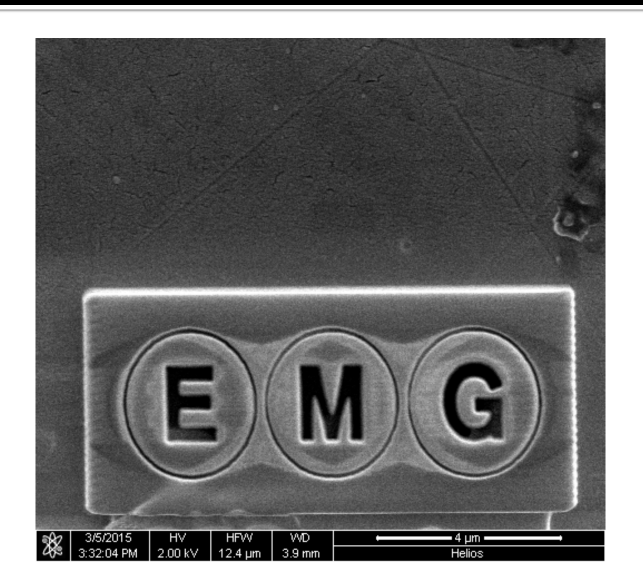
FIB-SEM Feb. 5, 2018

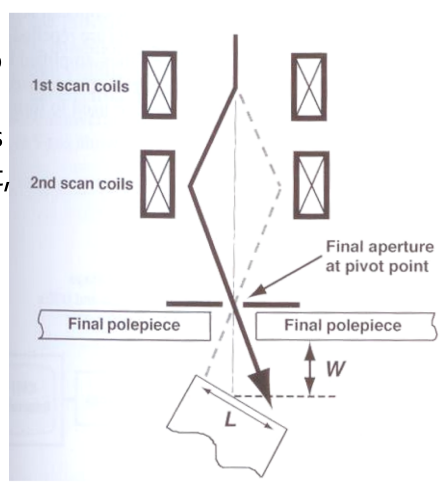


Outline

- SEM basics
- FIB Basics
- Application to room-temperature (conventional) specimens
- Use as a cryo-prep tool for cryo-TEM
- Use as a general prep tool (liftout) for TEM

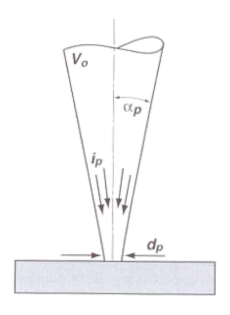
SEM Basics

- Electron probe is focused to a sharp point
- The probe is scanned across the specimen point by point, with each point producing signal
- Scan coils deflect beam to move across sample



SEM Beam: probe size

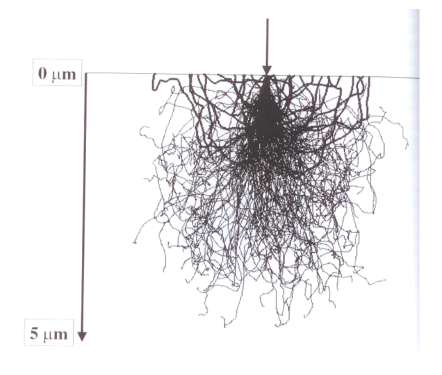
- Ideally want as small a probe as possible, relative to pixel size
- Probe size is determined by voltage, current, divergence angle
- Lens distortions
 - Spherical aberration (focus different at center and edge of lens) – instrument dependent
 - Aperture diffraction
 - Astigmatism (user correctable)
 - Chromatic aberration voltage dependent



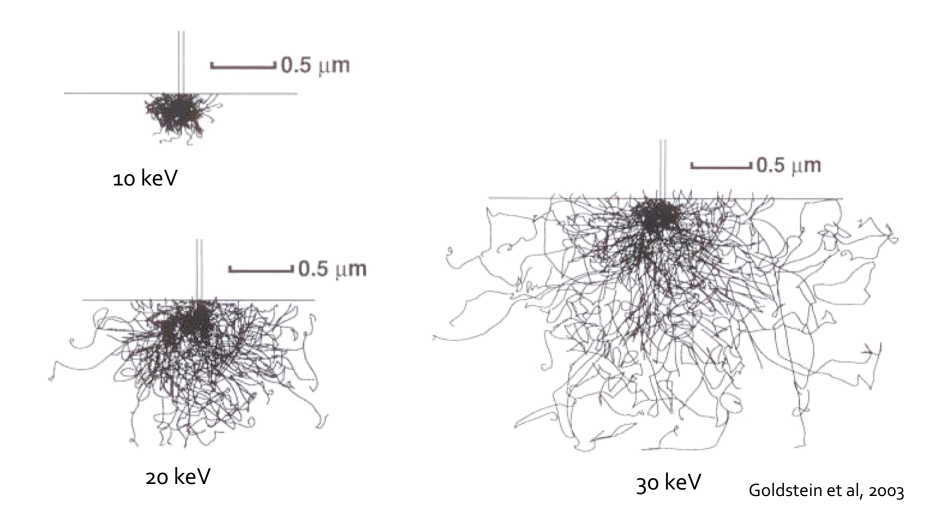
Goldstein et al, 2003

Beam-Specimen Interaction

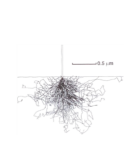
- Monte Carlo simulation of a 20 keV beam in Si
 - Dark traces: electrons which left the sample (BSE's)
- Electrons may be scattered elastically or inelastically
- Probability of elastic scattering ~ Z²
- Inelastic scattering:
 - Secondary electrons
 - X-rays

