

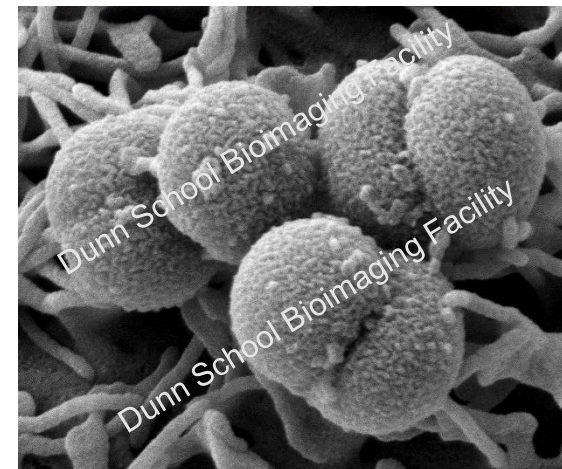
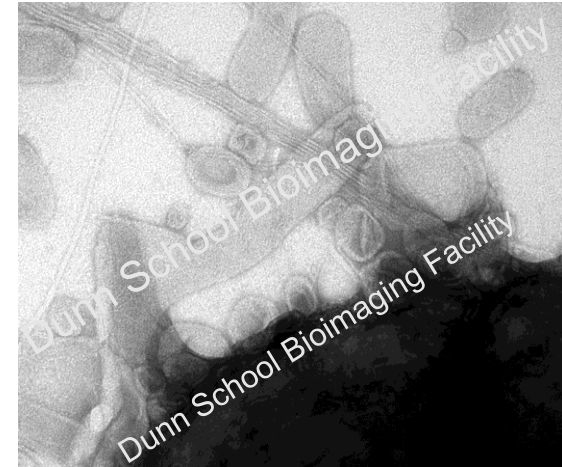


Applications of Electron Microscopy

Dr Errin Johnson

Head of Electron Microscopy

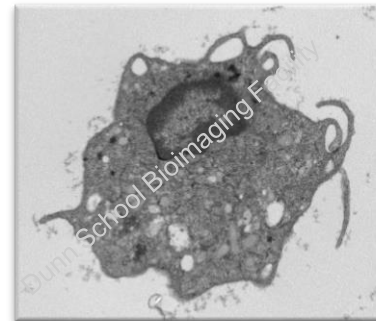
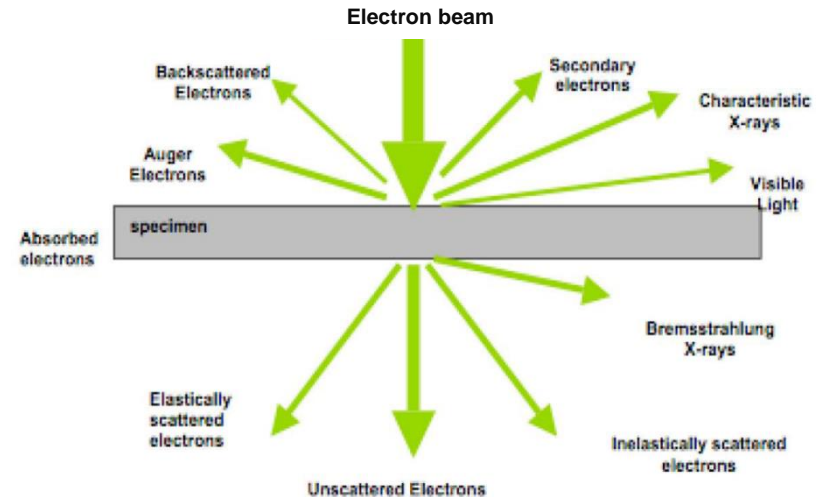
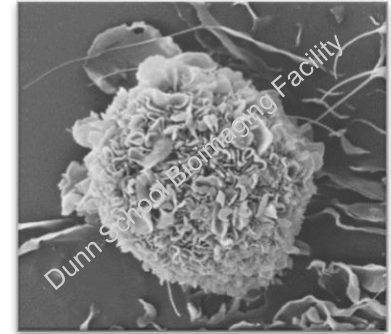
Sir William Dunn School of Pathology



Lecture Overview

- Introduction to Electron Microscopy (EM)
- Transmission Electron Microscopy (TEM)
 - Overview of the microscope
 - Biological specimen preparation for TEM
 - TEM applications
- Scanning Electron Microscopy (SEM)
 - Overview of the microscope
 - Biological specimen preparation for SEM
 - SEM applications
- Advanced EM techniques
 - Volume EM, protein localization, correlative microscopy & cryo-EM
- EM @ the Dunn School

Scanning Electron
Microscopy

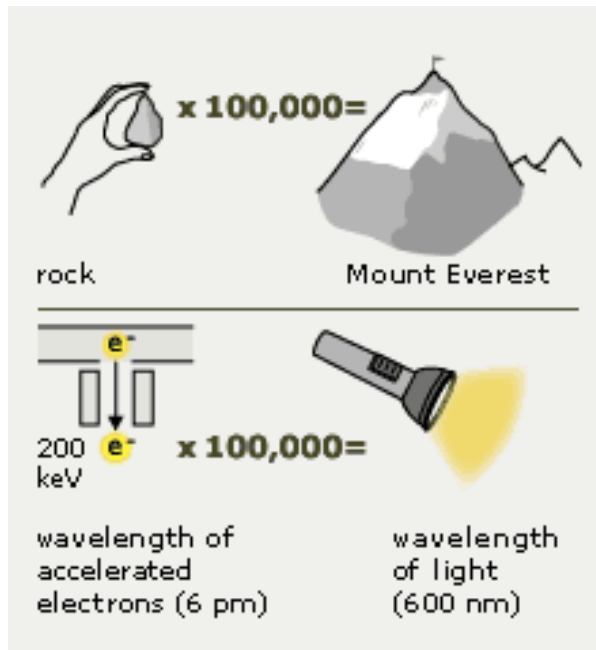


Transmission Electron
Microscopy

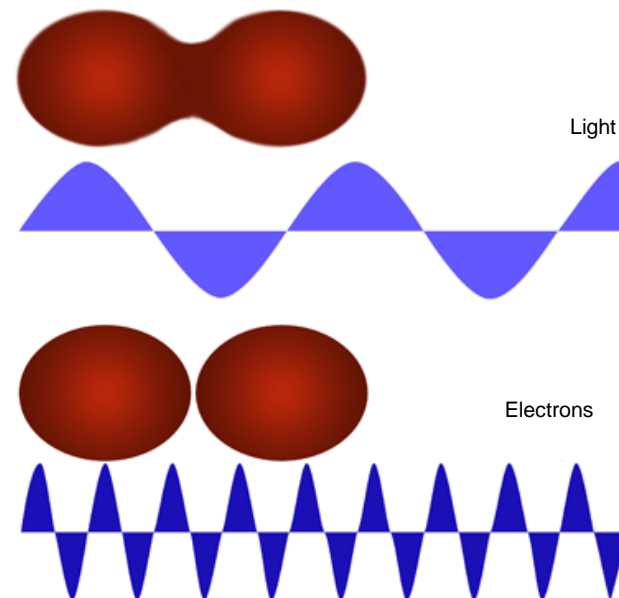
Introduction to Electron Microscopy

Resolution

- Resolution is the smallest distance at which two neighbouring points can be distinguished and is dependent on wavelength
- The wavelength of accelerated electrons is several order of magnitude shorter than that of light



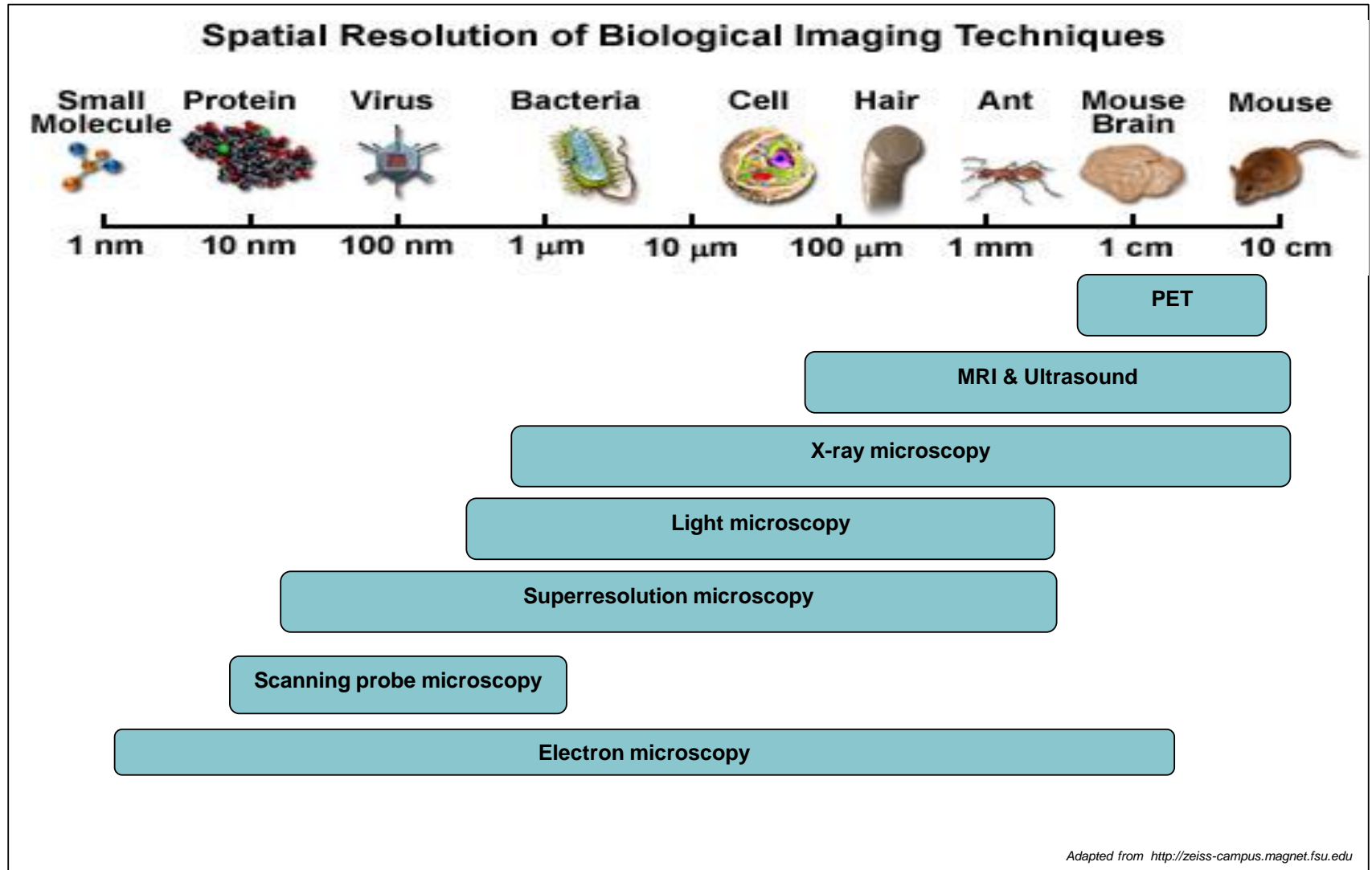
www.nobelprize.org



www.ammrf.org

Introduction to Electron Microscopy

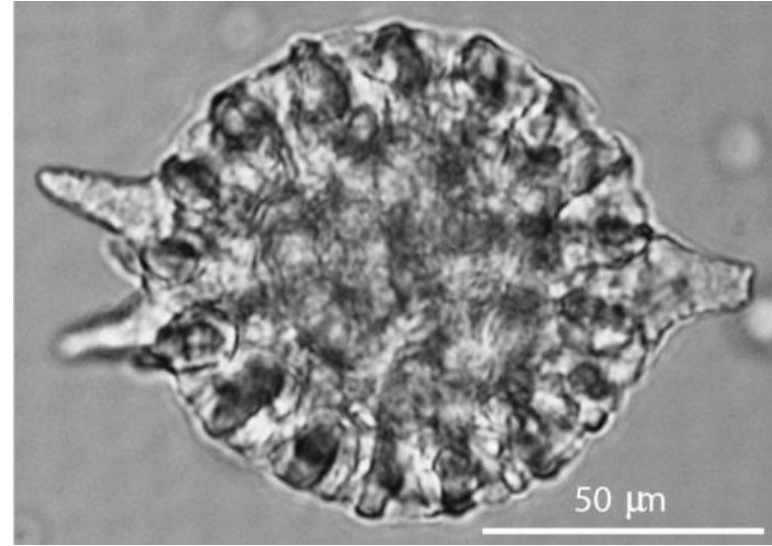
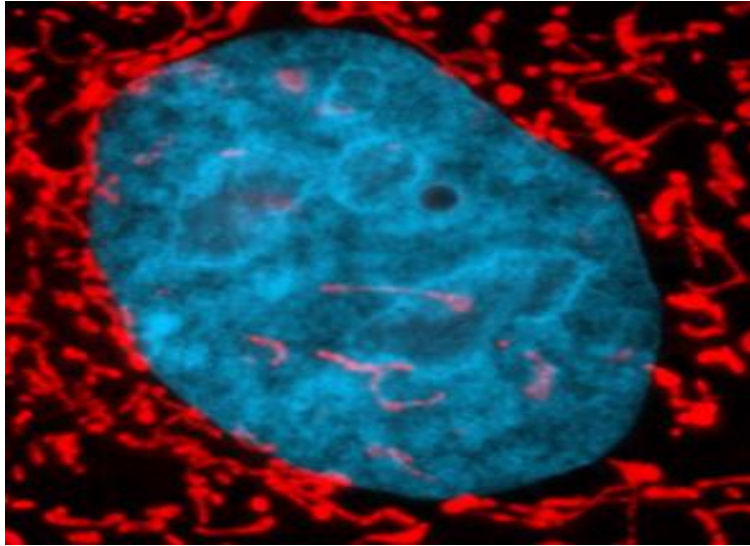
Resolution



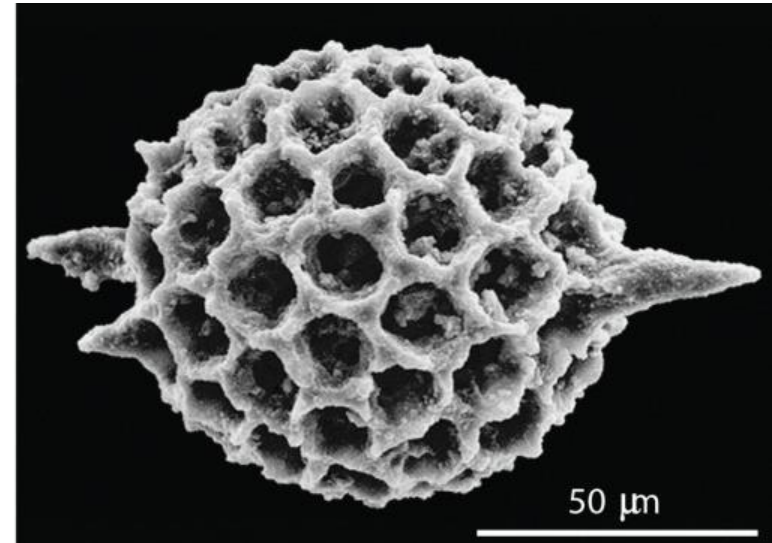
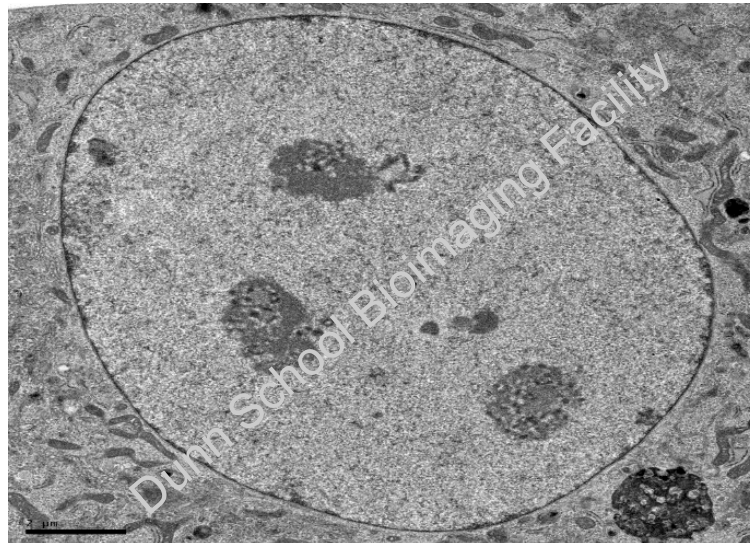
Introduction to Electron Microscopy

Resolution and contrast

Light microscopy



Electron microscopy



TEM image of fibroblast cell (E Johnson). Bottom: Confocal image of a kidney cell stained with DAPI (blue) and MitoTracker (red) (Hammamatsu.magnet.fsu.edu)

Radiolarian imaged with both SEM (top) and light microscopy (bottom). From: General Chemistry: Principles, Patterns, and Applications, B. Averill & P. Elderege

