td>
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<td>Pyrite with calcite</td>
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<td></td>
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<tr>
<td>Rutile inside quartz</td>
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<tr>
<td>BIF</td>
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800 Ma from late Archean to mid-Proterozoic (Klein, 2005). It was reported by Simonson and Schubel (1993) that carbonates were related to the microbial activity in the early history of Earth, and they were associated also with BIFs, and therefore, the BIFs can provide insights for astrobiologically relevant sites and locations on Mars (Crowley et al., 2008). The BIF sample studied here is from the Sherman mine in the Tegami greenstone belt in the Superior Province of Ontario Canada. These Algoma-Type BIFs have an age of ~2700 Ma, adjacent to the slightly younger and more heavily studied Abitibi Greenstone Belt (Ginley, 2016).

3. Experimental Details and Data Analyses

3.1. Technical

We conducted our Raman tomographic imaging experiments at the Center for Planetary Exploration at Stony Brook University, New York, and at the Science and Technology Application and Research Center of Canakkale Onsekiz Mart University, Turkey. Both experimental setups consist of the same instruments, a WiTec alpha300 R (WiTec GmbH) confocal Raman imaging system equipped with a 532-nm Nd:YAG laser, spectrometer with a CCD camera (peltier cooled to –60 °C), and 50X objective (NA = 0.8). The laser power on the sample surface was kept between 0.2 and 2.4 mW measured at the objective to avoid thermal alteration of the samples. After locating a region of interest in the field of view of the microscope, two-dimensional Raman intensity maps of the surface were collected from predefined areas with either 0.5- or 1-μm spatial resolution, and a full Raman spectrum was collected at every pixel within the measured area with ~0.05- to 0.08-s integration time. Experimental parameters set before each data collection session included area to be measured, spatial pixel size desired, integration time, depth to be scanned, and the number of layers desired within the defined depth, which determines the depth resolution. During the data collection, the predetermined area of each layer was measured starting from the surface of the samples down to the deepest layer with either 0.5- or 1-μm depth resolution. Low laser power and very short exposure time resulted no laser-induced damage or heating on the samples. Experimental parameters specific to each studied sample are given in Table 1.

3.2. Data Analyses

Data processing included several steps and software packages. After the data collection, we manually removed the artificial cosmic rays as well as the background. Using WiTec’s Project software, chemical distribution maps of individual chemical components were generated by integrating the signal between the endpoints of observed Raman peaks. This procedure was repeated for each layer collected at different depths (Figure 1, left). The distribution maps of all layers were then imported to Fiji-ImageJ software package. Using ImageJ, distribution maps extracted from all layers (40 layers for MET 00526, 30 layers for the pyrite, and 20 layers for the coesite, rutile, and BIF samples) were stacked individually and saved as stacked TIFF images (Figure 1, right), which were then imported to a commercial software package (Avizo) for the visualization of chemical distributions in three dimensions. In Avizo, all components in each sample were color coded and visualized together, and movies of various orientations were generated.

4. Results and Discussions

4.1. MET 00526

A 100-μm × 100-μm area and 20-μm depth from the MET 00526 was measured with 0.5-μm depth resolution and 1-μm spatial resolution, which resulted in 40 layers of Raman data cubes. Using the first layer out of 40 (the top surface layer), we determined the chemical components that are present in the meteorite. The matrix of the meteorite appears to be rich in carbon. While most ordinary chondrites lack carbon, the presence of carbon in the MET 00526 is not surprising considering it is a type 3.05
Figure 5. (a) Visible micrograph of the coesite mineral within glass. (b, c) Two-dimensional intensity distribution maps of coesite and glass. (d) Combined distribution map of the two components (yellow/orange is coesite and blue is SiO₂ glass). (e) Combined distribution map overlaid on the visible micrograph. The middle panel presents the Raman spectra of coesite and glass. See Table 2 for peak positions. The bottom panel presents reconstruction of coesite within the glass matrix (Movie S5) on the left, and clipping-plane cross-section view of this reconstruction (Movie S6) in the right. See supporting information files for movies of the rotations.
meteorite, and unequilibrated ordinary chondrites normally present relatively higher amounts of carbon.