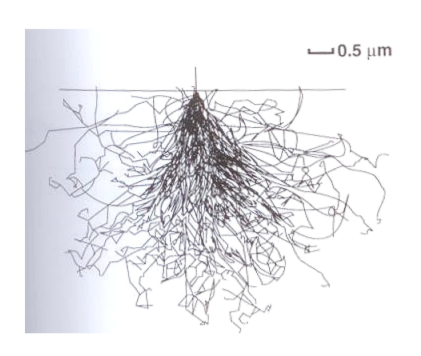


Simulations at different voltages



Material Dependence of Interaction Volume





Iron, 20 keV

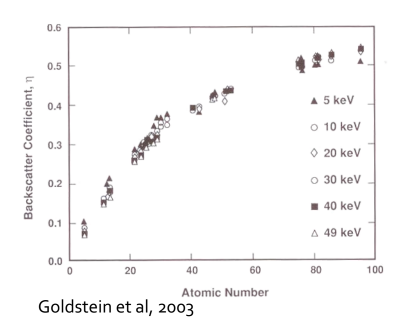
Carbon, 20 keV

Back Scattered Electrons and Secondary Electrons

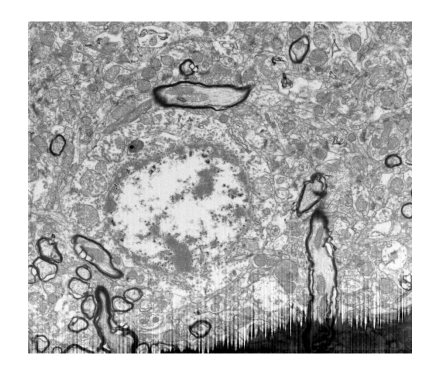
Detectors can be tuned for either one, or for both

BSE efficiency is material dependent, voltage independent

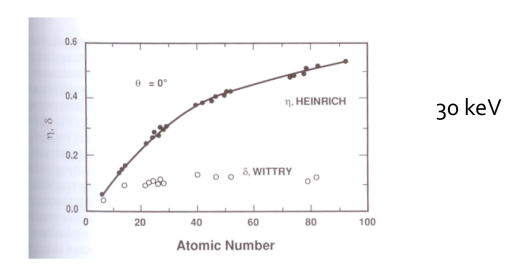
Fraction of e's that backscatter



BSE's give contrast between light and heavy elements



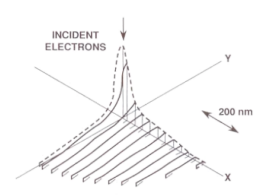
Specimen Dependence of BSE, SE



SE's are less sensitive to atomic number than BSE's (may be more sensitive at lower beam energies)

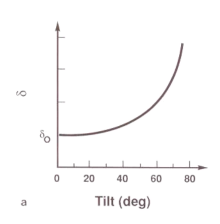
Goldstein et al, 2003

Emission shape of BSE's



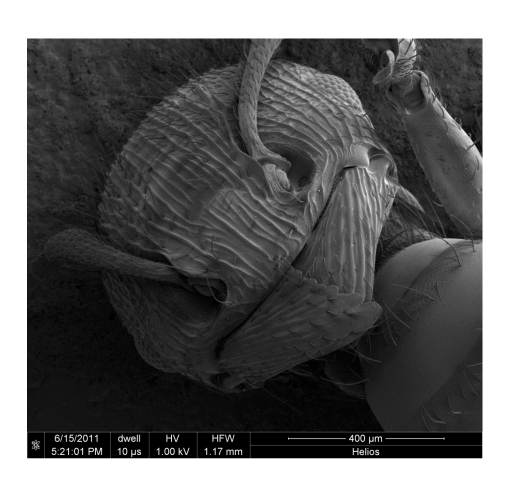
Most BSE's are released close to origin ("high quality" BSE's) Higher atomic number elements have sharper central peak

Angular Dependence of SE's



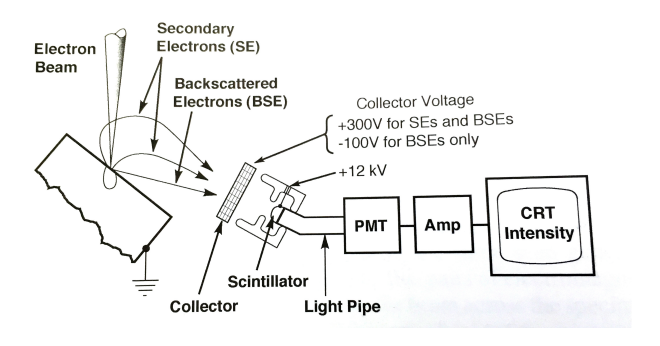
Goldstein et al, 2003

SE's also give topographic information



Detection of BSE's, SE's

Everhart-Thornley (ET) Detector



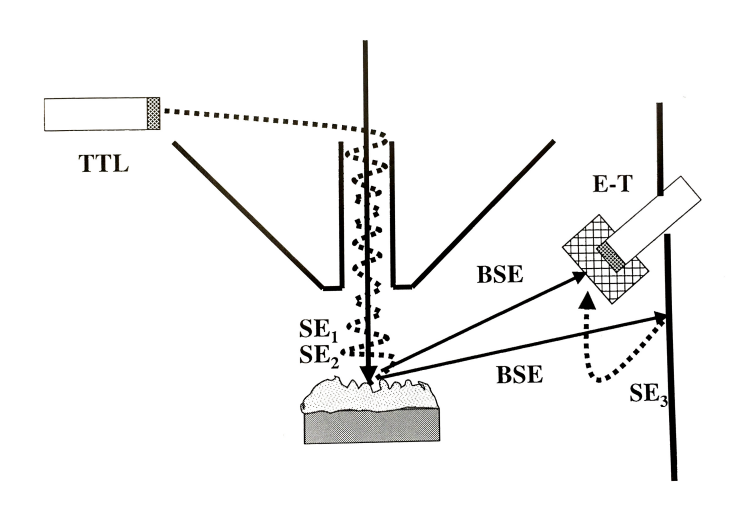
Electrons strike scintillator, releasing photons
Photons travel to photomultiplier tube
Eventually converted to electric signal, storing intensity values
Combined SE/BSE detector

Apply bias

- Bias can be applied to detector, directing electrons toward or away from it
 - Negative bias (< -50 V): detect only BSE's
 - Positive bias: collect more SE's, indirect BSE's: greater total signal

Positive bias

Through-Lens Detector (TTL)