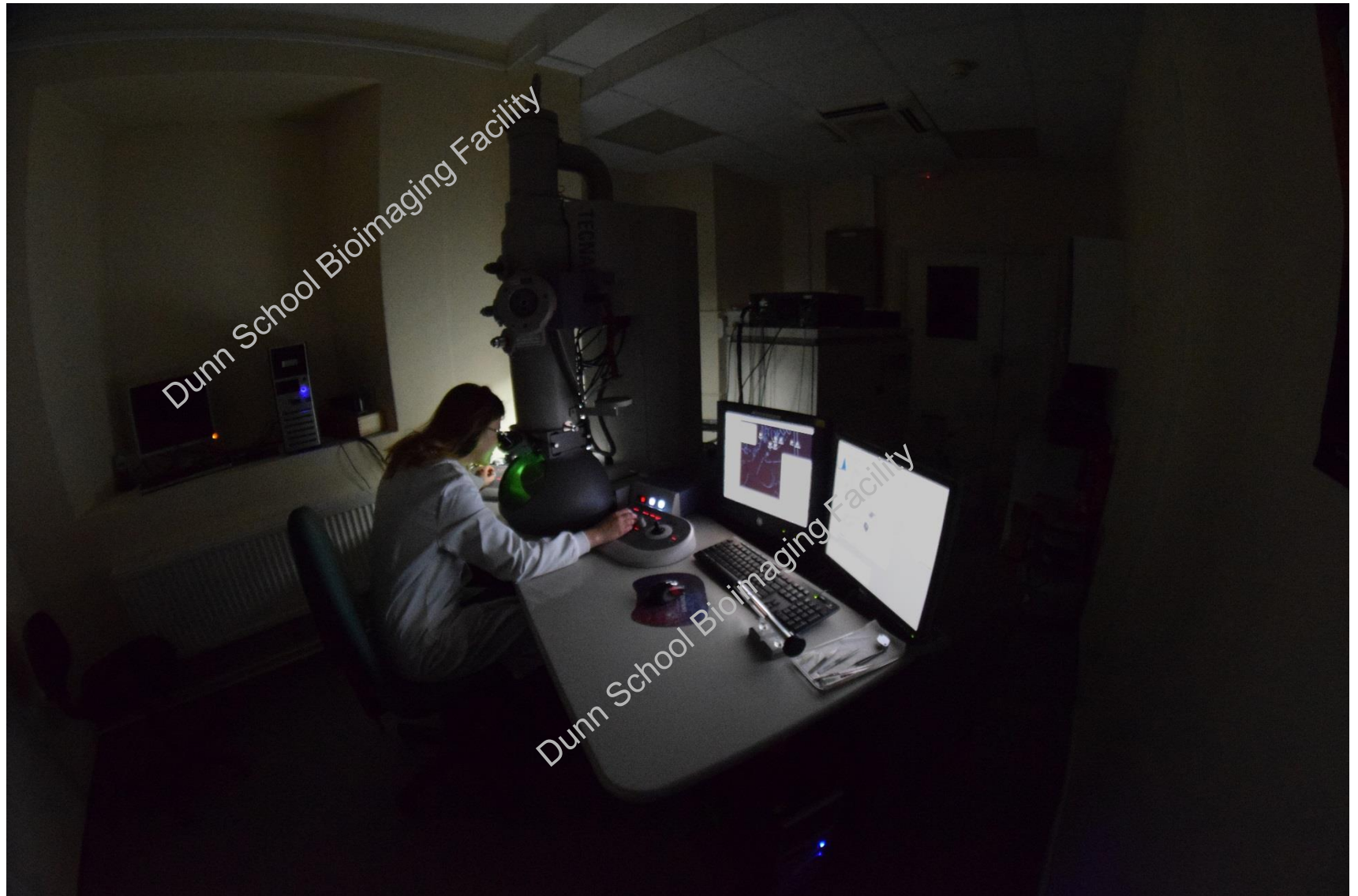
Introduction to Electron Microscopy

Electron microscopes

- Electron microscopes share the following main components:
 - An electron gun – source of electrons, operated at high voltages to accelerate the electrons
 - Electromagnetic lens system – to manipulate the electron beam
 - Vacuum system – protect beam integrity and prevent electrical discharging
 - Camera/detector – fast and sensitive signal detection
 - Computer – microscope control



Transmission Electron Microscopy (TEM)



The TEM

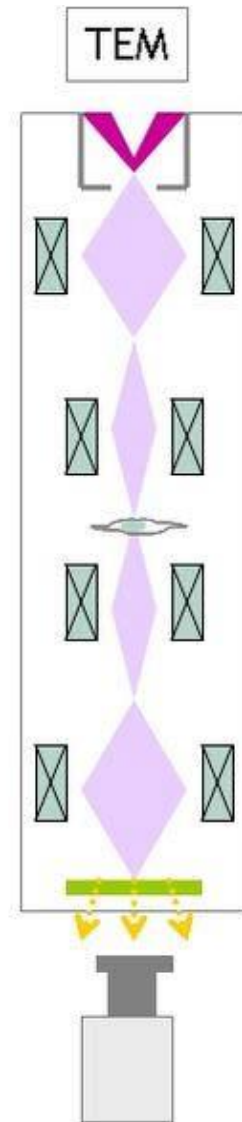
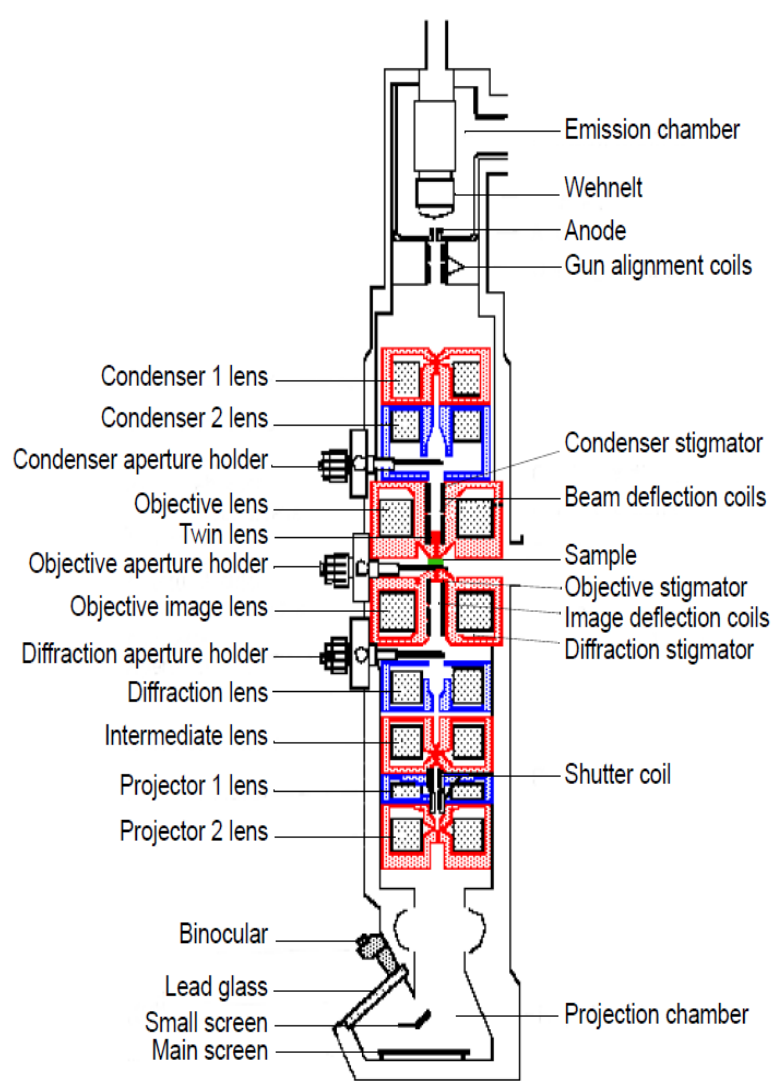
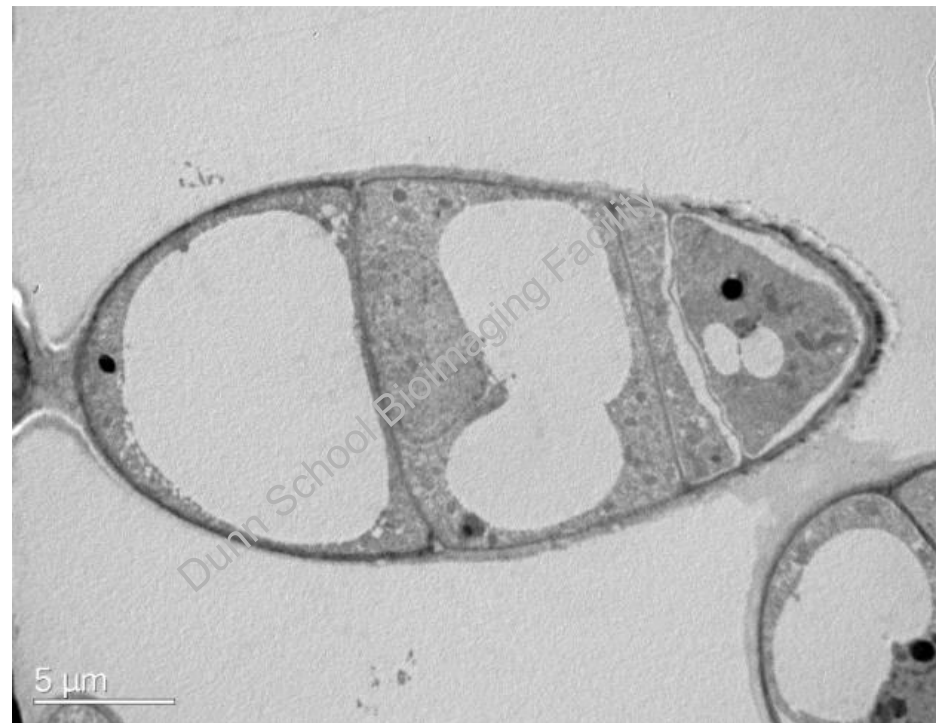
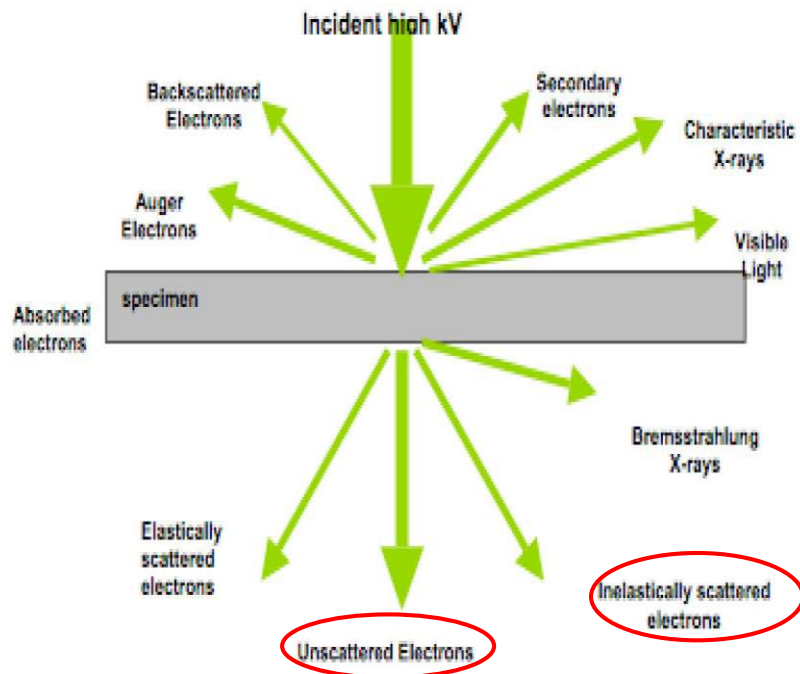


Image contrast in the TEM

- Contrast is generated by density differences within the sample.
- Darker areas in the image are where few electrons have been transmitted through the sample, due to thickness or high atomic number.



Lavender trichome, E Johnson

Biological TEM

Specimen requirements

TEM

Stable in the vacuum

Well preserved internal structure

Electron dense staining

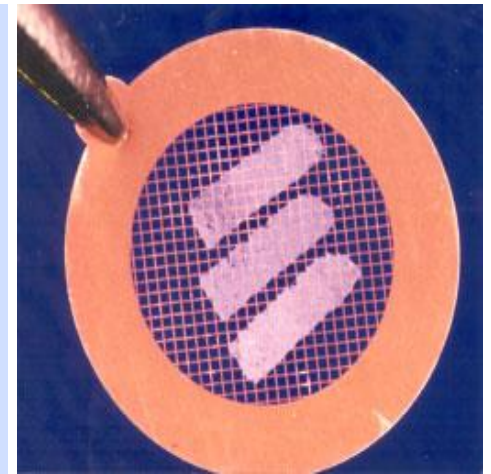
Very thin (eg: 70 nm)

Particulate samples can be stained
and viewed quickly

Cells and tissue require extensive
specimen preparation



*Mouse heart
~7 mm wide*

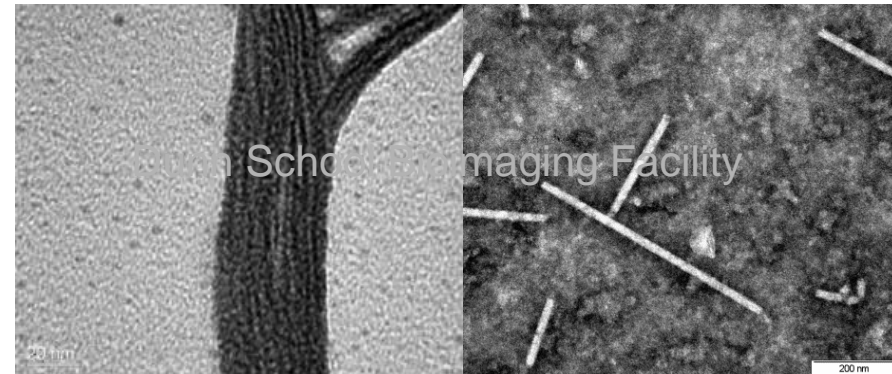
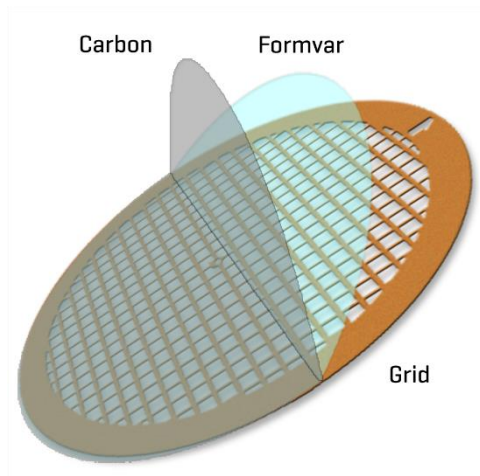
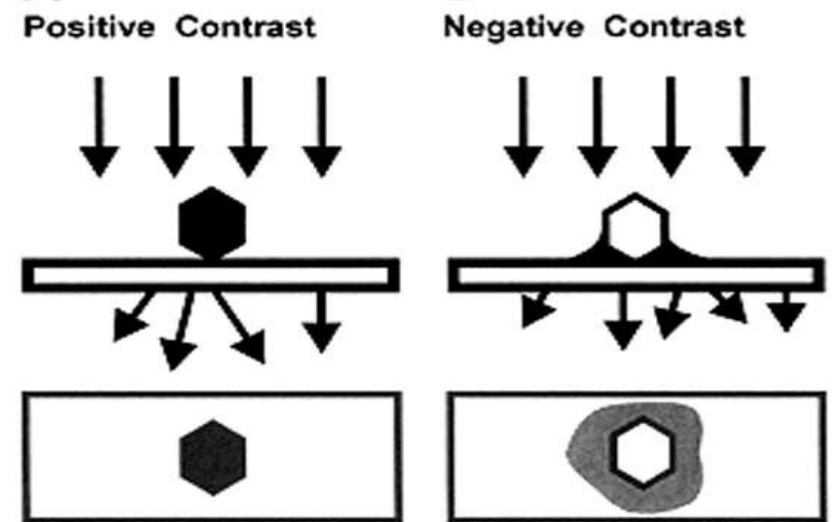