Figure 2 presents reflected plane-polarized light microscope images of MET 00526. We measured a region that contains silicates and carbon-rich matrix, indicated in Figure 2b with a red rectangle. A visible reflected light micrograph of the measured region was collected with our Raman microscope and is shown in Figure 2c. We collected two-dimensional Raman data cubes from an area of $100 \mu m \times 100 \mu m$ with 1-\mu m pixel size and 0.05-s integration time between 100 and 3,700 cm$^{-1}$. We scanned a depth of 20 \mu m with 0.5-\mu m depth resolution, which resulted in 40 layers. Each layer consists of 10,000 pixels, therefore a total of 400,000 Raman spectra were collected for this sample. We generated the two-dimensional chemical distribution maps for these components, as shown in Figures 2e–2h. Figure 2d shows the color-coded distribution of these components overlaid on the visible micrograph to present how and where they are located in the measured area.

Figure 2 also presents Raman spectra of the observed chemical components (see Table 2 for peak positions and their assignments). Raman peaks at 1,341 and 1,601 cm$^{-1}$ are due to disordered (D band) and graphitic (G band) carbon bands, respectively. We also observed the presence of anhydrous silicates such as olivine and pyroxene. A doublet near 845 cm$^{-1}$ is characteristic of olivine. Pyroxene presents Raman peaks near 352, 680, and 1,020 cm$^{-1}$. Impurities in the sample (Kagan & McCreery, 1994) can intrinsically generate fluorescence due to absorption by the molecule at the wavelength of the laser probe. In the case of MET 00526, the fluorescence appears in the region of 2,500–3,500 cm$^{-1}$ and its Raman spectrum is also shown in Figure 2.
Using the Raman spectra and distribution images obtained from the first layer, we investigated the succeeding layers and generated the distribution maps of the components for all layers. During the data analyses of the MET 00526 meteorite, another component was observed to appear around 10-μm depth and continue until the depth of 20 μm in our measurement. The Raman spectrum of this component (Figure 3) presents various C-H bending and stretching modes both in the fingerprint region as well as in the higher-frequency region. This component is in fact the epoxy resin used during the sample preparation. The meteorite thin section was prepared by gluing it on a glass slide using epoxy, and it was vacuum impregnated when it was first placed in epoxy. Therefore, it is possible for the epoxy to penetrate into the meteorite sample to this extent (Roger S. Harrington, private communication, May 2016). The analyses of MET 00526 shows that one has to be extra cautious when studying organic compounds in meteorites that are prepared in any way in the laboratory.

We combined distribution maps from all layers using Avizo and generated the three-dimensional tomographic reconstructions of the distributions of all observed components in MET 00526. Figure 4 presents selected 2-D projections of these reconstructions. See supporting information files for movies of the rotations.

### 4.2. Coesite in Silica Glass

We measured a 80-μm × 80-μm area of the coesite sample with 1-μm depth resolution. The coesite exhibits a strong peak at 521 cm⁻¹, which reflects the Si-O-Si stretching mode. Coesite also exhibits weaker peaks at 113, 173, 267, and 429 cm⁻¹. The SiO₂ primarily exhibits a broad peak near 449 cm⁻¹, with a substantial drop-off in intensity at 494 cm⁻¹. The 2-D and 3-D mapping of this sample shows that the coesite occurs as ~30-micron spherical aggregates that are actually composed of smaller, micron-sized, individual microcrystallites and nanocrystallites. This is particularly apparent in the 3-D maps that show SiO₂ glass underneath many of the coesite grains (Figure 5).