SEM summary

- Images formed by scanning points across sample
- For higher resolution, want to minimize both probe size and interaction volume
 - Low voltage operation
 - But still need enough signal for detection
- Apply negative bias to detect mainly BSE's
- Backscattered imaging gives elemental contrast
- Secondary imaging gives more signal and topographic images
- Through Lens Detector for better resolution

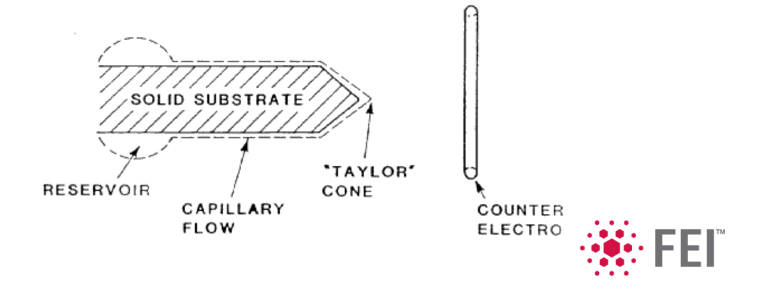
Biological Imaging

- Biological specimens are mostly light elements: little elemental contrast
- Standard procedure, as with negative stain TEM, is to stain with heavy metal salts (lead, uranium, tungsten) and look at the stain
- Long procedure involving:
 - Tissue fixation
 - Substitution of water with organic solvent
 - Infiltration with resin
 - Staining of biological components
 - Polymerization of resin
- Worked out over past 50 years, many protocols for different cells, tissues, organelles
- Can get very fine ultrastructural detail

FIB Operation

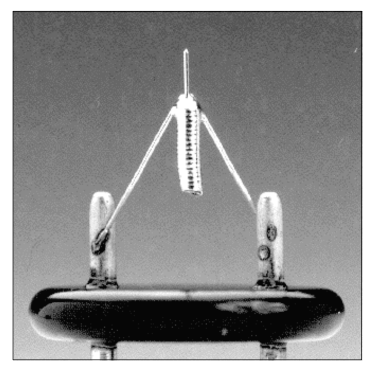
Basic Mechanism

- Liquid Flow from Reservoir
- Ion Formation
- External Beam Interactions



Gallium is the Most Popular LMIS

- A liquid metal
- Room temperature operation
- Long lived (500-1500 hr sources)
- High vacuum compatible
- Large ion for sputtering





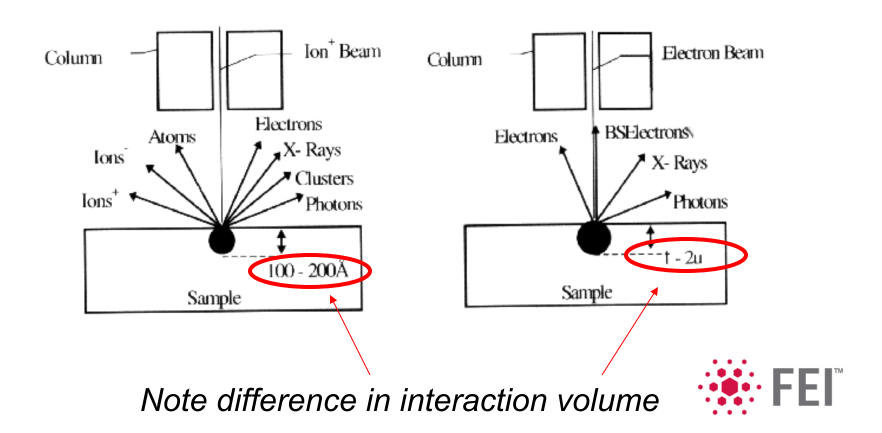
Ion Column

- Source LMIS at top
- Focusing Optics
 - Use Electrostatic lenses since ions are heavier than electrons.
- Deflection Electronics/Pattern Board
- High-speed Blanking
 - Need to prevent milling while blanking