*70 nm thick resin-embedded tissue
sections on a TEM grid*

Specimen Preparation for TEM

Negative staining of particulate samples

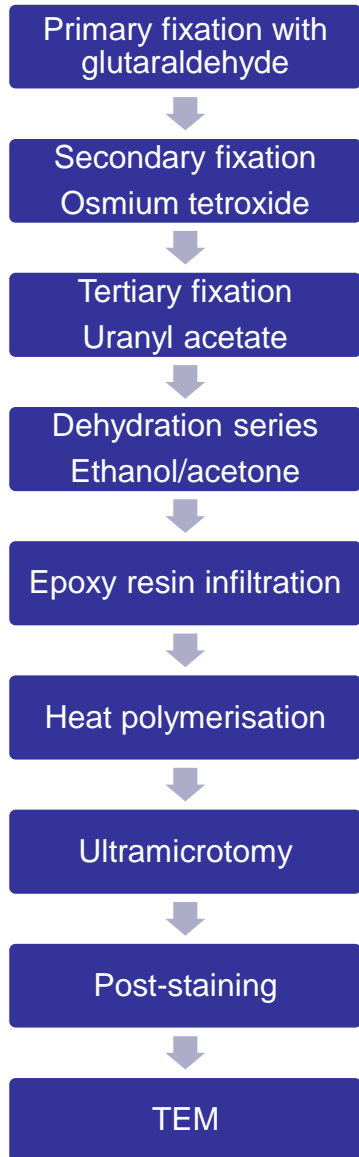
- For proteins, liposomes, DNA and viruses:
 - Coat grids with plastic film and carbon
 - Apply the particulate specimen
 - (eg: proteins, viruses, DNA)
 - Stain with heavy metal solution – this is most commonly uranyl acetate
 - Blot dry and view in the TEM



Bacterial protein stained with uranyl acetate; Tobacco mosaic virus negatively stained with sodium silicotungstate (E. Johnson)

Specimen Preparation for TEM

Standard protocol for cells and tissue



Estimated time required

Prep: 1 week

Ultramicrotomy: ~3-9 hrs

Microscopy: ~3-9 hrs

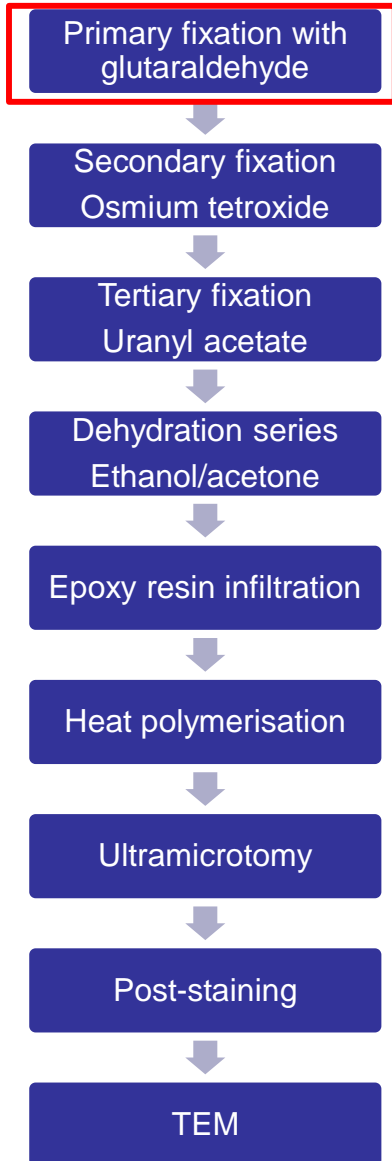
Data analysis: Dependent
on number of samples

Factors affecting prep quality

Type of fixative, mode of
fixation, type of heavy metals
used, tissue size and density
(vibratome sections or 1-2 mm³
pieces)

Specimen Preparation for TEM

Primary Fixation



- Fixation stops cellular processes and aims to preserve the specimen as close as possible to its natural state.
- Characteristics of a good fixative:
 - Permeates cells readily and acts quickly
 - Is irreversible
 - **Does not cause fixation artifacts**
- Methods of fixation include:
 - Chemical fixation with aldehydes
 - Immersion (cells, *Drosophila*)
 - Perfusion (mouse, rat)
 - Microwave-assisted
 - Cryo-fixation with liquid nitrogen

Standard chemical fixation:
2.5% glutaraldehyde +
2-4% PFA in 0.1M PIPES or
sodium cacodylate buffer

Specimen Preparation for TEM

Chemical fixation

Primary fixation with
glutaraldehyde



Secondary fixation
Osmium tetroxide



Tertiary fixation
Uranyl acetate



Dehydration series
Ethanol/acetone



Epoxy resin infiltration



Heat polymerisation



Ultramicrotomy



Post-staining

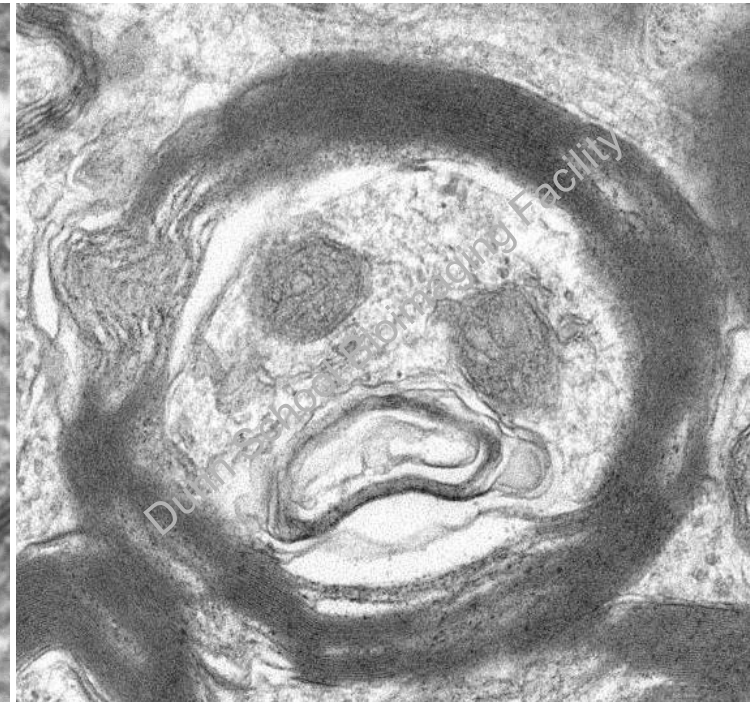


TEM

Good fixation



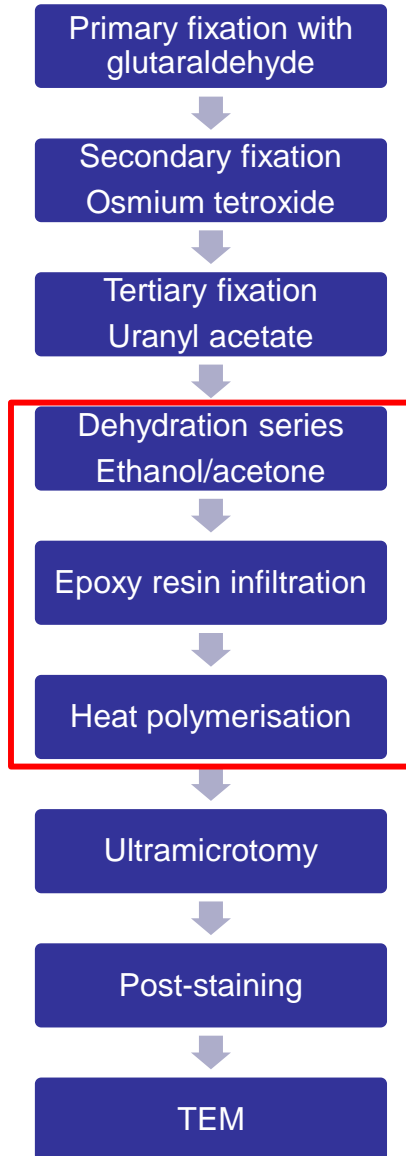
Bad fixation



Specimen Preparation for TEM

Dehydration & resin infiltration

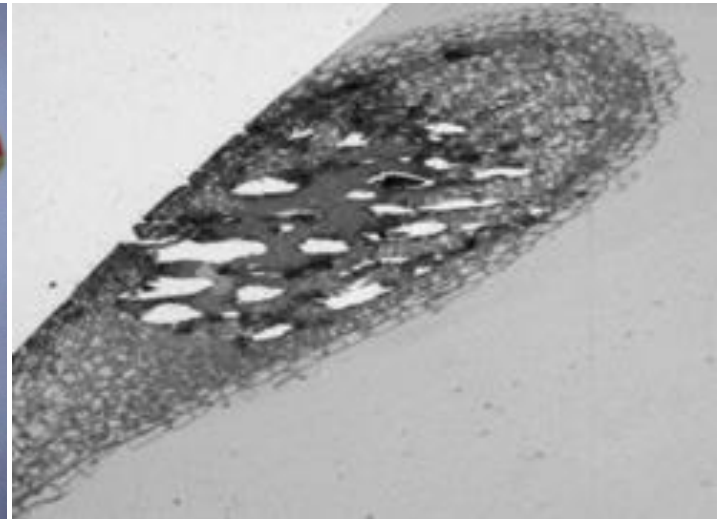
- Dehydration is the process of gradually replacing water in the sample with a solvent (usually acetone or ethanol).
- The solvent is then gradually replaced with resin. This process can be lengthy and depends on both the sample and type of resin used. Dense nerve tissue can be problematic.



Resin blocks



Poor resin infiltration



Specimen Preparation for TEM

Cryo-fixation and freeze substitution

- Tissue can be cryo-fixed using LN₂ in the High Pressure Freezer (Leica EM ICE, Leica EM PACT)



Estimated time req.

Prep: 1 week

Ultramicrotomy: ~3-9 hrs

Microscopy: ~3-9 hrs

Data analysis: Variable

Cryo-fixation
High pressure freezing



Freeze substitution
Osmium tetroxide + Uranyl acetate in acetone



Resin infiltration



UV/heat polymerisation



Ultramicrotomy

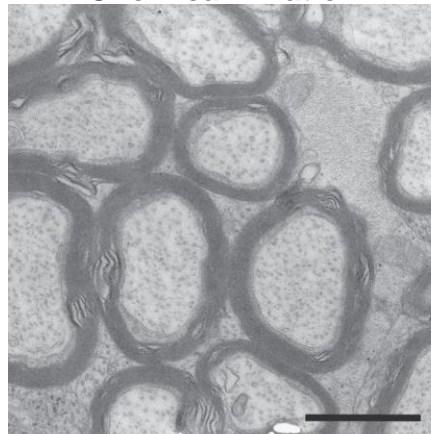


Post-staining

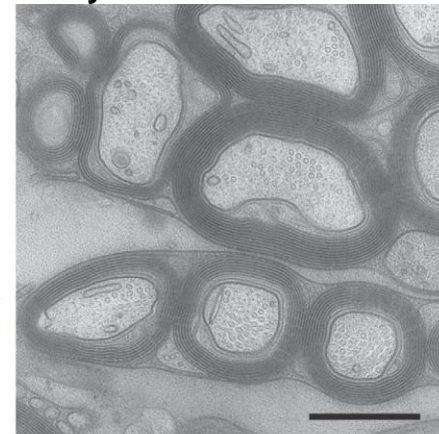


TEM

Chemical fixation

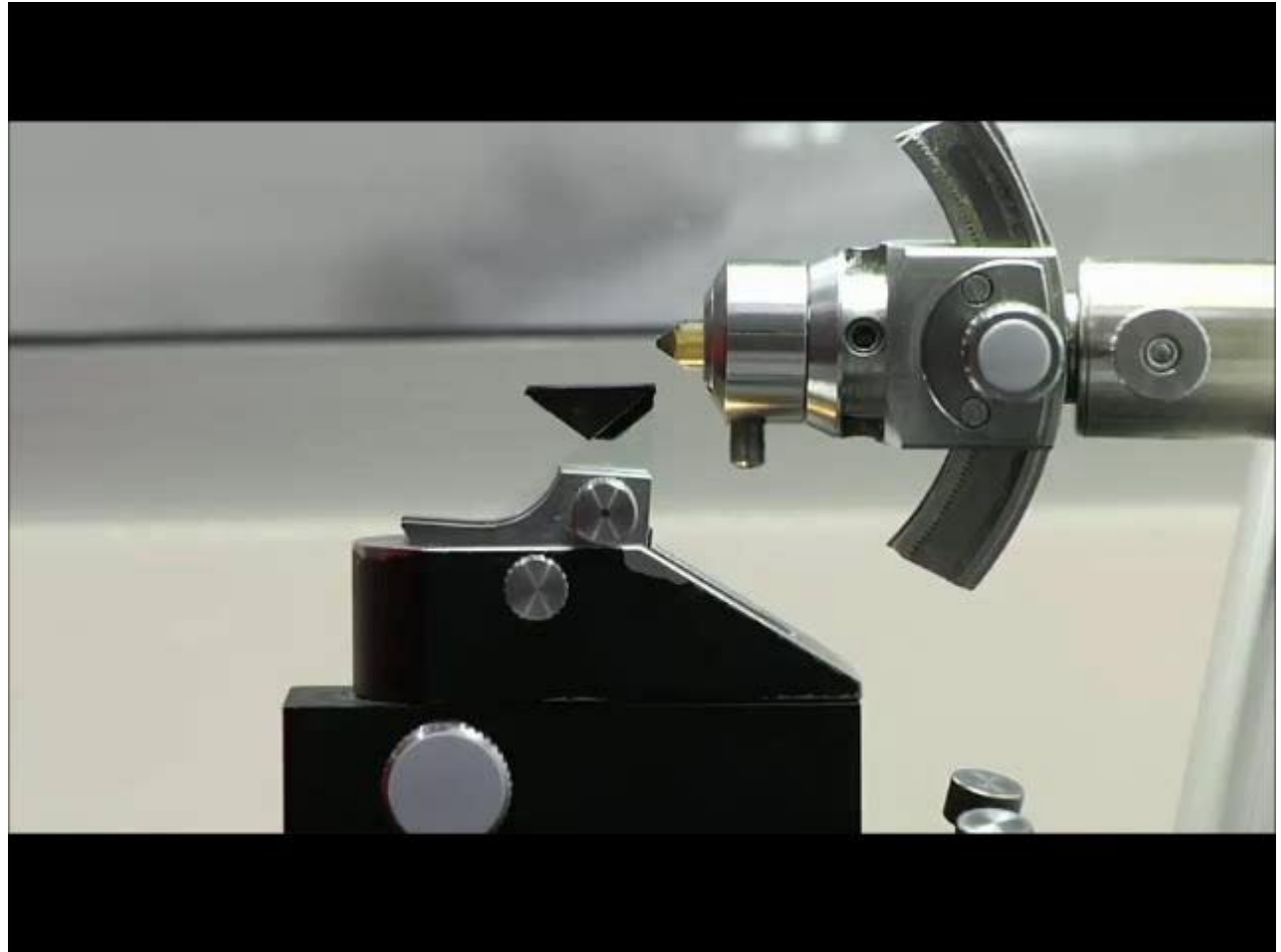
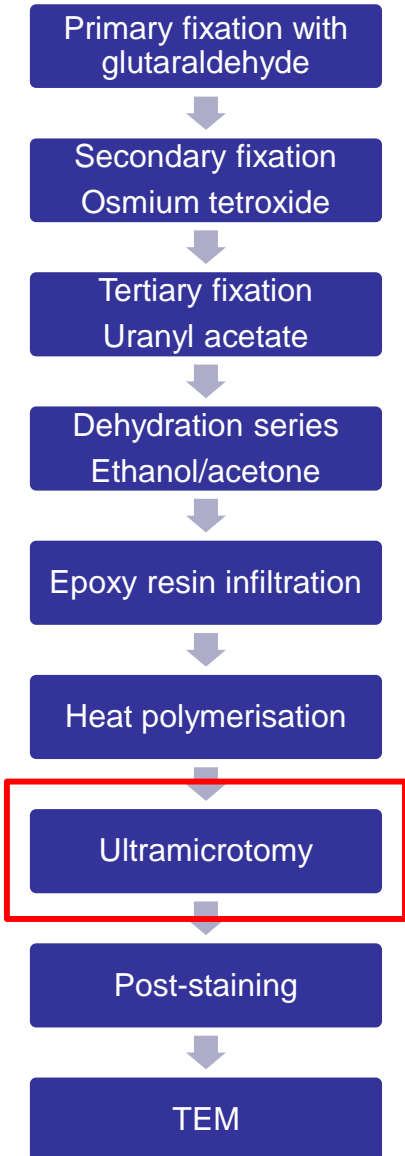


Cryo-fixation with HPF



Specimen Preparation for TEM

Ultramicrotomy



Introduction to ultramicrotomy video, University of Sydney

Electron microscopy techniques

Which one should I use?