### 4.3. Rutile Inside Quartz

We measured a 80-μm × 80-μm area on the rutile sample, where the rutile appears as a cylindrical bar that extends into the quartz matrix at an angle (Figure 6). The quartz is identified by the strong peak at
464 cm$^{-1}$ and weaker peaks at 205 and 125 cm$^{-1}$. Rutile exhibits a strong peak at 612 cm$^{-1}$ and a weaker peak at 447 cm$^{-1}$. Smaller grains of rutile are identified by the peak at 612 cm$^{-1}$, but because the rutile needles are smaller than the sampling volume (single pixel) of the Raman, these spectra are not pure. In particular, the mixtures show both the 464-cm$^{-1}$ peak of quartz plus the additional 612-cm$^{-1}$ peak. The 447-cm$^{-1}$ peak in rutile is masked by the 464-cm$^{-1}$ peak from quartz. Although small individual nanocrystals of rutile can be identified by mapping of the 612-cm$^{-1}$ peak, quantification of the amounts and size of the rutile cannot be determined. Mapping in 3-D is useful in this case because it illustrates the 3-D sampling volume of the Raman imaging instrument.

This particular rutile sample also contains many thin needle-like rutile occurrences inside the quartz matrix; they are randomly oriented inside the quartz and are ~0.5 to 1.0 μm in diameter, and 5 to 20 μm in length (Figure 7a). These rutile needles are spatially resolved in high-resolution three-dimensional Raman tomographic imaging experiments. Figures 7c-7f show the 3-D distribution of the needles at different orientations.

4.4. Pyrite Within Calcite

An area of 80 μm × 80 μm was measured to resolve the cubic pyrite crystals in three dimensions. The 3-D mapping clearly shows the cubic shape of the pyrite grain, which is composed of two cubes adjacent to one
Figure 9. (a) Visible micrograph of the measured BIF sample with the false color combined distribution maps overlaid. Colors correspond to the individual components (green: dolomite; blue: quartz; and yellow: hematite). (b) False color combined distribution map of the components. (c–e) Distribution maps of dolomite, hematite, and quartz, respectively. The middle panel presents Raman spectra of the components observed in the BIF sample. See Table 2 for peak positions. The bottom panel presents reconstruction of the BIF sample shown at different orientations (Movie S15), and the clipping-plane cross-section view of the reconstruction (Movie S16). See supporting information files for movies of the rotations. BIF = banded iron formation.
another inside a calcite matrix. The calcite exhibits a strong peak at 1,088 cm$^{-1}$ and weaker peaks at 713, 283, and 155 cm$^{-1}$ (Figure 8), while pyrite is represented by the two strong peaks at 337 and 370 cm$^{-1}$. Our 3-D Raman imaging data reveal that the pyrite is pure and did not cocrytallize with other sulfides, which is a common trait of hydrothermal pyrite and distinct from metamorphic pyrite (Craig et al., 1998).

4.5. BIF

We have measured a 150-μm × 60-μm region on the BIF sample and observed the presence of quartz, dolomite, and hematite, whose Raman spectra are shown in Figure 9. Quartz presents a prominent peak at 472 cm$^{-1}$ as well as smaller features near 200 cm$^{-1}$. The dolomite presents well-known carbonate peaks near 186, 305, and 1,100 cm$^{-1}$. Hematite is also present in this sample. It has a noisier spectrum, which presents a peak at 1,325 cm$^{-1}$. The reason for the lower signal-to-noise ratio in the case of hematite is that relatively few pixels were averaged to compose the spectrum.

Figures 9a–9e presents the two-dimensional intensity distribution maps for the observed components. The brighter material near the middle of the visible image (Figure 9a) is dolomite within the silicate (quartz) and iron oxide (hematite) matrix, which are distributed throughout the sample Figure 9 (bottom) presents the images of the three-dimensional distributions at different orientations for a depth of 20 μm.