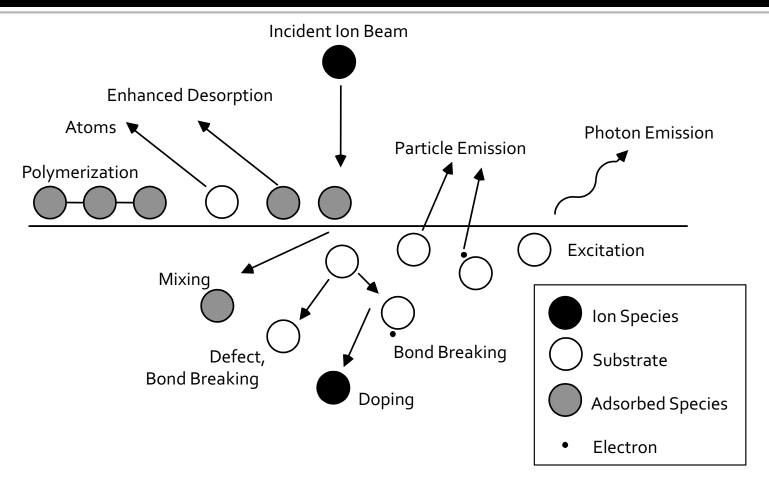


Using the System

Beam Interactions



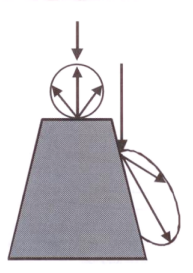
Ion Beam to Sample Interactions





Sputtered Particles

Sputtered Particle Ejection Behavior



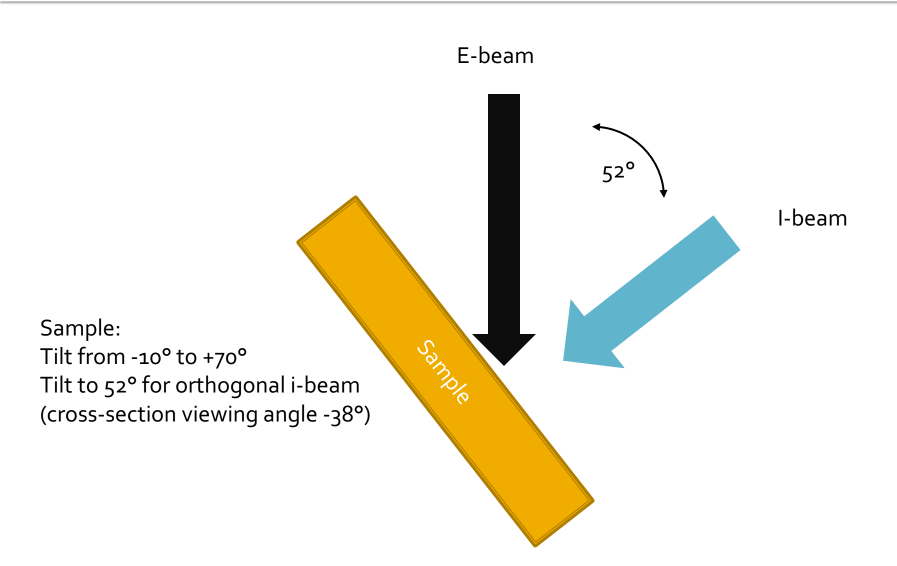


Geometry

E-beam I-beam Sample

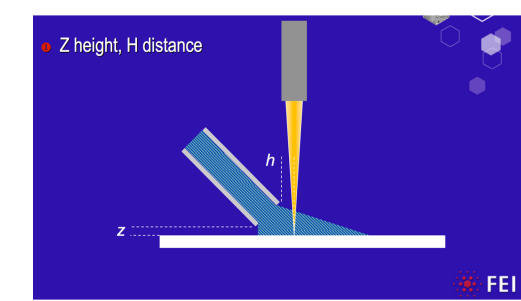
Sample: Tilt from -10° to +70°

Geometry



Deposition

- (Methylcyclopentadienyl) trimethyl platinum
- Warm to gas, spray over sample with needle
- I-beam or e-beam interactions break it apart, deposit metal onto sample
 - Protection
 - Hard surface for mill



Applications to Resinembedded tissue

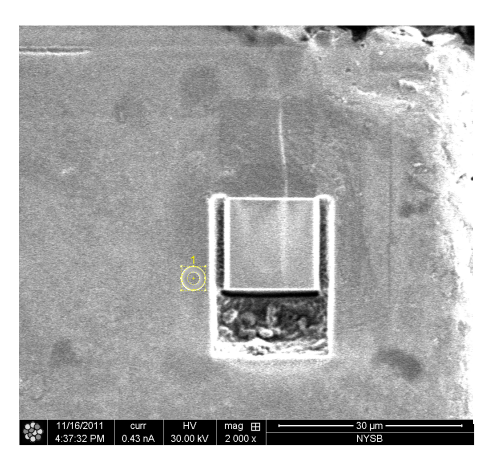
Tissue or Cells

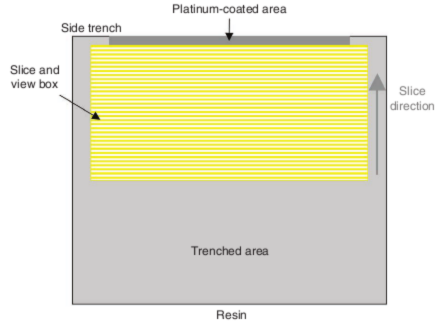
- Sample Prep
 - High Pressure frozen (optional)
 - Chemically fixed, freeze substituted
 - Resin embedded
 - En bloc staining
 - OsO₄, U Ac, Pb citrate
 - Osmium impregnation (OTO)
 - Want to make samples more conductive, more heavily stained
 - Thin conductive layer (C, Pt, Au-Pd) coated just before insertion

Imaging conditions

- Low voltage (2 keV or less)
 - Want to image only the surface
 - Minimal depth penetration (slice as thin as 5 nm)
 - No topographic information
 - Elemental contrast (C vs Os)
 - Through-lens detector for highest resolution
 - BSE mode (positive bias)
 - Stained parts will show up as bright on dark