- **Particles:**

- Screening (eg: checking purification, aggregation) -> TEM
- Protein localisation -> Immuno TEM
- Structural characterisation -> TEM, cryo-TEM

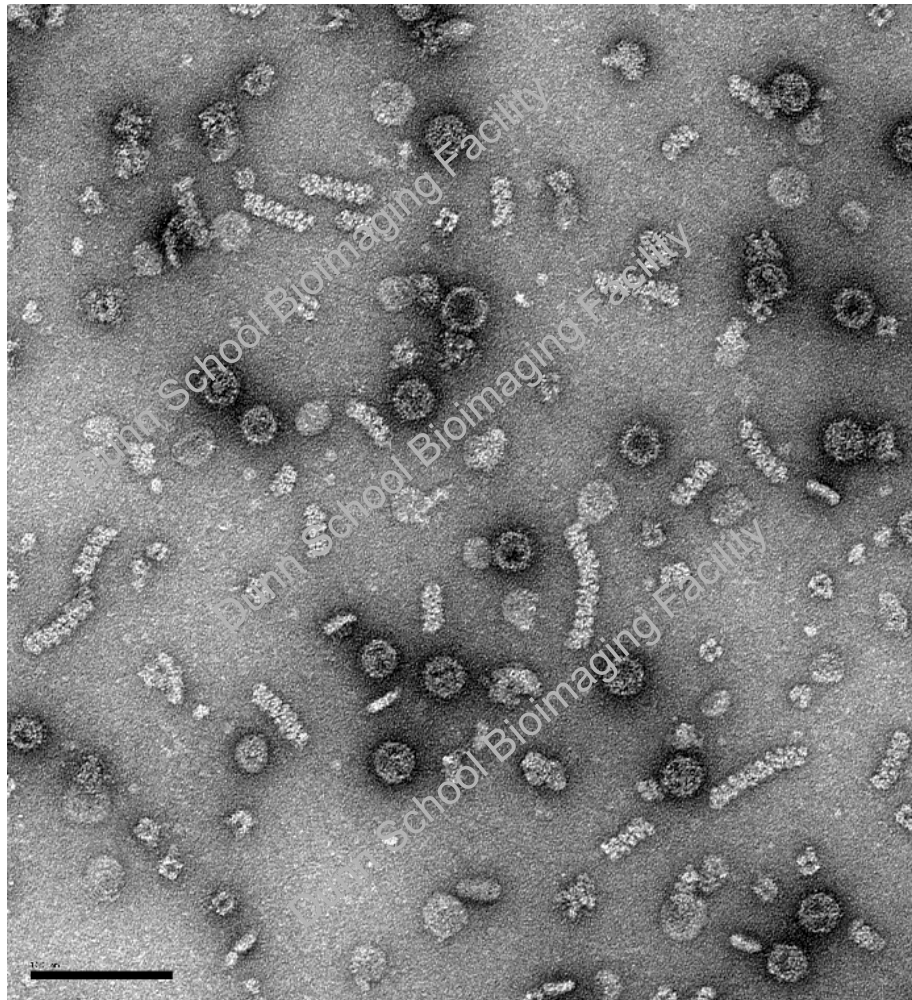
- **Cells and tissue:**

- Assessing ultrastructure in 2D (drug treatment, knockouts etc) -> TEM
- Assessing morphology (drug treatment, knockouts etc) -> SEM
- Assessing 3D ultrastructure -> Volume EM: serial sectioning TEM, 3View, FIB-SEM, EM tomography, array tomography
- Localising proteins of interest -> immuno TEM, EM genetic tags, CLEM
- Identifying specific cells -> immuno TEM, CLEM, EM genetic tags

Particulate samples

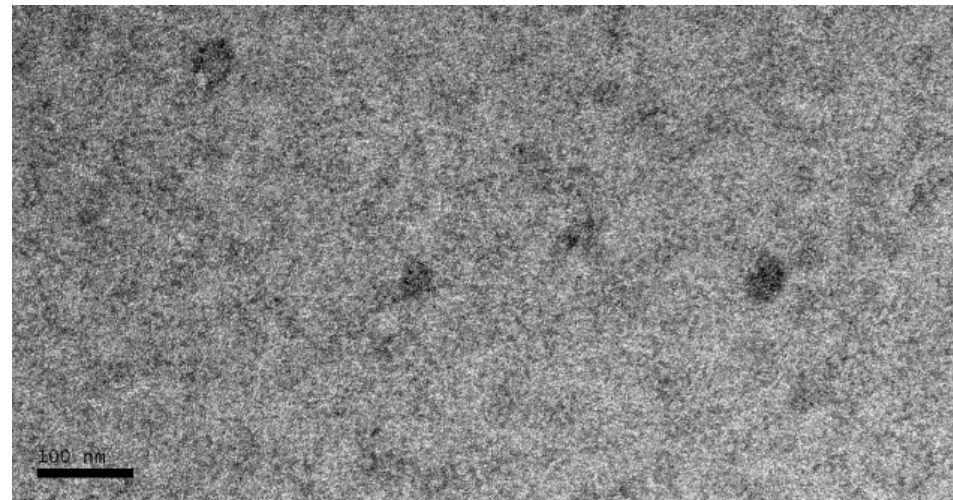
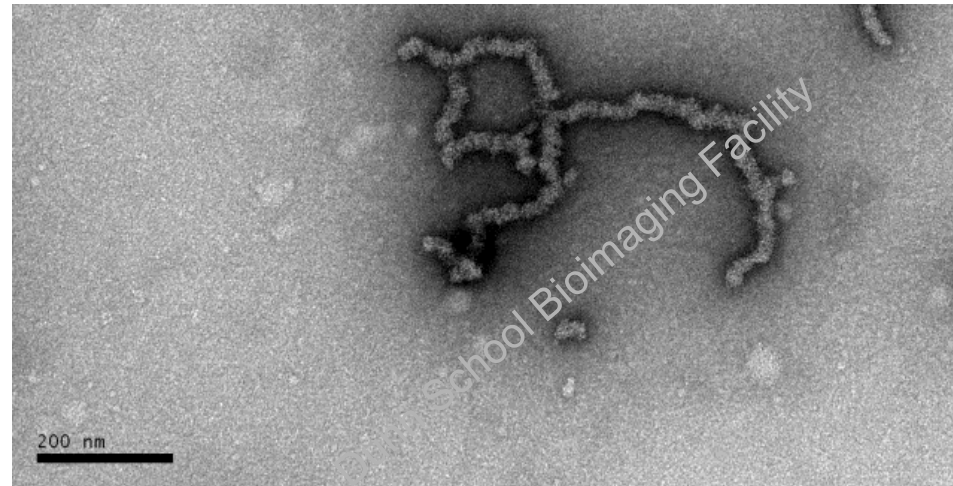
Screening preps using negative staining TEM

Viruses



*Negatively stained virus-like particles
(D Leneghan/E Johnson)*

Proteins

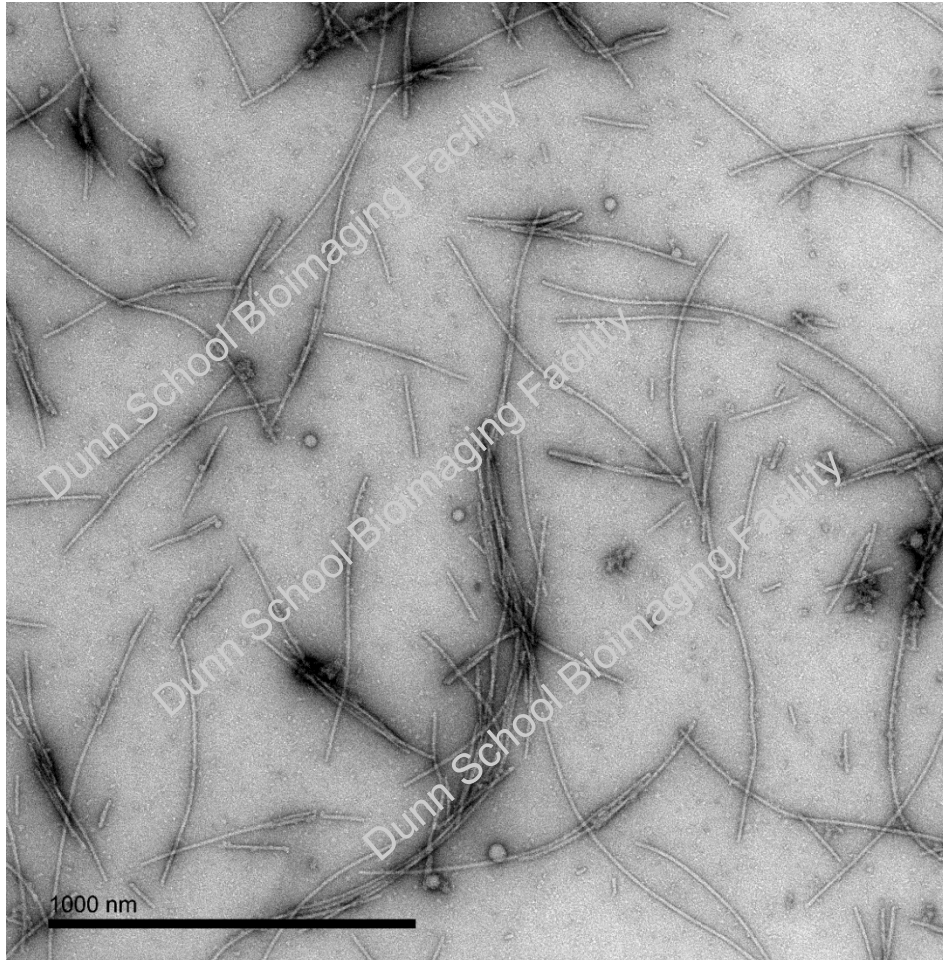


*SAS-6 protein WT aggregates (top) & mutant dimers (bottom)
(M Cottee/E Johnson)*

Particulate samples

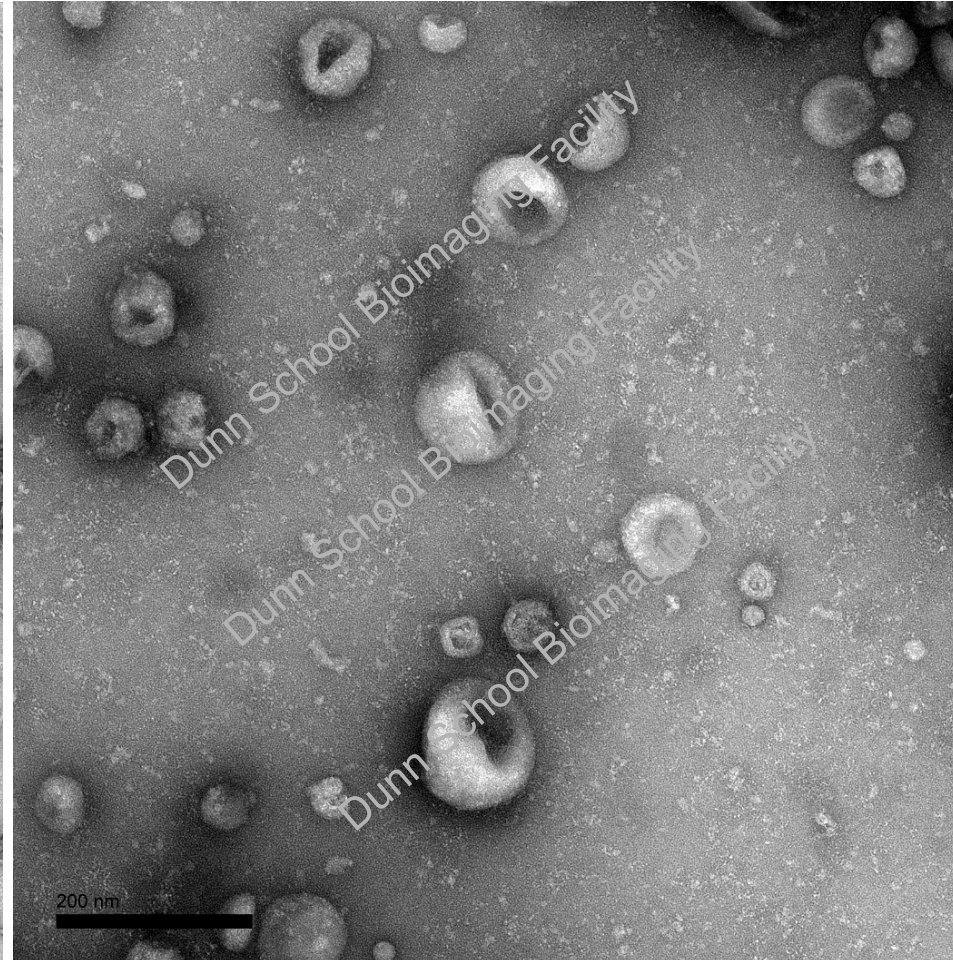
Screening preps using negative staining TEM

Fibrils



Negatively stained α -synuclein fibrils

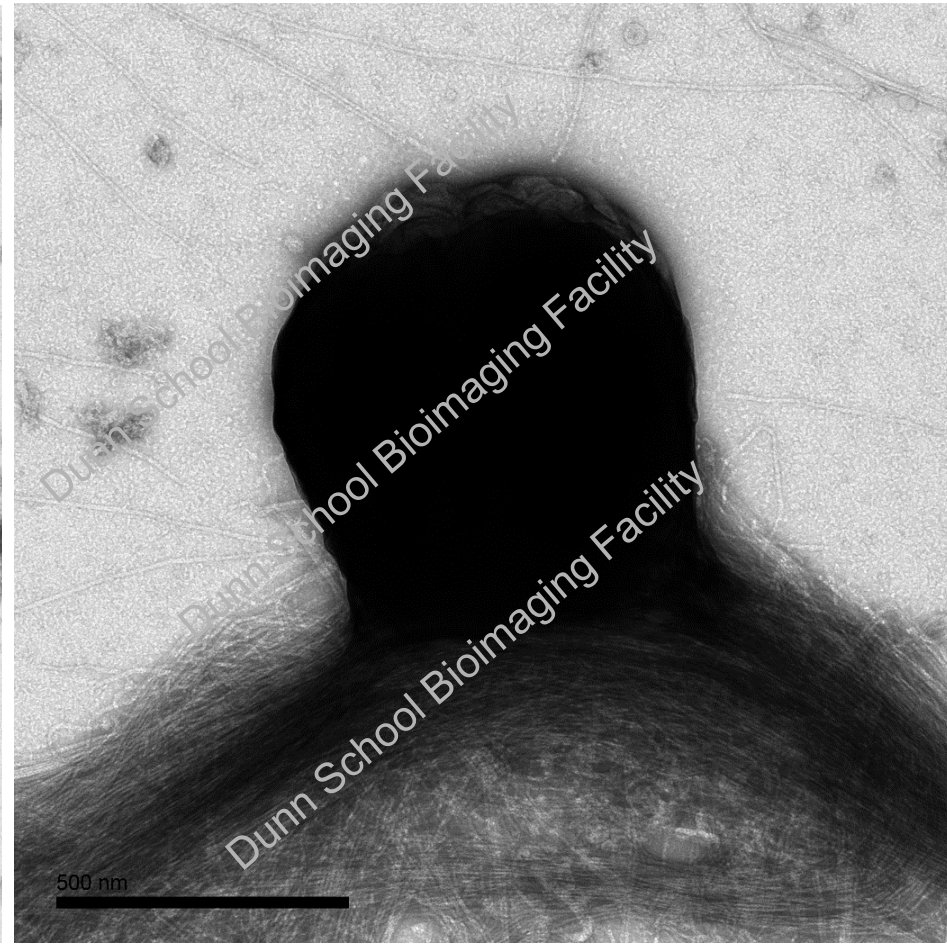
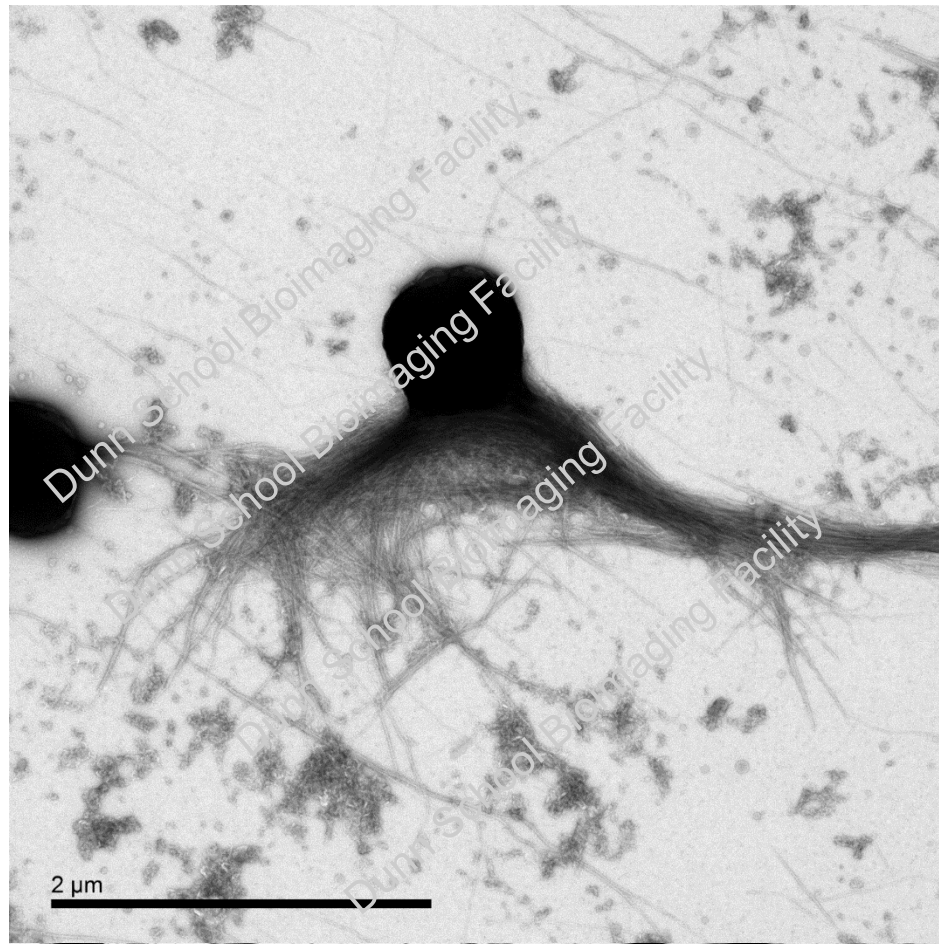
Vesicles



