Keysight Technologies
Imaging Organic and Biological Materials with Low Voltage Scanning Electron Microscopy

Application Note
Introduction

Scanning electron microscopy has become a popular imaging tool in different areas of science and engineering\(^1\)\(^-\)\(^7\). Being able to elucidate the structure of a material at the micro- and/or the nano-scale level is indeed crucial to characterizing the material, understanding its mechanism and mode of formation, and explaining/predicting its properties and performance under a given set of environmental or load conditions\(^1\)\(^,\)\(^2\)\(^,\)\(^3\)\(^,\)\(^8\)\(^,\)\(^9\). Secondary electron imaging is commonly used to reveal surface topography, grain morphology and size, phase composition, and fracture profile. In such cases, highest resolution and contrast are needed to illustrate the finest structural features. However, imaging of organic and biological specimens represents a real challenge to investigators. We discussed briefly in a previous note\(^10\) the options available to overcome these difficulties. With the Keysight Technologies, Inc. 8500 low voltage scanning electron microscope (FE-SEM), imaging of organic and biological materials is strongly facilitated\(^11\)\(^,\)\(^12\). For this application note, we present here imaging data obtained with Keysight 8500 FE-SEM for different materials: (1) chemical and physical-chemical gelling of bioinjectable copolymers NC-NCA being explored as an alternative treatment for brain aneurism; (2) HeLa cervical cancer cells used to study the effect of irreversible electroporation on adherent cells; (3) biotically-precipitated calcium carbonate induced by CO\(_2\)-producing bacteria and enzyme for soil improvement.

Methodology

Data presented in this note concern samples from different areas of engineering. This includes: bioinjectable copolymers for medical application; culture cancer cells; biotic (both enzymatic- and bacterial-induced) precipitation of calcium carbonate for soil consolidation. Samples were processed and prepared in different ways, but preparation for FE-SEM imaging was minimal. Specimens are simply cut or fractured from original samples and adhered via carbon tape to aluminum stubs and loaded onto microscope stage. In the case of the HeLa cervical cancer cells, the entire electrodes plate (1 sq. in.) was adhered to the aluminum stub via C-tape and mounted onto the microscope stage. The samples examined here come from the following experiments:
Figure 1. SE Image sequence (a–f) obtained with Keysight 8500 for NC-NCA chemical gel. Images (a–c) clearly show the gel product is porous, but homogeneous, with large pores and channels sever microns across. Images (d–f) show the gel has plenty of the nano-pores (< 200nm).

(1) Copolymers: Chemical gel as well as physical-chemical gel systems were investigated. The experimental procedure is described in detail in reference [8]. The copolymers investigated here are poly(NIPAAm-co-cysteamine), abbreviated NC and poly(NIPAAm-co-HEMA-acrylate), abbreviated NCA. For SEM imaging, samples were cut with a razor blade and mounted directly onto carbon tape adhered to an aluminum stub. Physical-chemical gel material imaged here with FE-SEM is compared to material from a similar experiment observed with E-SEM after coating with gold. Chemical gel material is imaged first here. No previous images are available for comparison.

(2) Cervical cancer cells: These are the famous HeLa cells. The cells were grown on a microelectrode array and tested with variable voltage pulses. The aim of the experiment was to study the effect of irreversible electroporation on adherent cells. However, there might be a failure in the chemical fixing process of the cells at the end of experiment that caused damage to the membrane of most cells.

(3) Biotically-precipitated calcium carbonate: Samples studied here come from different experiments. In the first, quartz sand was consolidated by calcium carbonate (calcite) precipitated by the action of a CO$_2$ producing enzyme [9]. The sample was a fragment of a consolidated core sample that was adhered via carbon tape onto the aluminum stub. The second comes from an experiment done with CO$_2$-producing bacteria [12]. Peculiarly, calcite did not precipitate in the sand core, but as a pink crust that formed somewhere else in the system. The sample studied here was a fragment broken from this pink crust that was divided into two pieces of which face and back were imaged.
Results and Discussion

1. FE-SEM Imaging of copolymer chemical and physical-chemical gelling systems:

The gelling systems investigated here as possible treatment for cerebral aneurism are made of the copolymers poly(NIPAAm-co-cysteamine) and poly(NIPAAm-co-HEMA-acrylate). These hydrogels undergo gelation through dual mechanisms: temperature sensitivity (physical gelation) and chemical crosslinking (chemical gelation). Full description of the preparation and characterization of these gels are given in details[8]. The aim of imaging is to envision the evolution of micro- and nano-structure of chemical gel and compare with that of the physical-chemical gel. Figure 1 shows SE image sequence obtained with Keysight 8500 for NC-NCA chemical gel. Images clearly show the gel product is porous and homogeneous, with large pores and channels several microns across. At higher magnification, the gel is shown to have plenty of nano-pores (< 200nm). In contrast, the physical-chemical gel, while still having a spongy structure with micro pores and channels, has a heterogeneous composition. A minor grainy porous phase has formed in the cavities of the major (matrix) phase. The latter has only sporadic nano-pores. These observations suggest that structural (phase) changes during chemical gelation are different from those occurring during physical-chemical gelation. In the former, gelation occurs through chemical reaction (cross-linking) of the copolymers at constant temperature which is below their LCST (lower critical solution temperature). Whereas in the physical-chemical gelation, the temperature rises and allows physical and chemical gelation processes to occur simultaneously. This means some of physical gel gets trapped in the cavities of the chemical gel without having the chance for the copolymers to cross-link completely. This is in complete accord with X-ray diffraction and SAXS studies.
2. FE-SEM Imaging of HeLa cervical cancer cells:

These are cultured cells tested with variable electrical pulse. The whole electrode array with the tested and “fixed” cells was observed. The chemical fixing process might have been incomplete and caused destruction of the cells on most electrodes (control, 4V, 8V, and 12V). However, our main goal was to demonstrate that Keysight 8500 LV-SEM can be used efficiently to image biological materials, including cancer cells. Successful imaging, indeed, help bioengineers and microbiologists optimizing the experimental procedure. These both goals were served by the data acquired. Figure 3 shows SE images taken with Keysight 8500 FE-SEM of the aforementioned HeLa cervical cancer cells. The images show some of the cells were healthy at the end of the pulse experiments and survived the fixing process. These are indicated by the letter H on the images. Other cells, even sitting on the control electrode or on the substrate away from any electrode, have been killed. They are indicated by the letter K on the images. In the latter, the cell membrane has ruptured and the nucleus separated from the rest of the cell. The experiment shall be optimized and further imaging is needed to elucidate how the cell membrane interacts with an increasing electric pulse and whether nano-pores will open in it, thus allowing gene delivery into the cancer cells.

Figure 3. SE Images (a-f) obtained with Keysight 8500 for HeLa cervical cancer cells from an electroporation experiment. Cells indicated by H are considered healthy cells after the fixing process is finished. Whereas those indicated by K are cells killed by the fixing process or by the application of the electric pulse. In the latter case, cell membrane is being broken and nucleus separates from the rest of cell; some remnant connection may still be seen, as indicated by the double arrows.
3. FE-SEM Imaging of calcium carbonate precipitation for soil improvement:

Data presented here relate to two experiments[9]. In the first experiment, an enzyme extracted from plants is injected into the sandy soil. The enzyme convert urea into CO₂ and NH₃. Ammonia raises the pH and CO₂ combines with calcium ions to form calcium carbonate (mostly calcite), which acts as a cement to consolidate the soil. In the second experiment, denitrifying bacteria is injected with “its food” in the soil. CO₂ is produced as a result of bacterial metabolism and combines with calcium ions to form calcium carbonate, which serves as a cement in sandy soil improvement. The first experiment worked perfectly and quartz sand was cemented by the precipitated carbonate (calcite). The other experiment, however, gave some peculiar results. Calcite did precipitate in the sand core, but somewhere else in the system. Geotechnical engineers are investigating this issue. But, as far as the FE-SEM imaging is concerned, the sample examined met perfectly our goal of imaging bacterial cells without the need for sample coating. In both cases, high resolution images with excellent contrast were obtained for the biological and organic samples with minimal charging and no damage to them.

Figure 4 shows images of enzymatically-induced calcium carbonate precipitation. Full comment on these images is given in a previous application note[10]. FE-SEM imaging has permitted to better understand the mechanism of the reaction process and the different aspect of the role played by the enzyme[10].
In Figure 5 a set of SE images obtained with the Keysight 8500 FE-SEM is presented for the experiment with denitrifying bacteria. These images reveal many features of the biotic process. Bacterial cells are very abundant in the carbonate product. Some bacterial cells are spherical, some ellipsoidal, and some others rod-shaped. Some bacterial cells are actually in dividing stage or spore stage. It is worth noting the normal bimodal size of bacteria cells is 0.5–5 μm with most being 1μm and nanobacteria size being 100–300 nm. This is consistent with observation of fossilized bacteria in hydrothermal travertine (a rock composed of calcium carbonate, too)[2]. Note also the intergrowth of calcite crystals and bacterial cells. In many locations, bacterial cells or their remnant skeletons are buried inside the carbonate crystals. The latter have therefore a spongy porous internal structure. It was also observed that the outside surface of this carbonate product crust is well sealed, with carbonate crystals intermixed with bacterial bodies and crystal edges well rounded. Full interpretation of the entire body of data acquired for both carbonate precipitation experiments and explanation of the different implication to geotechnical engineering, biomineralization, and sedimentology shall be published soon in a peer-reviewed journal.
Conclusions

Using Keysight 8500 FE-SEM, it was quite possible to image, with high resolution and excellent contrast, insulating materials such as organic (copolymers and enzyme) and biological (cancer cells and bacteria), without the need for a metal coating or radiation damage to the samples. These analyses are otherwise not as feasible using other FE-SEMs: cold field emission SEM operating at low voltage and E-SEM, respectively. It is fair to say also that Keysight 8500 FE-SEM is more helpful in studying sensitive organic or biological samples such as enzymes, cell membranes, neurons, bacteria, biofilms, and biogels, when compared to other FE-SEMs currently available.

References