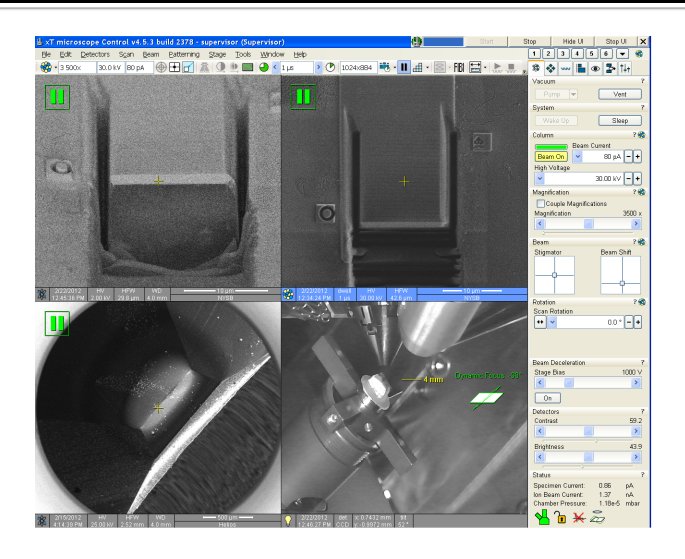
Milling Samples



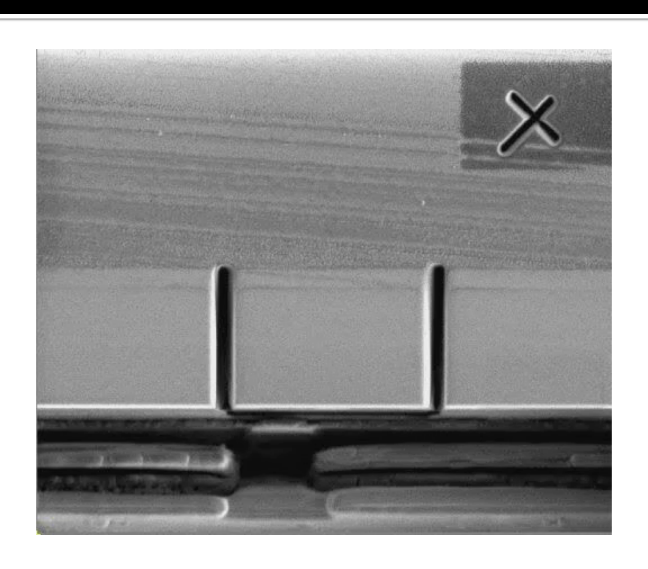


Bushby et al, 2011

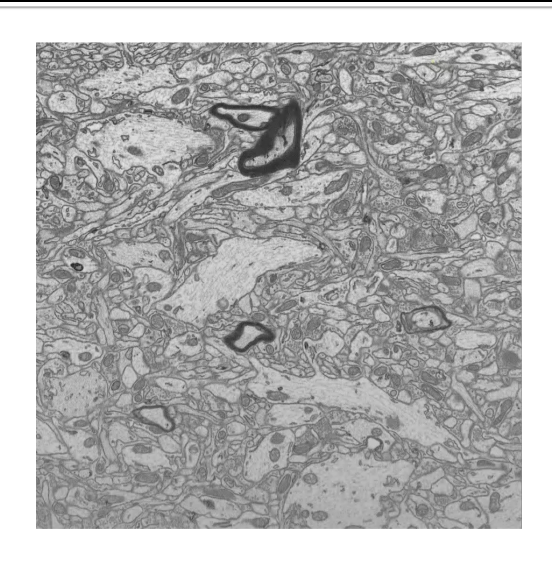
Set up for Slice and View



Milling: i-beam view



Example Movie: Neural Tissue

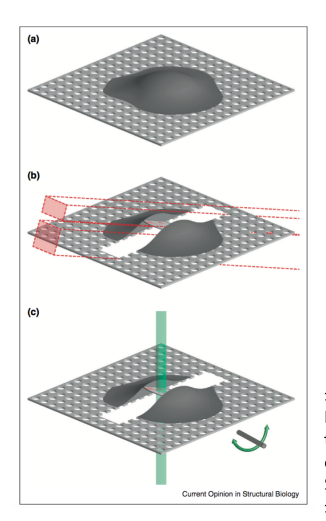


Ideal workflow

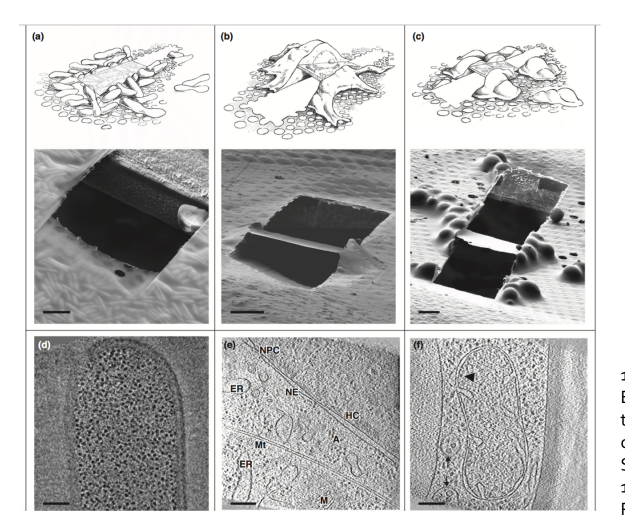
- Samples stained and embedded
- Thin slice for overall map make easier to find features (LM or ultrathin EM)
- Face of block polished
- Set up for slice and view (1 day)
- Collect slices (1-5 days)
- Align, process (IMOD, Amira) (1 day)
- Segmentation (IMOD, Amira) (weeksmonths)
 - Neural network automation : EMAN 2.2

FIB/SEM for Cryo Prep

Mill a thin slice through a cell



1: Villa E, Schaffer M, Plitzko JM, Baumeister W. Opening windows into the cell: focused-ion-beam milling for cryo-electron tomography. Curr Opin Struct Biol. 2013 Oct;23(5):771-7. doi: 10.1016/j.sbi.2013.08.006. Review. PubMed PMID: 24090931.



1: Villa E, Schaffer M, Plitzko JM, Baumeister W. Opening windows into the cell: focused-ion-beam milling for cryo-electron tomography. Curr Opin Struct Biol. 2013 Oct; 23(5):771-7. doi: 10.1016/j.sbi.2013.08.006. Review. PubMed PMID: 24090931.

Setup for standard Lamellae Preparation

FEI autogrid loading station with stereo microscope

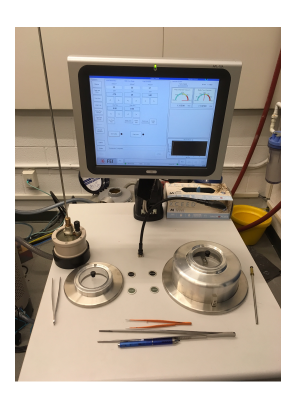


Shuttle with 2 grids



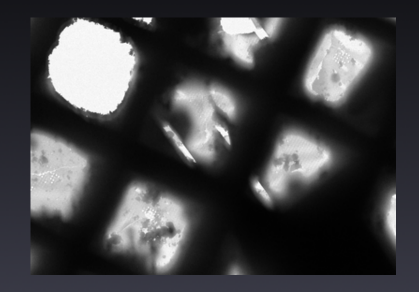


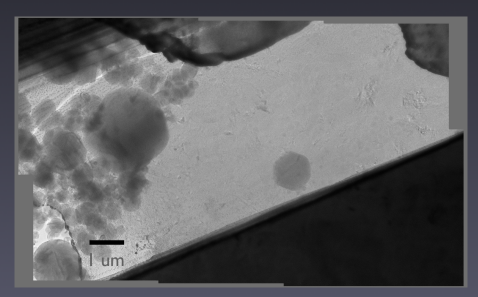
Quorum cryo loading station



Cutting windows into cells and tissues

Find lamella in TEM





Ideal workflow

- Cells grown on gold grid, then plunge frozen
- Image by cryo-LM to find features (1 day)
- Load into FIB/SEM, mill slices (1 day)
- Load into TEM, collect tomograms (1-2 days)
- Align, process (Protomo, IMOD) (1 day)
- Segmentation (IMOD, Amira)
- Sub-tomogram averaging