*Negatively stained exosomes
(H Rodriguez Caro/E Johnson)*

Particulate samples

Whole-mount negative staining TEM

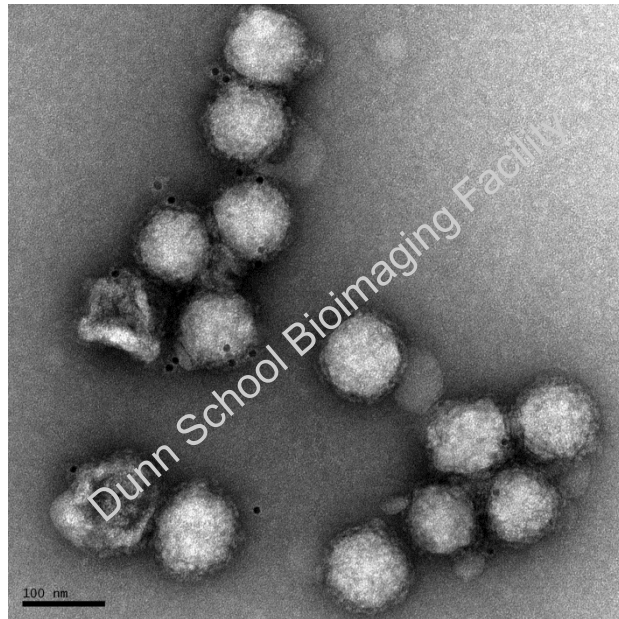


Negatively stained N. meningitidis (Tang lab/EJohnson)

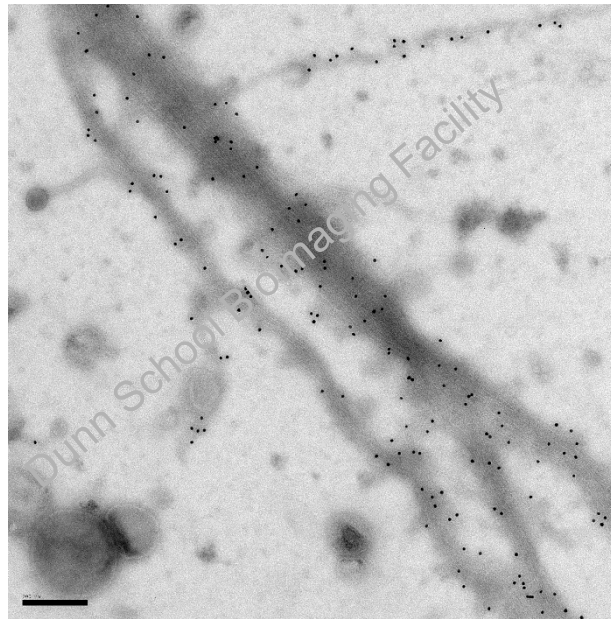
Particulate samples

Protein composition – immuno-negative staining

- As for immunofluorescence labelling, but the secondary antibody is conjugated to a small (1-4 nm) colloidal gold particle instead of a fluorophore



*Immunolabelled influenza
(Ed Hutchinson/E Johnson)*



*Immunogold labelled Type IV pili from Neisseria
meningitidis (M Woermann/E Johnson)*



*Whole mount immunolabelled Trypanosome
cytoskeleton (S Dean)*

Electron microscopy techniques

Which one should I use?

- **Particles:**

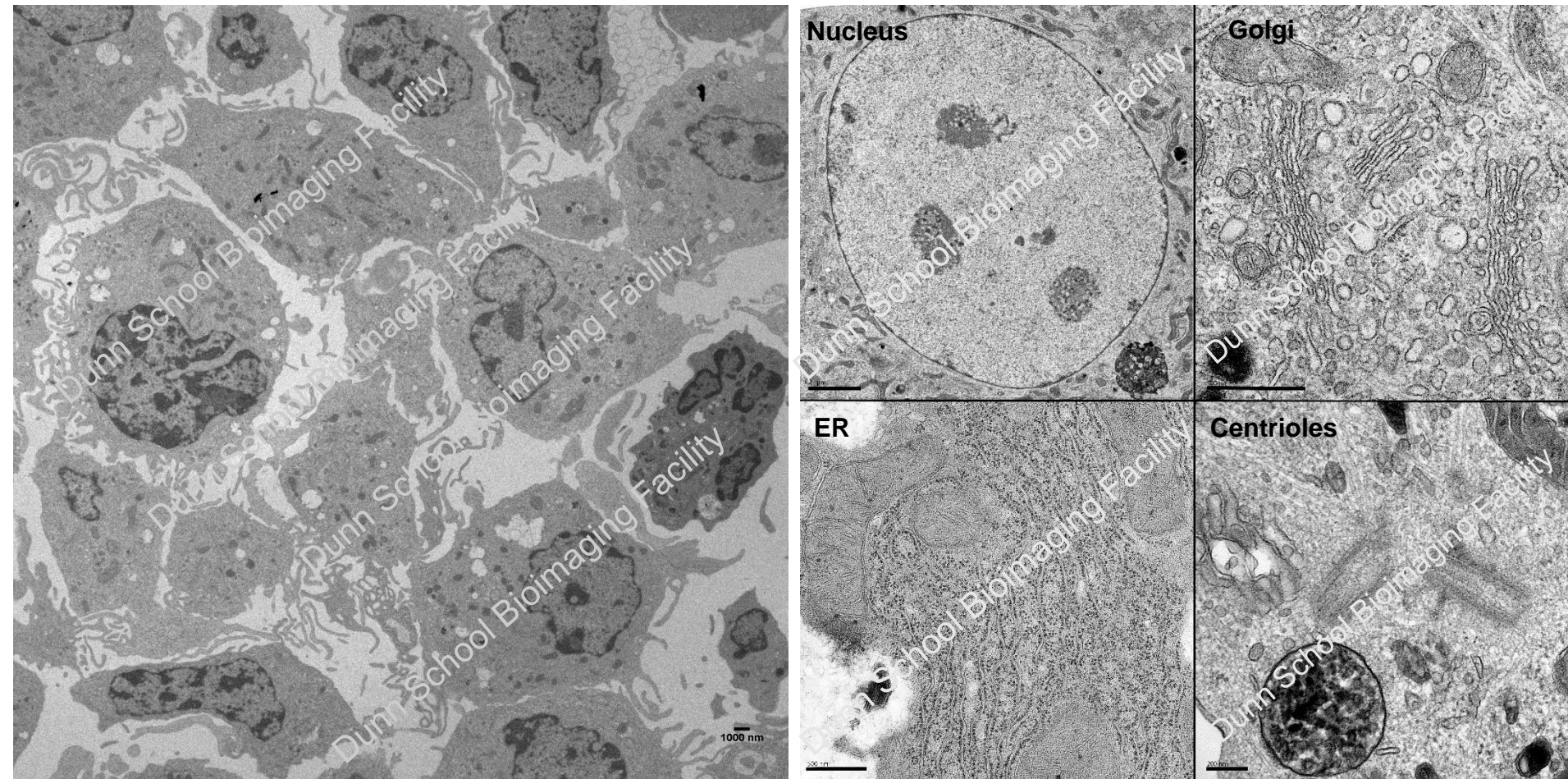
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TEM Ultrastructure

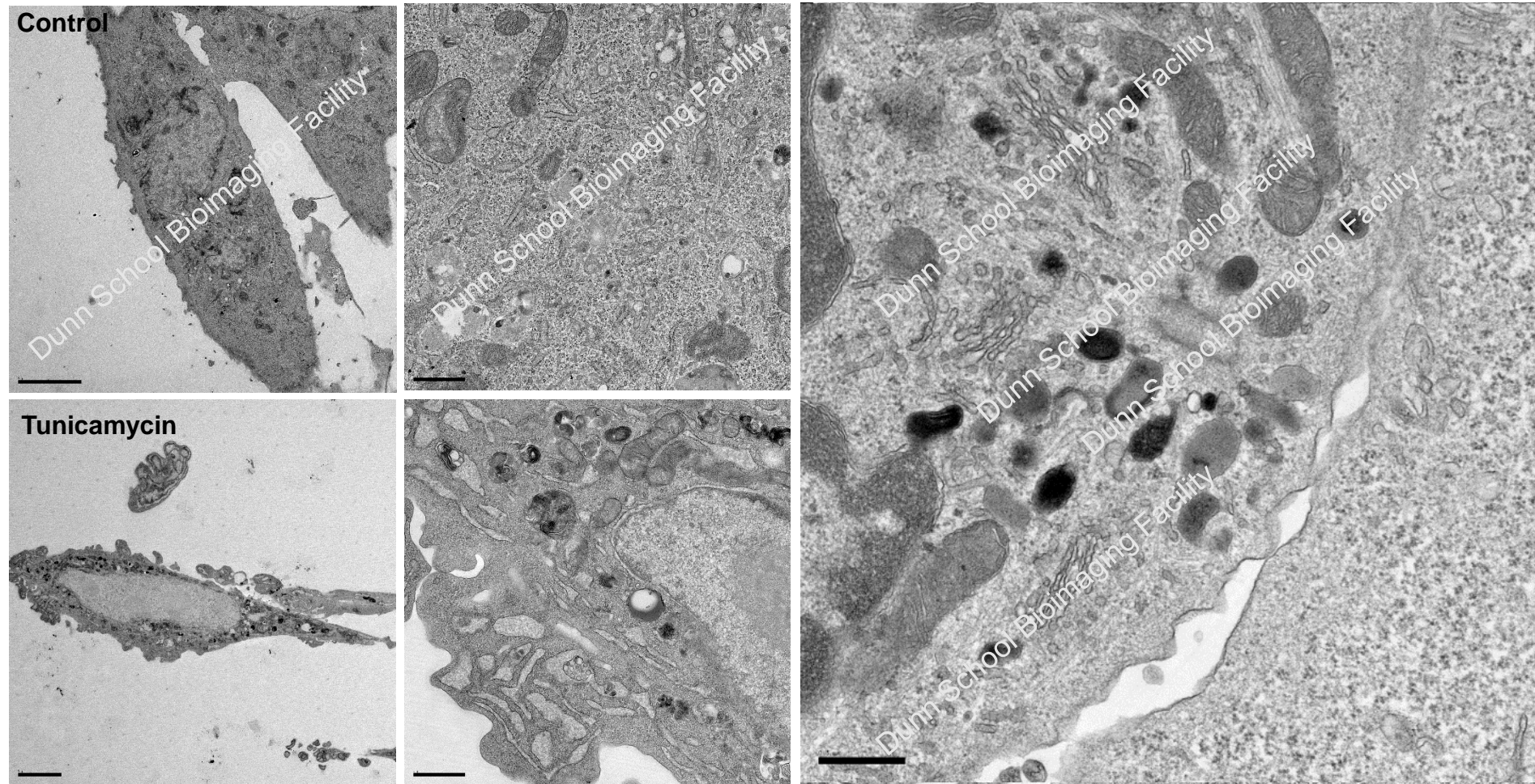
Cells



Mammalian culture cells (E Johnson)

TEM Ultrastructure

Cells

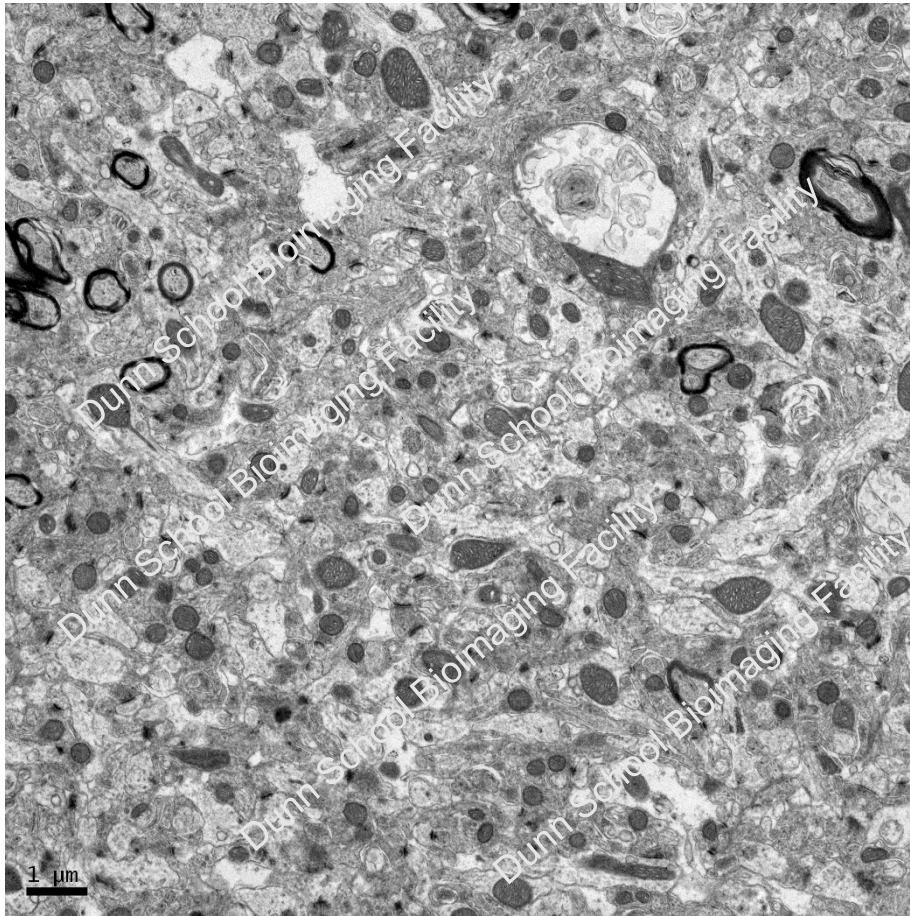


Mouse fibroblasts controls (top) and treated with tunicamycin (bottom)
E Johnson/V Liebe

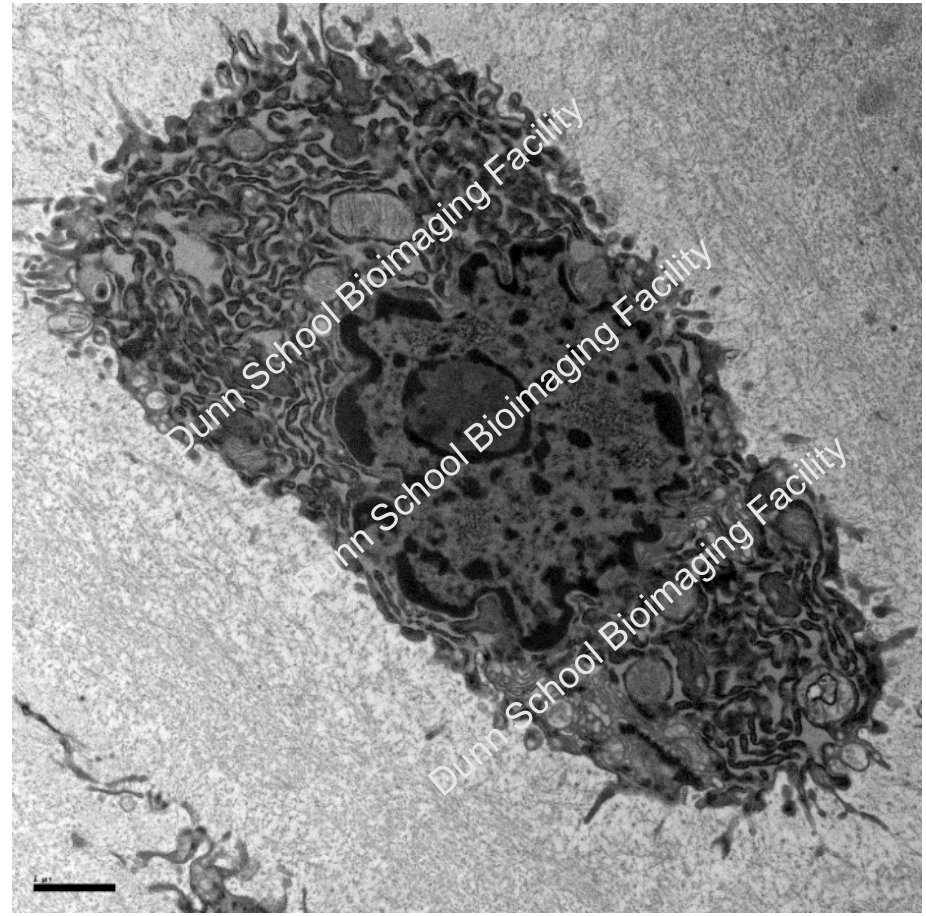
HRP labelled T-cell interacting with a melanoma cell
E Johnson/G Bossi

