Biological Sciences Tutorial: Freeze-fracture, Deep-etch and 3D Anaglyphs

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The advent of freeze-fracture electron microscopy in the late 1950’s provided a major breakthrough for cell biologists, enabling en face views of cell membranes and membrane leaflets1. Continued developments through the 1960’s and 70’s featured ultrafast freezing techniques2 that complemented parallel advances in metal coating units3. Rapidly frozen specimens, subjected to etching and rotary metal application gave birth to “deep-etching”, rendering detailed replicas of cell surfaces, organelles, cytoskeletons, and macromolecules, with ~1-2 nm resolution and striking three-dimensional perspective4,5.

Each wave of imaging advancement, whether in sample preparation, LM, TEM, SEM, CryoEM, ultramicrotomes, and molecular biology, bring with it an exciting new twist to the understanding of biological processes. Enjoying it’s 60th year of invention, freeze-fracture deep-etch EM holds up as a robust method of sample preparation, that is complementary to other contemporary approaches. The procedure is relatively quick, start to final imaging is typically a work day6. Recent use of Tokuyasu sectioning techniques, FRIL and STORM have given freeze-fracture a new life, providing additional cytochemical information to ultrastructure that was once elusive1,7-9. This tutorial will provide an overview of today’s freeze-fracture deep-etch EM technique.

References:

**Figure 1.** Purified molecules on mica. Freeze-fracture and deep-etch.

**Figure 2.** Freeze-fracture deep-etch of cell pellets and tissues.

**Figure 3.** Freeze-dried preparations of sonicated cell culture.