Cryo-SEM imaging

Technical Note

Cryo FIB-SEM: Volume imaging of cellular ultrastructure in native frozen specimens



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^b Cellular Neuroscience, Max Planck Institute of Experimental Medicine, Hermann-Rein-Straße 3, D-37075 Göttingen, Germany

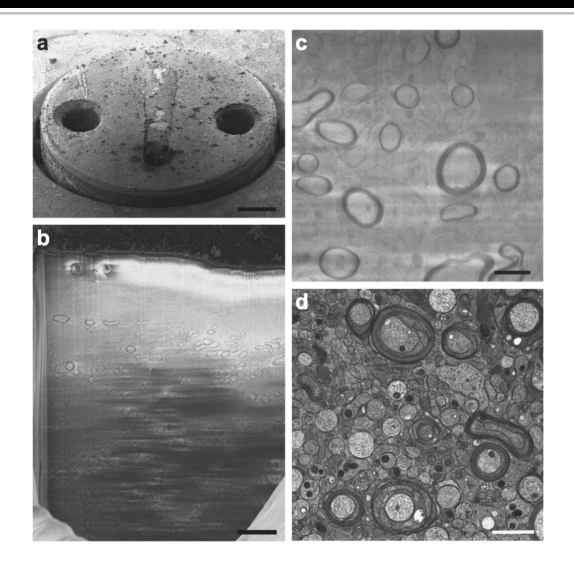
^c Department of Systemic Cell Biology, Max-Planck-Institute of Molecular Physiology, Otto-Hahn-Straße 11, D-44227 Dortmund, Germany

d Department of Neurogenetics, Electron Microscopy Facility, Max-Planck-Institute of Experimental Medicine, Hermann-Rein-Straße 3, D-37075 Göttingen, Germany

e Advanced Light and Electron Microscopy, Centre for Biological Threats and Special Pathogens, Robert Koch Institute, Nordufer 20, D-13353 Berlin, Germany

^fCenter for Nanoscale Microscopy and Molecular Physiology of the Brain (CNMPB), Göttingen, Germany