TEM Ultrastructure

Tissue



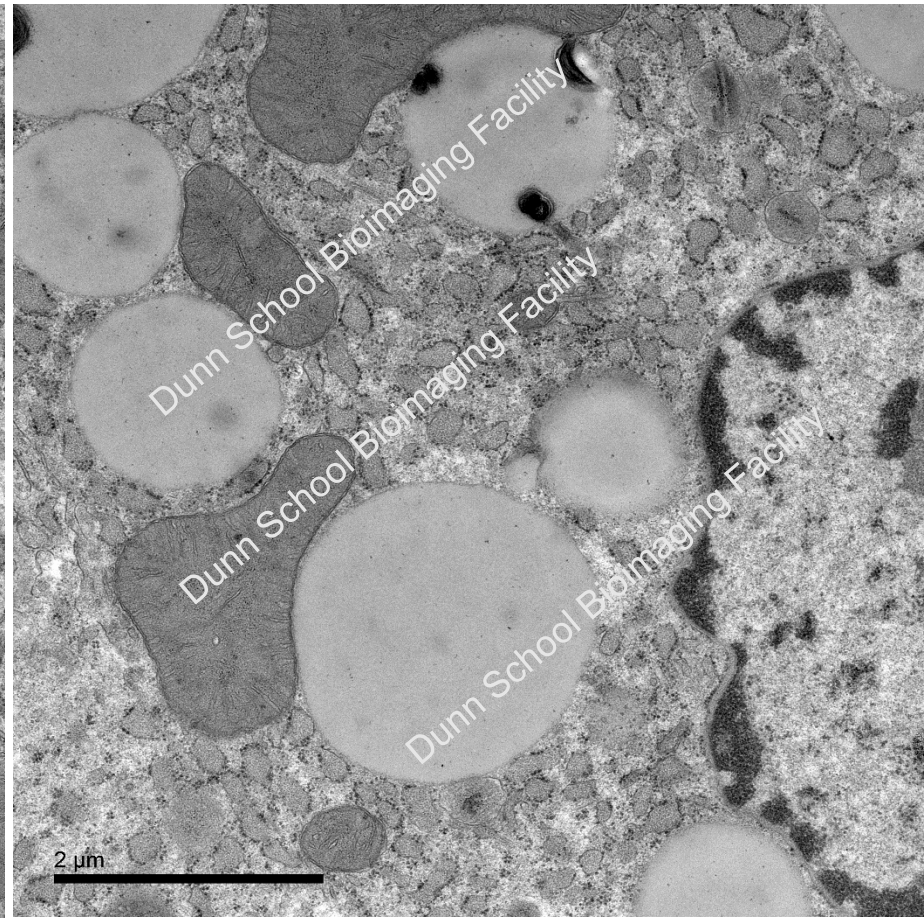
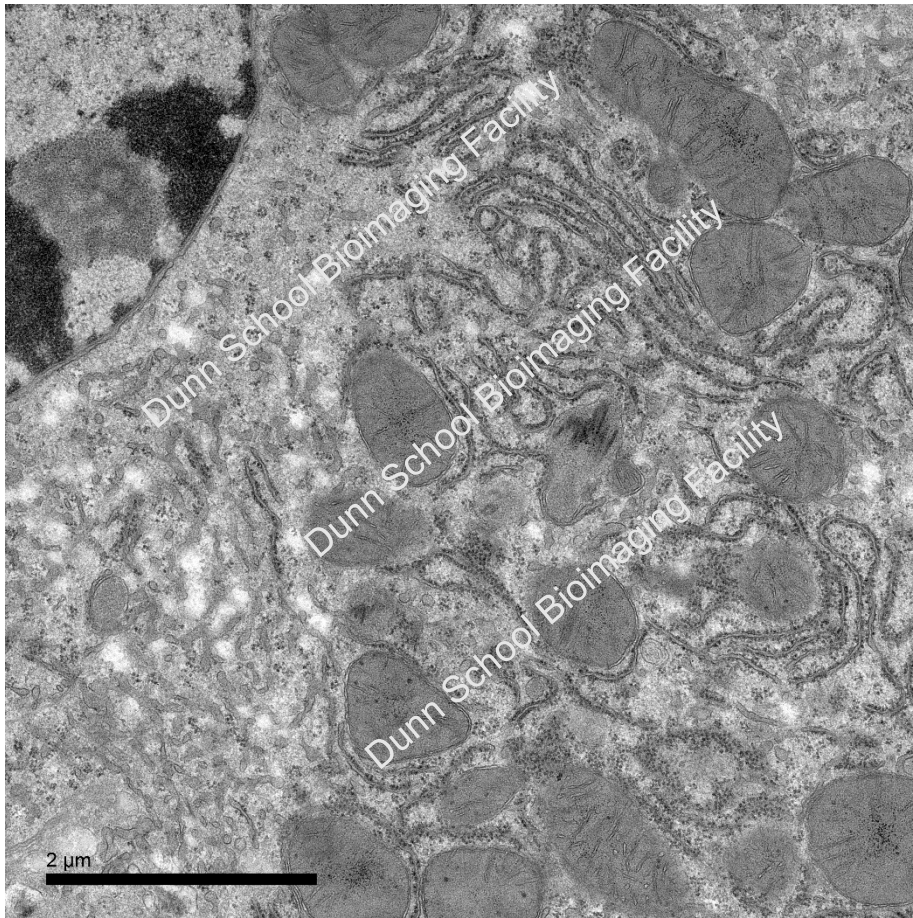
Mouse brain tissue
Tecnai12 TEM, E Johnson



Chondrocyte in mouse cartilage tissue
Tecnai12 TEM, P Sacitharan/A Pielach

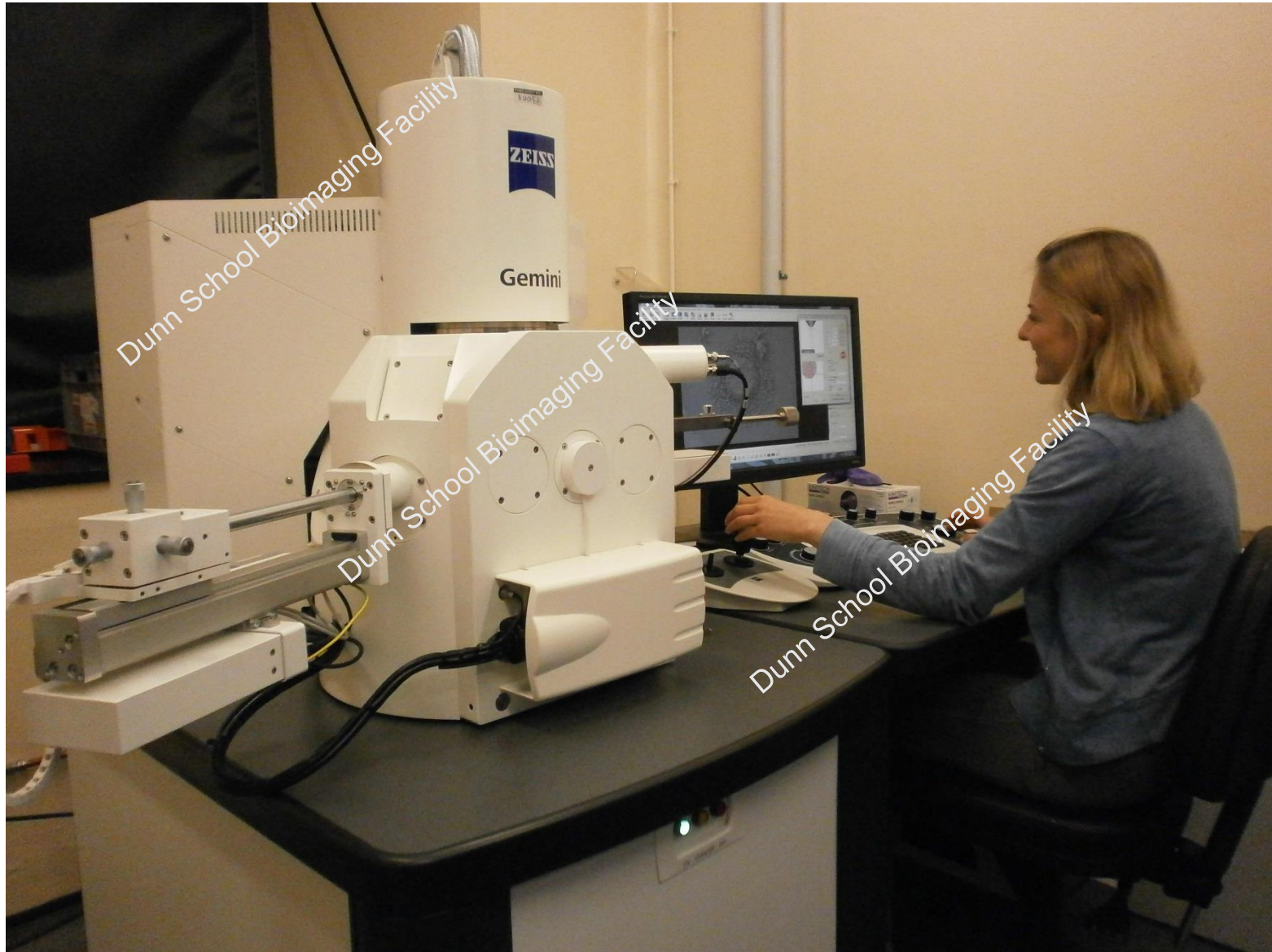
TEM Ultrastructure

Tissue



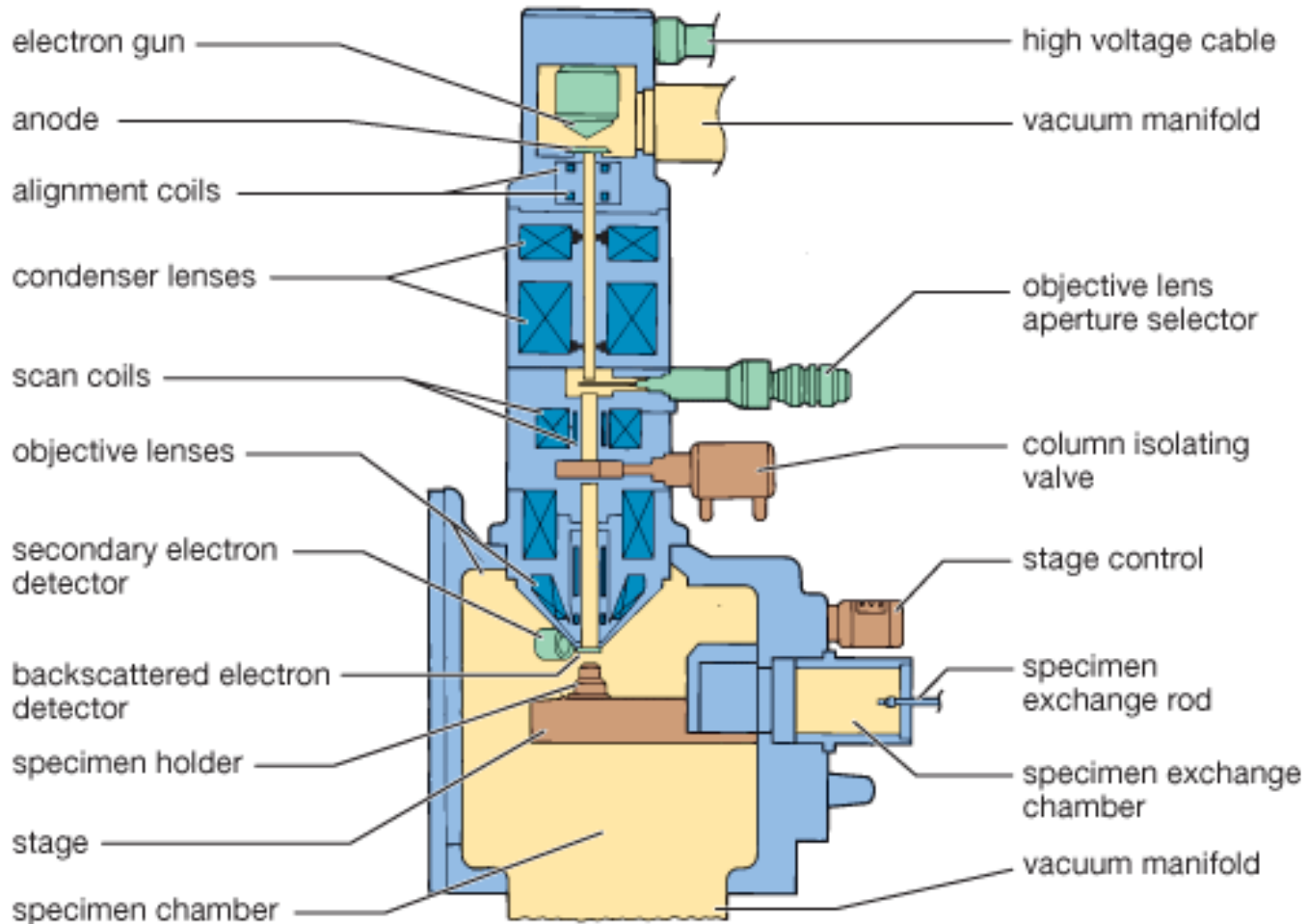
Mouse liver tissue, untreated (left) and under ER stress (right) (V Liebe & E Johnson)

Scanning Electron Microscopy (SEM)



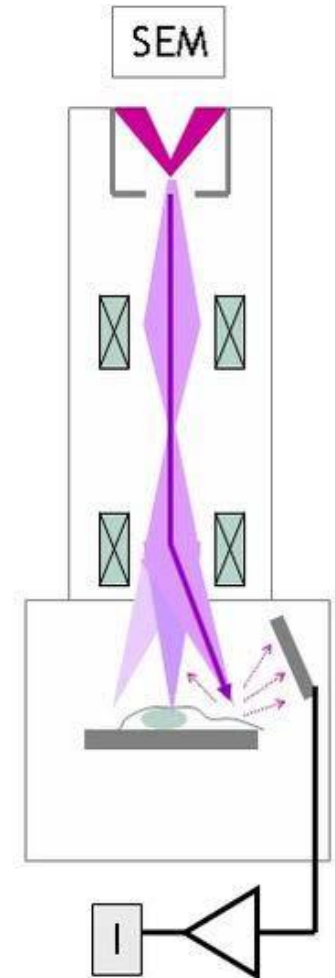
The SEM

Parts of a scanning electron microscope



Source: JEOL U.S.A., Inc.