5. Conclusions

Three-dimensional Raman mapping is a powerful tool for the identification of various organic and inorganic phases in a wide range of terrestrial and extraterrestrial geologic materials. This technique especially has promise in the fields of meteoritics, high-pressure/temperature mineralogy, and for the identification of mineral inclusions because of its ability to spatially map various molecular functional groups and to investigate statistical correlations that can potentially reveal past chemical processes. Three-dimensional textural and chemical observations through this Raman tomographic mapping are particularly useful for our understanding of formation pathways and scenarios as well as parent body alteration processes in the extraterrestrial samples. For instance, spectral parameters of D and G carbon bands can be used to infer peak metamorphic temperatures that the parent asteroid of meteorites experienced; however, since 2-D Raman imaging collects the signal from the surface, such spectral parameters may yield artificially modified parameters because of the modified carbon structures on the surface of samples during the sample preparation. The 3-D tomographic imaging allows the collection of carbon D and G band spectral parameters from the interior of samples where carbon phases have not been altered in any way in the laboratory, thus leading to more correct understanding of the thermal histories of parent asteroids. Organic and inorganic compounds in extraterrestrial materials are not always seen in the surface of thin sections, which is the most common form of meteorite samples; however, deeper regions of the sample may contain valuable compounds that can be easily observed and identified via 3-D Raman imaging. Oxidation processes on newly exposed surfaces of samples can be studied to understand reaction products as well as reaction kinetics and mechanisms. These reactions take place in very short timescales, and with 2-D Raman imaging the data collection is possible after the sample is cut; however, the oxidation reactions starts as soon as the surface is exposed. The 3-D Raman imaging allows the data collection even before the sample is cut to expose the fresh surface such that the oxidation processes and prereactions and conditions can be observed prior to the exposure of the interior of the sample to the oxidizing environment.

Similarly, as shown for the rutile sample, the context and orientations of small inclusions can be readily resolved and identified with 3-D imaging. This is important for precious samples where as little cutting of samples is necessary. Three-dimensional imaging can serve as a prescreening technique to better inform where or at what orientation to cut samples if that becomes necessary.

We have demonstrated the application of this novel nondestructive Raman imaging technique to the Antarctic carbonaceous chondrite MET 00526, which revealed the presence of nonindigenous organic material hidden underneath olivine. The carbonaceous phase appears to be distributed within the top 3–5 μm of the sample; however, in some locations it extends to deeper into the sample. The 3-D distribution, morphology, and shape of coesite have been revealed with high resolution in shocked basalts from the Lonar crater in India. The cylindrical rutile bar inside quartz as well as the thin randomly oriented 0.5–1 μm diameter needle-like rutile occurrences inside the quartz matrix have been observed. The 3-D tomography of pyrite in
carbonates show that the pyrite is composed of two adjacent cubes. The pyrite also appears to be pure and did not cosinter with other sulfides, which is a common trait of hydrothermal pyrite and distinct from metamorphic pyrite. Lastly, 3-D Raman mapping reveals the alternating layers of iron in the BIF, which may provide insights for determination of astrobiologically relevant locations due to its association with carbonates that may have been related to the microbial activity in the early history of Earth. The potential of 3-D tomographic imaging has also great potential for broad use such as solid-state devices, electronics as well as medical and biological investigations.

Acknowledgments
We are grateful to NASA’s Meteorite Working Group for the meteorite sample from the Antarctic meteorite collection, John Luczaj for the pyrite, Sidney Hemming and Troy Rasbury for the rutile, and Martin Schoonen and Hiroshi Ohmoto (Emeritus at Penn State) for the BIF sample. We thank reviewers and the Editor for their constructive comments. We also wish to thank Volkan Eskiýebek and Arife Aktas (Science and Technology Application and Research Center of Canakkale Onsekiz Mart University, Turkey) for their support during the Raman data collection. This work is funded by the RISÈ node of NASA’s Solar System Exploration Research Virtual Institute (SSERVI; PI: T. D. Glotch). The data presented here will be made publicly available via Turkish Spectral Database at http://tsd.klu.edu.tr/.

References