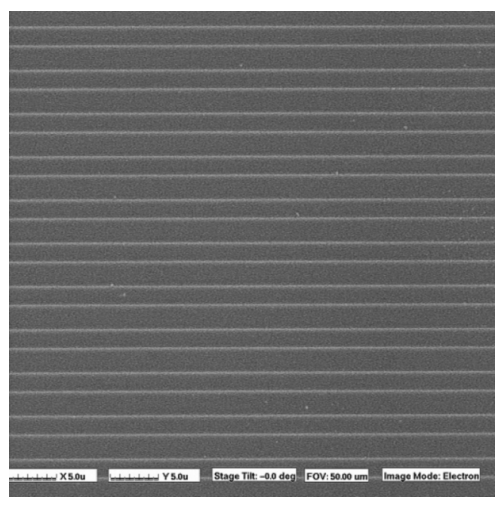
In-lens SE detector at 2.33 kV



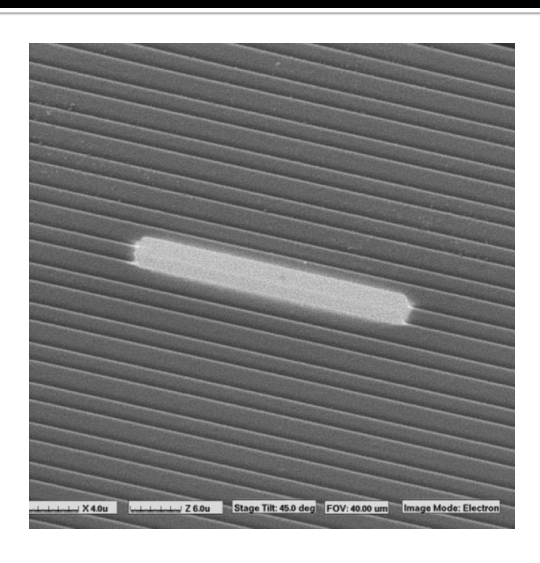
Lift Out

Use of FIB/SEM to prepare bulk material for TEM imaging

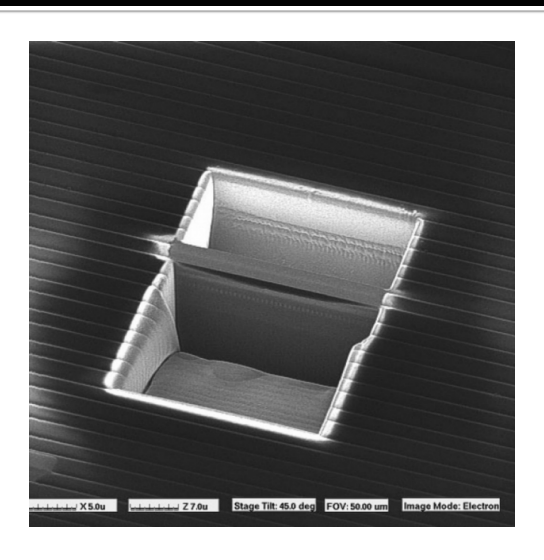
Area of Interest



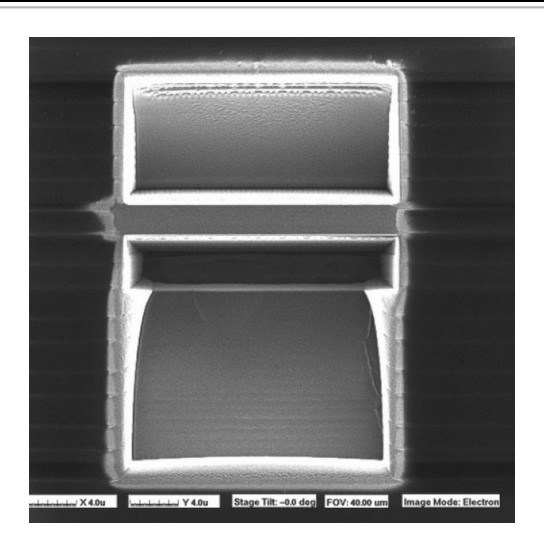
Protect Area of Interest



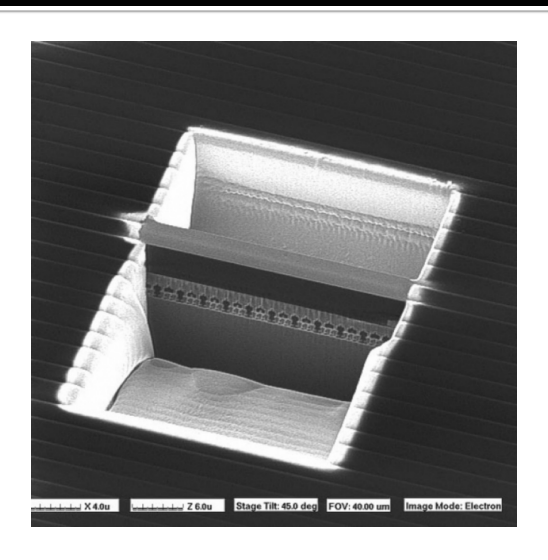
Mill Trenches around area



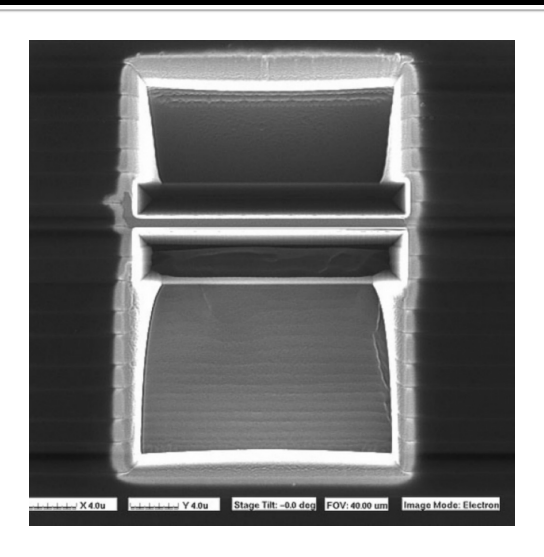
Top View of Trench



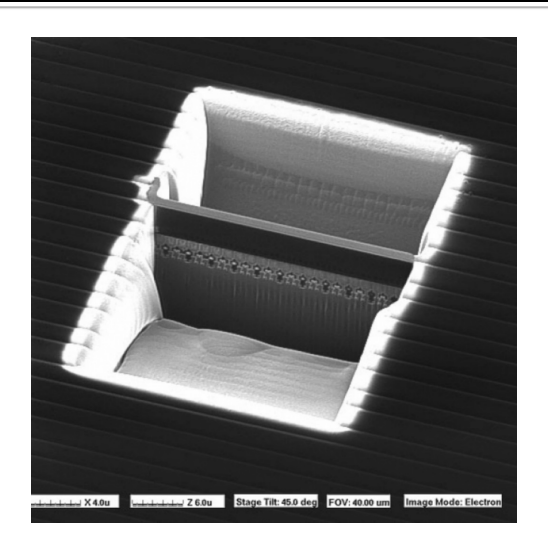
Polish Section



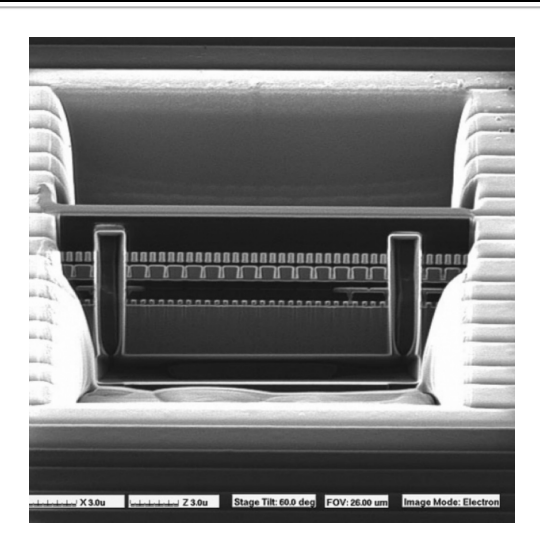
Further Polishing



Iso-View of Second Polish



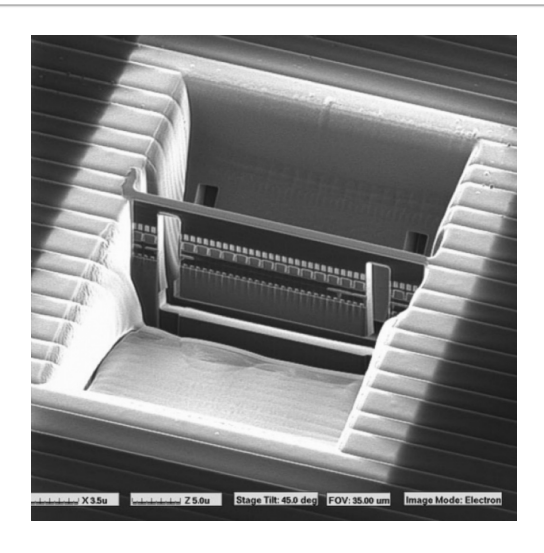
Frame Cuts to Define Area for Removal



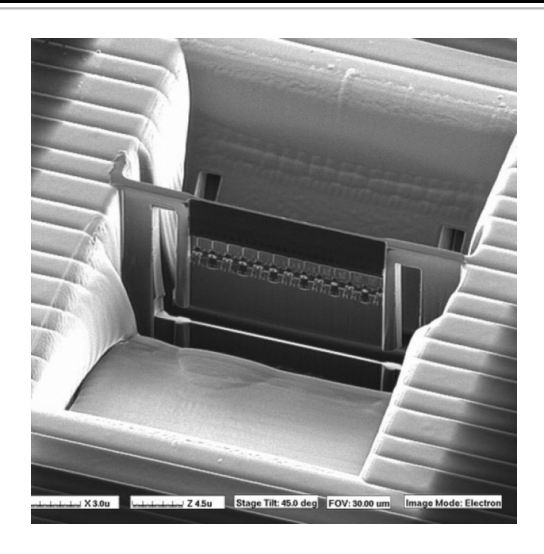