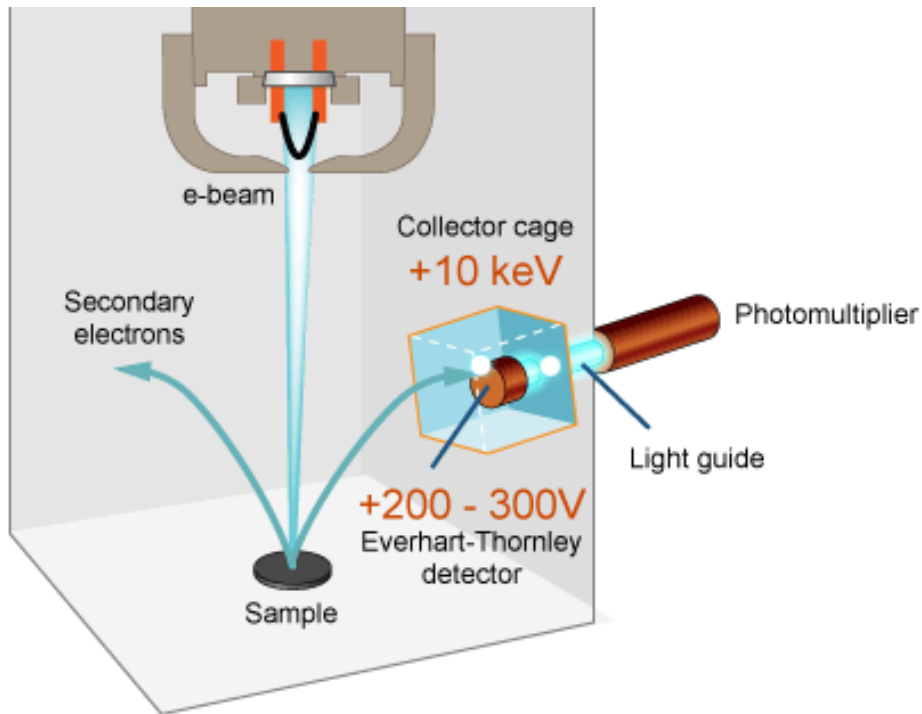
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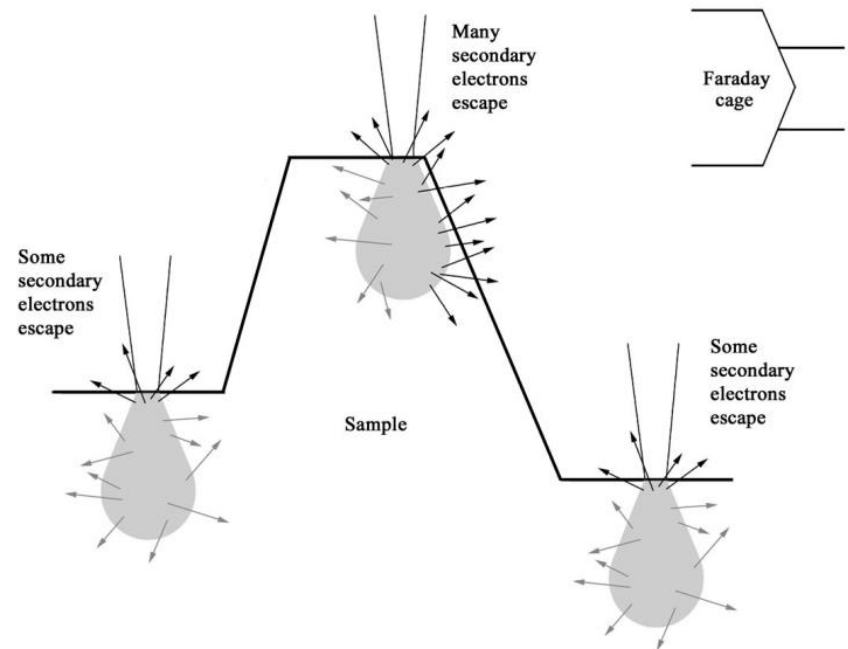
The SEM

Signal detection

- Secondary electrons (SEs) provides surface morphology and topology information.
- SEs are captured by the Everhart-Thornley detector



www.ammrf.org



Dept Biological Sciences, Smith College Northampton USA

SEM

Specimen requirements

SEM

Stable in the vacuum

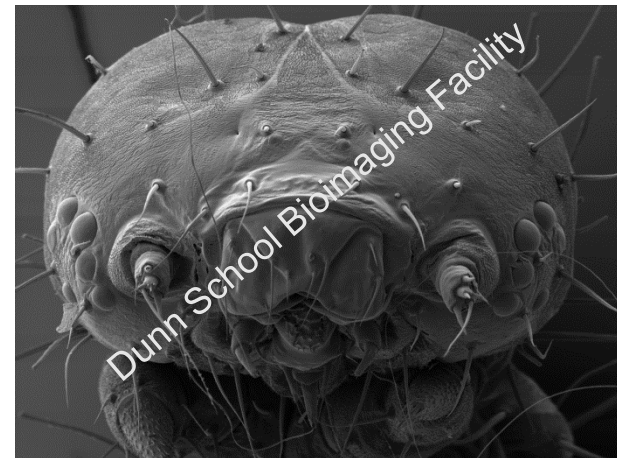
Well preserved surface structure

Conductive surface

Whole mount

Particulate samples can be coated and viewed quickly

Cells and whole organisms require some spec prep



Specimen preparation for SEM

Cells and tissue

Primary fixation
with
glutaraldehyde



Secondary fixation
Osmium tetroxide



Dehydration series
Ethanol/acetone



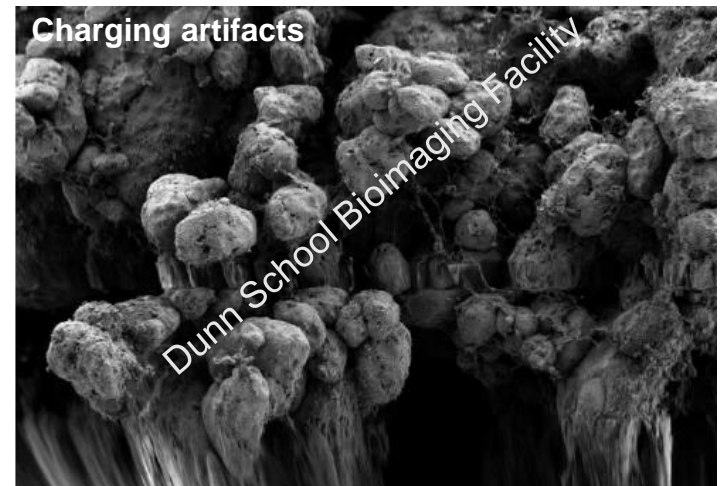
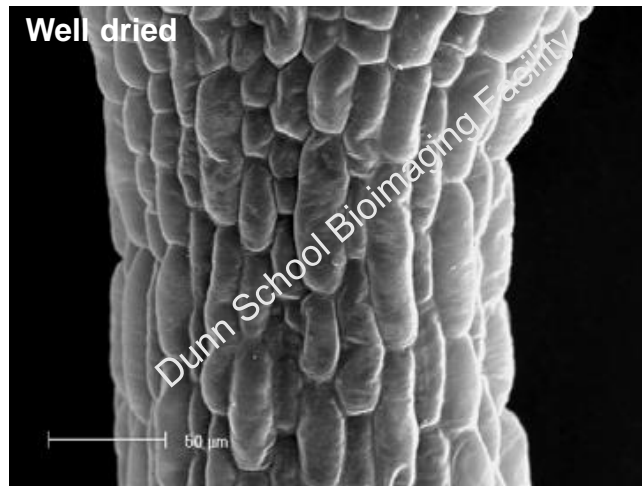
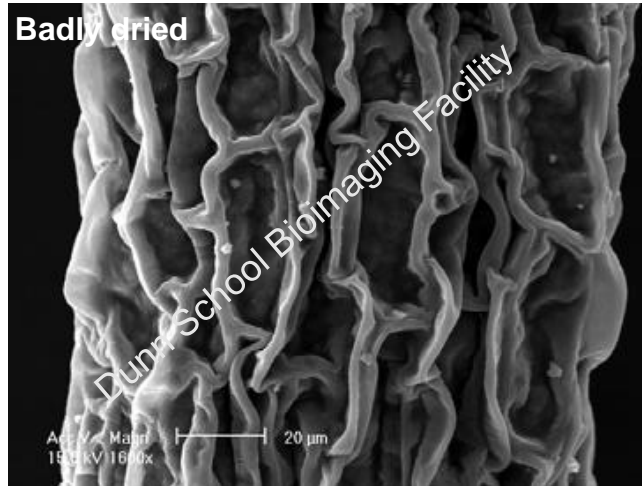
Dry using HMDS
or with the critical
point dryer



Mount & sputter
coat



SEM



Arabidopsis stem, Phillips XL30 SEM, E Johnson

*Arabidopsis xylem (top) & processed cheese bottom
Zeiss UltraSEM, E Johnson*

Electron microscopy techniques

Which one should I use?

- **Particles:**

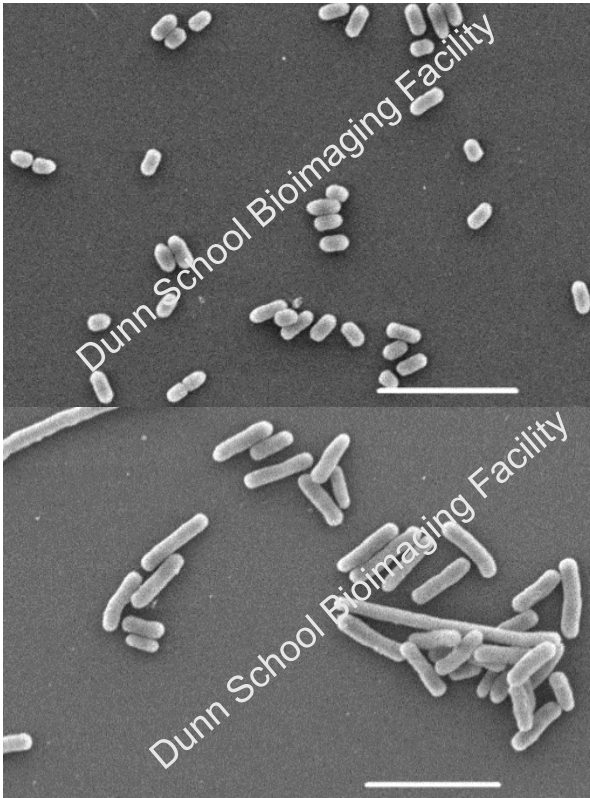
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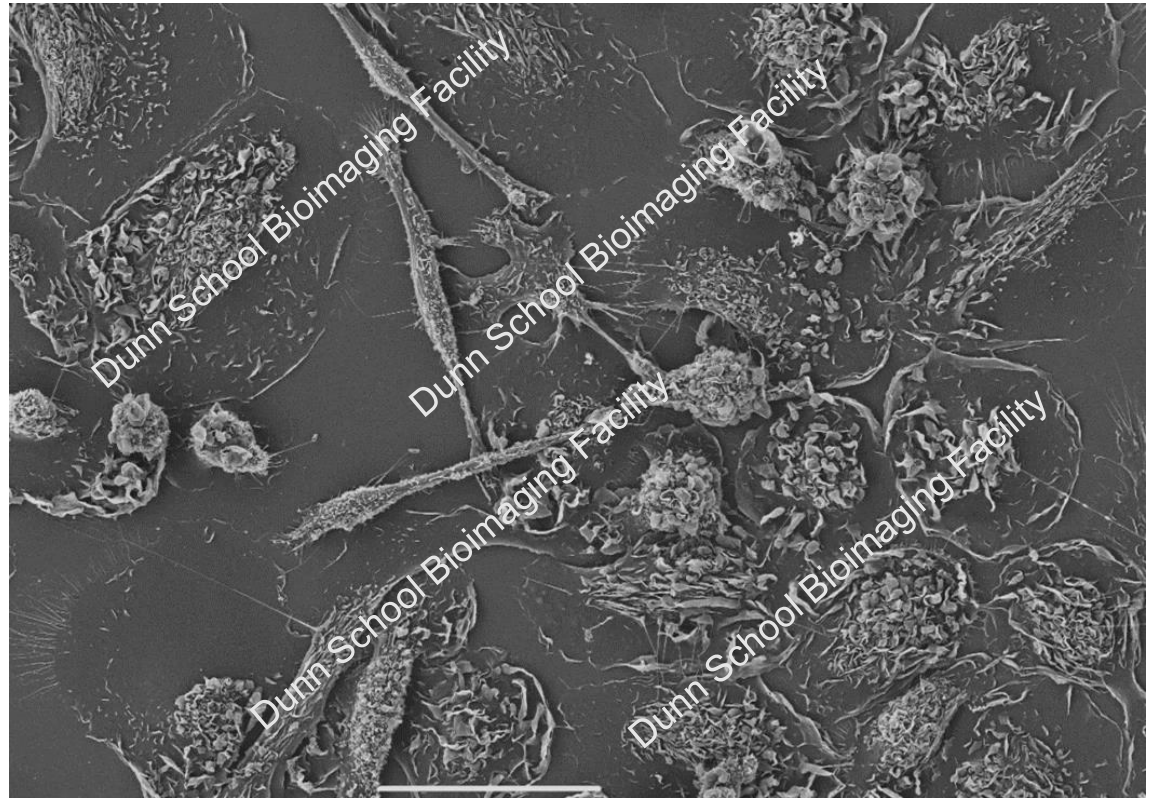
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SEM topography

Cells



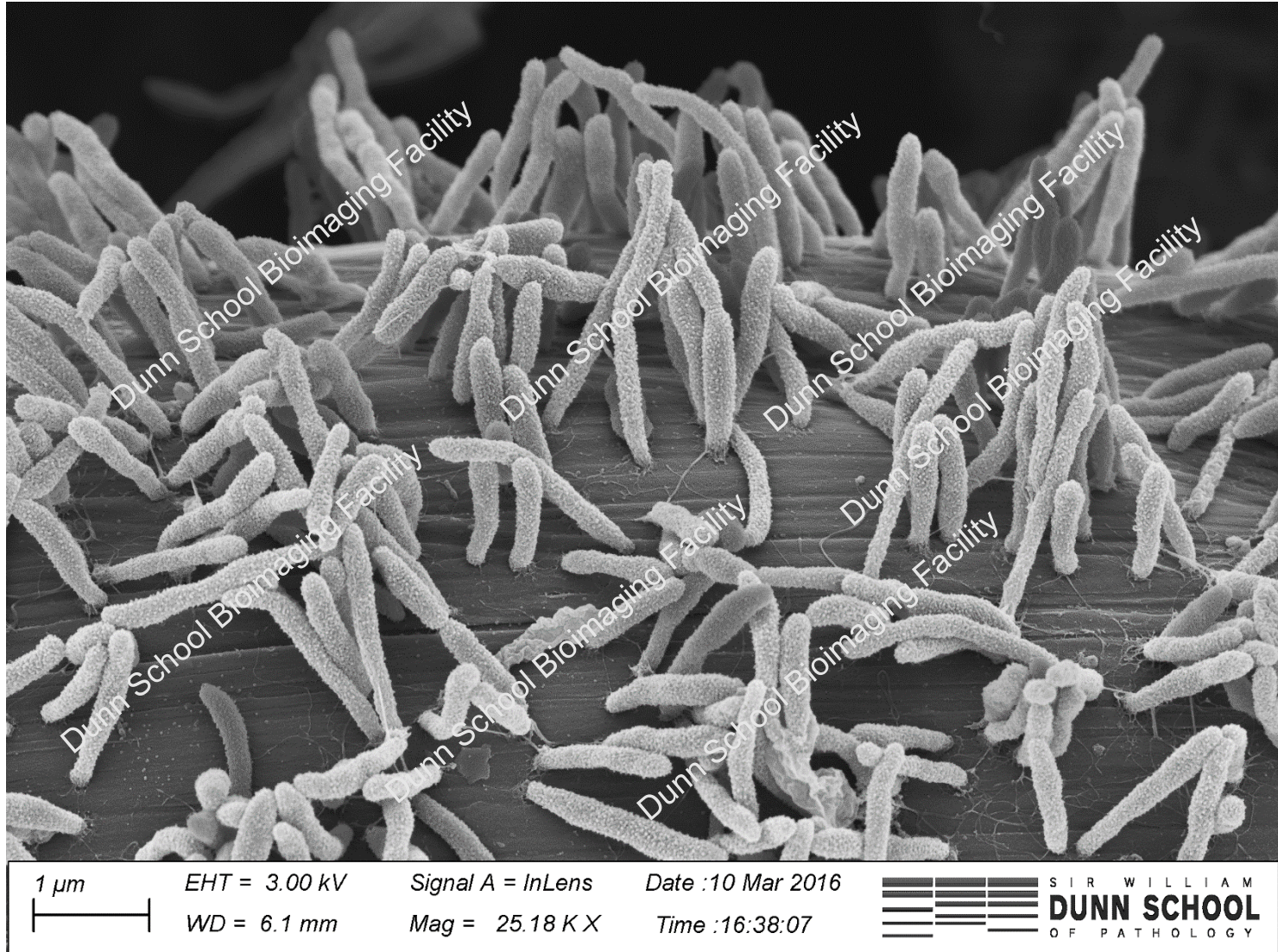
E. coli (WT at top, +vector at bottom)
Scale bar 5 μm (R Harding/E Johnson)



Monocytes and macrophages, scale bar 50 μm (B van Wilgenburg/E Johnson)