Top View of Frame Cut



Iso-View of Frame Cut



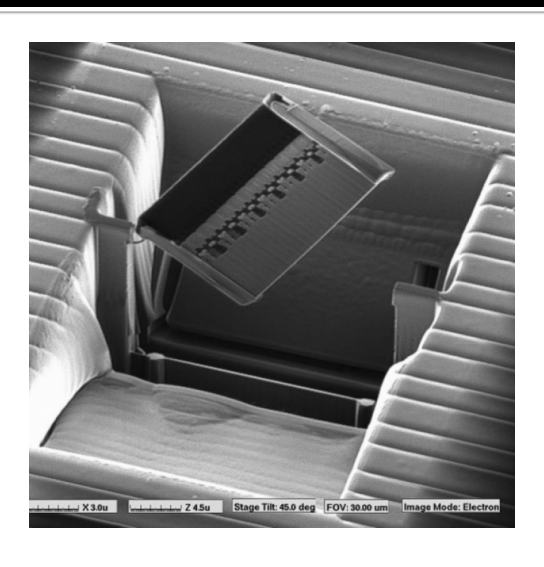
Thin to Electron Transparency



Top View just before Removal



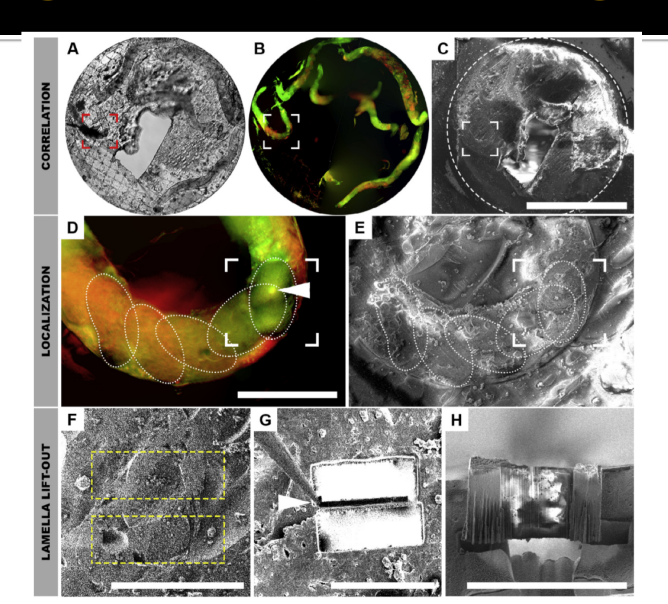
Remove Section and attach to manipulator (not shown)



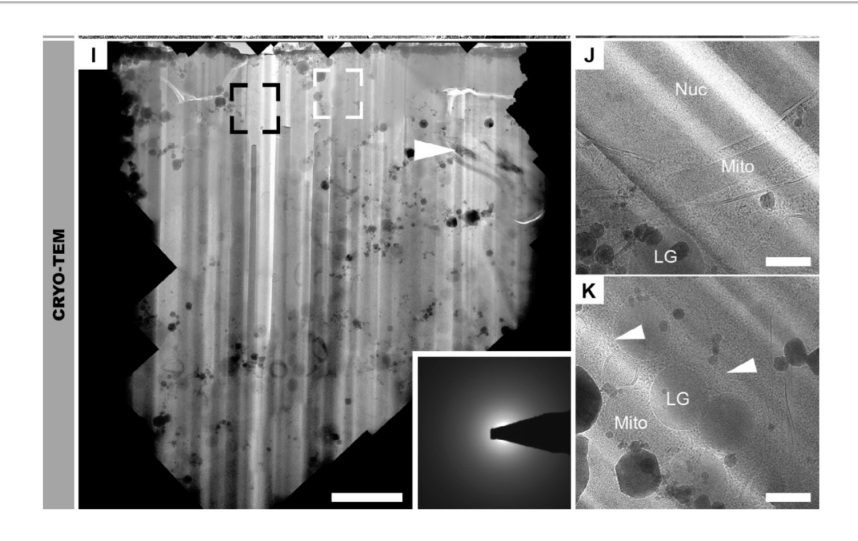
Applications to Cryo

Mahamid J, Schampers R, Persoon H, Hyman AA, Baumeister W, Plitzko JM. A focused ion beam milling and lift-out approach for sitespecific preparation of frozen-hydrated lamellas from multicellular organisms. J Struct Biol. 2015 Nov;192(2):262-9. doi: 10.1016/j.jsb.2015.07.012. PubMed PMID: 26216184.

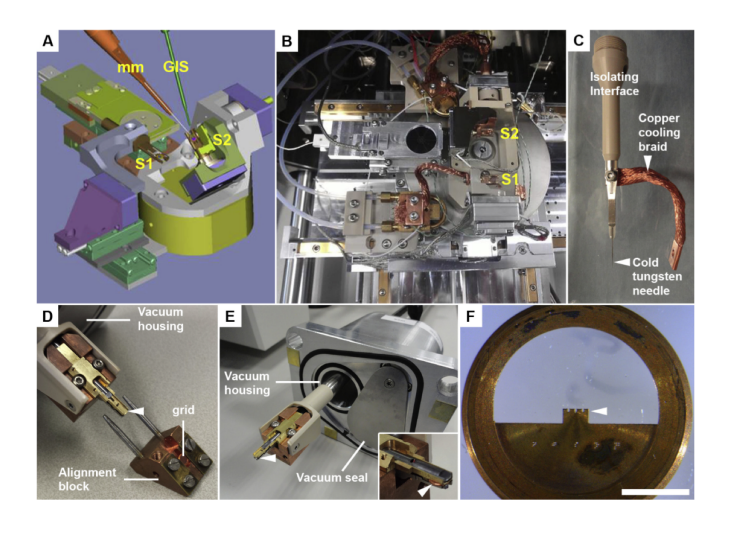
C Elegans Embryo HPF on grid



TEM



Equipment Needed



Questions