SEM topography

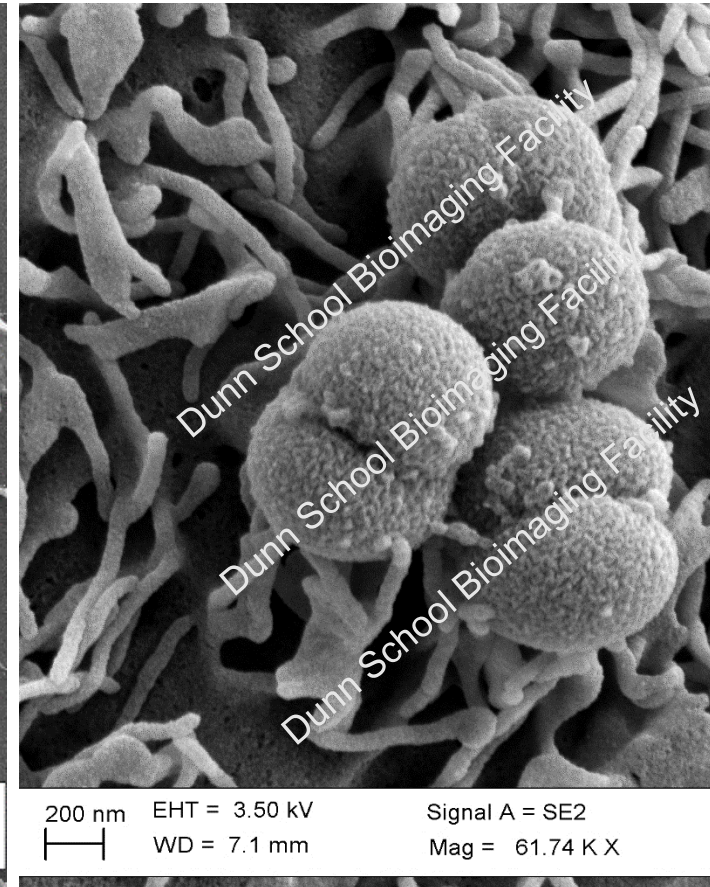
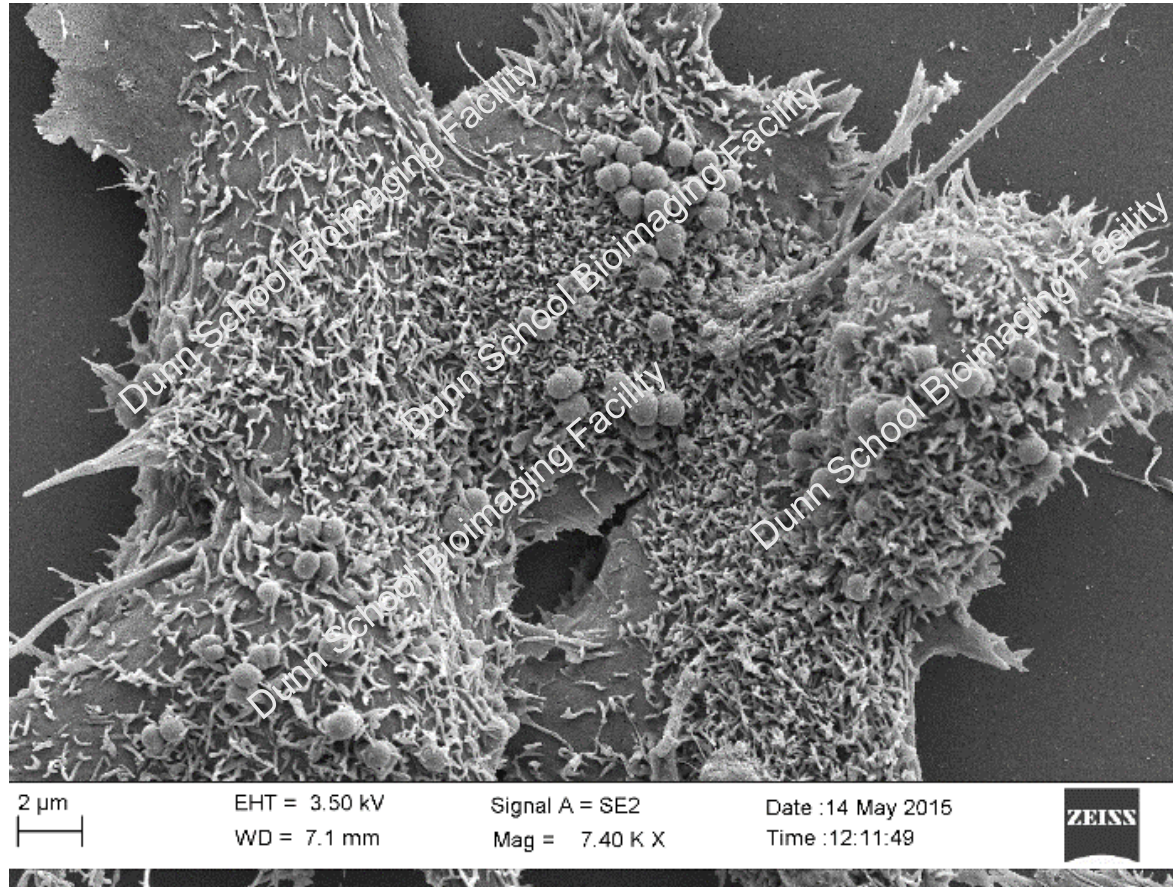
Cells



*Bacteria on carbon electrodes
(s Putra/E Johnson)*

SEM topography

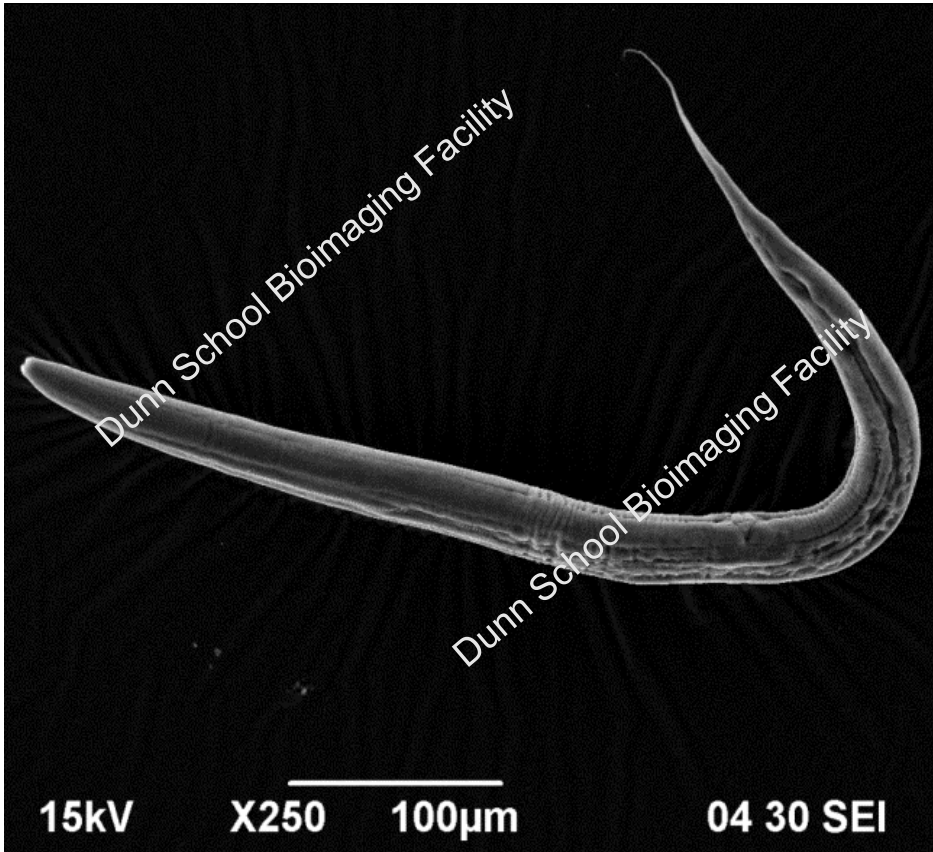
Cells



Neisseria sp. on epithelial cells (R Exley/EJohnson)

SEM topography

Organisms



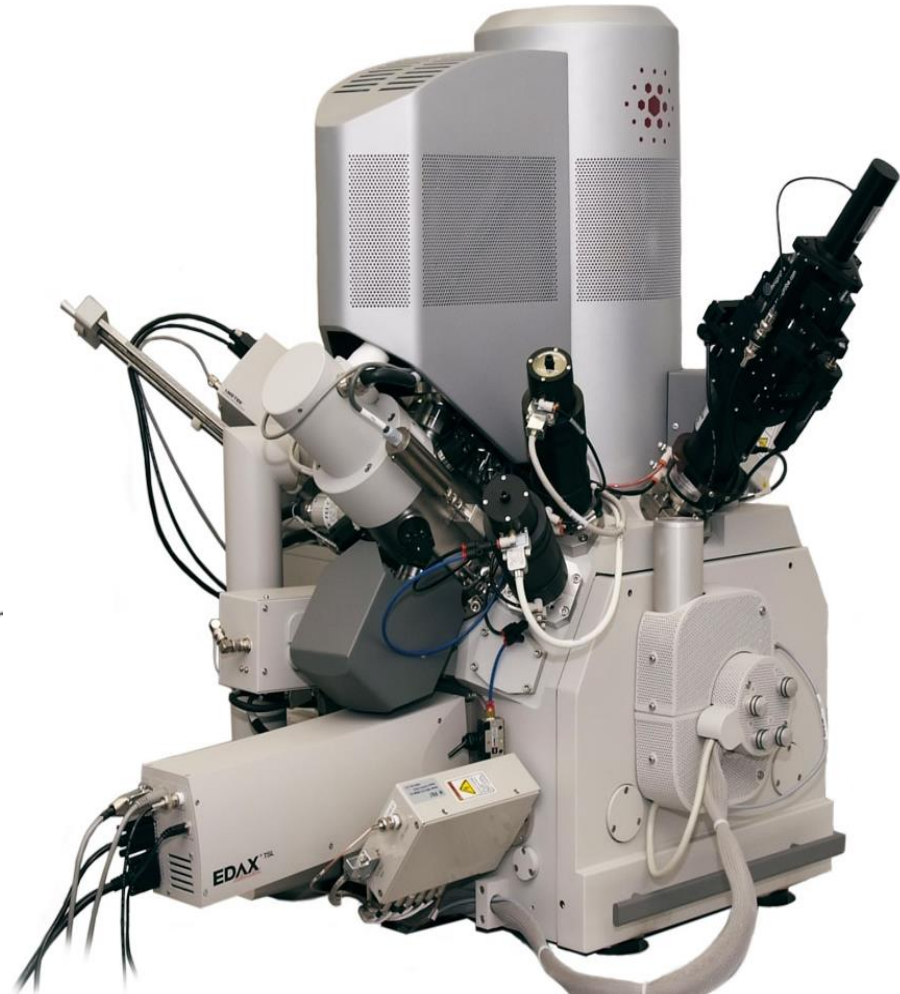
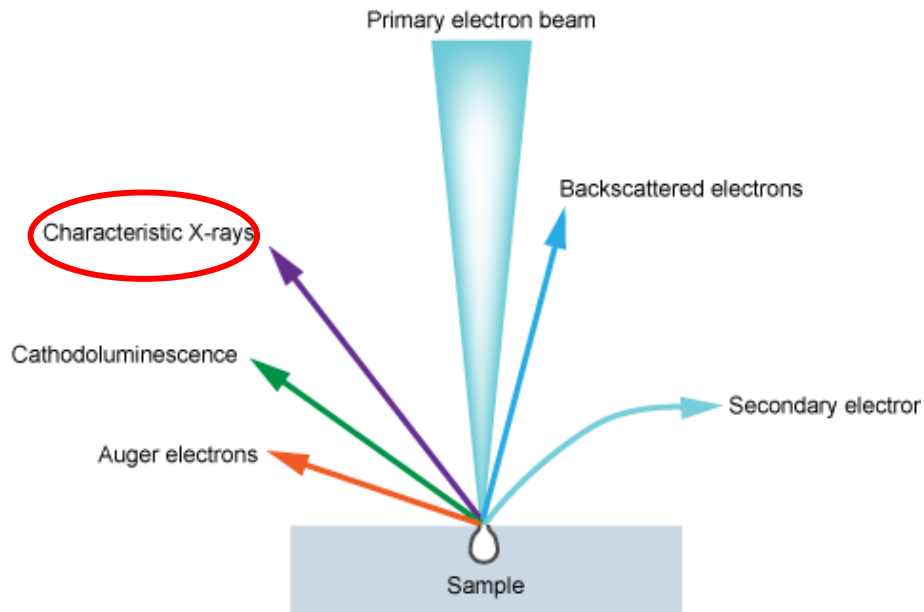
C. elegans
(E Johnson/A Moloney, Dunn School)



Drosophila rough eye phenotype
(M Elschami, NDCN)

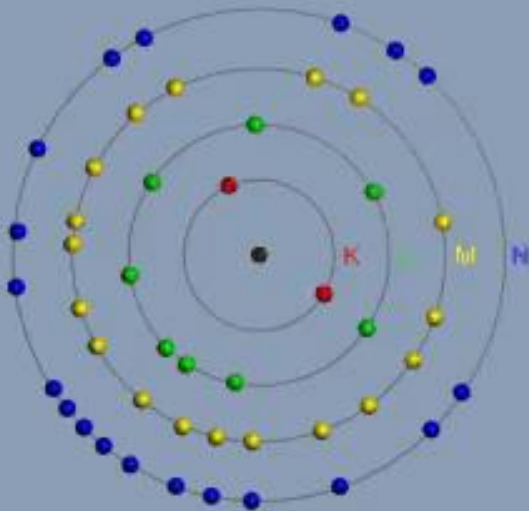
The SEM

Diverse imaging capabilities



Elemental mapping in the SEM

Energy Dispersive X-ray Spectroscopy (EDS)



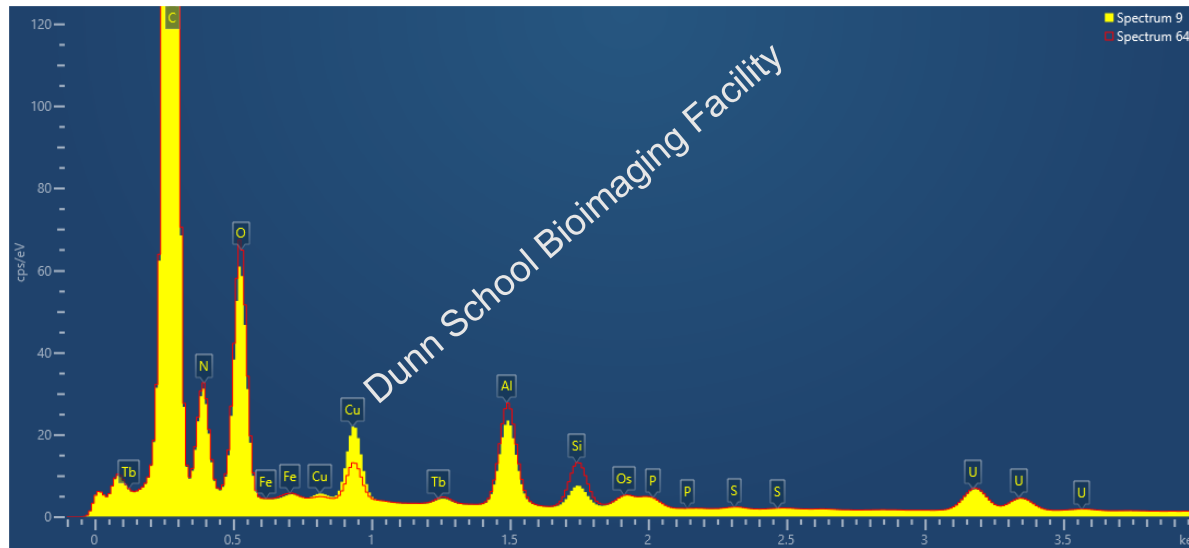
1. Interaction between a high energy beam electron and an atom on the sample's surface
2. Inner orbital electron ejected: ionisation
3. Vacancy filled by outer orbital electron
4. Release of energy as X-ray photon
5. An EDS detector is then used to measure the energy of the X-ray photon which is specific to the atom and therefore identify it

This enables the elements in the sample to be mapped and quantified

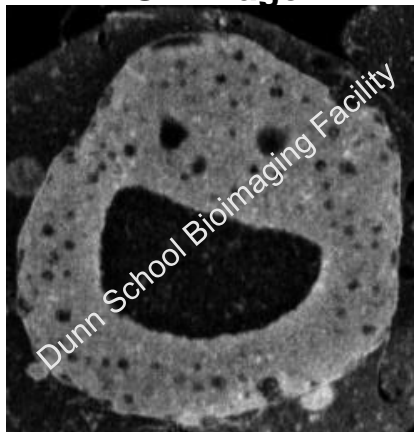
Slide modified from P Trimby, Oxford Instruments

Elemental mapping in the SEM

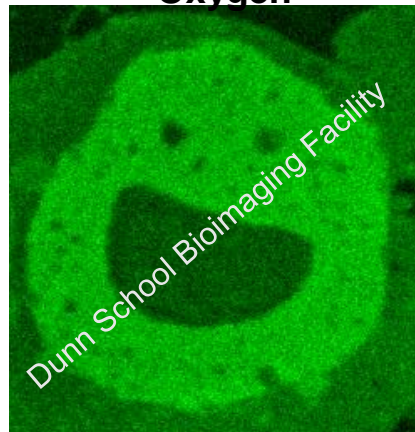
Energy Dispersive X-ray Spectroscopy (EDS)



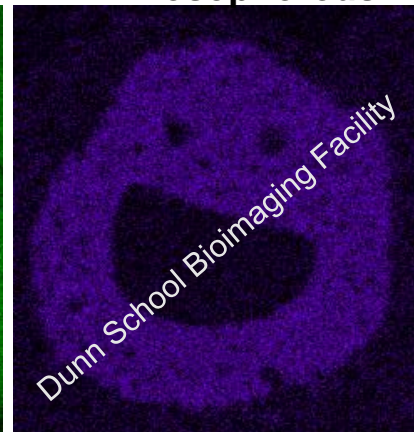
SE image



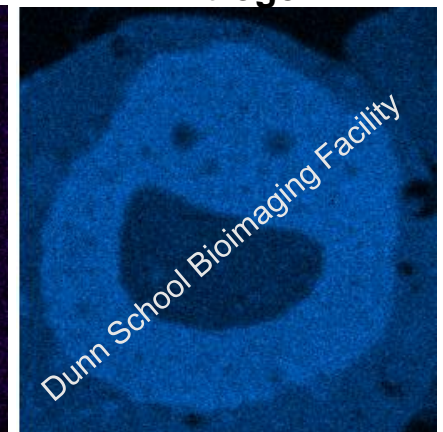
Oxygen



Phosphorous

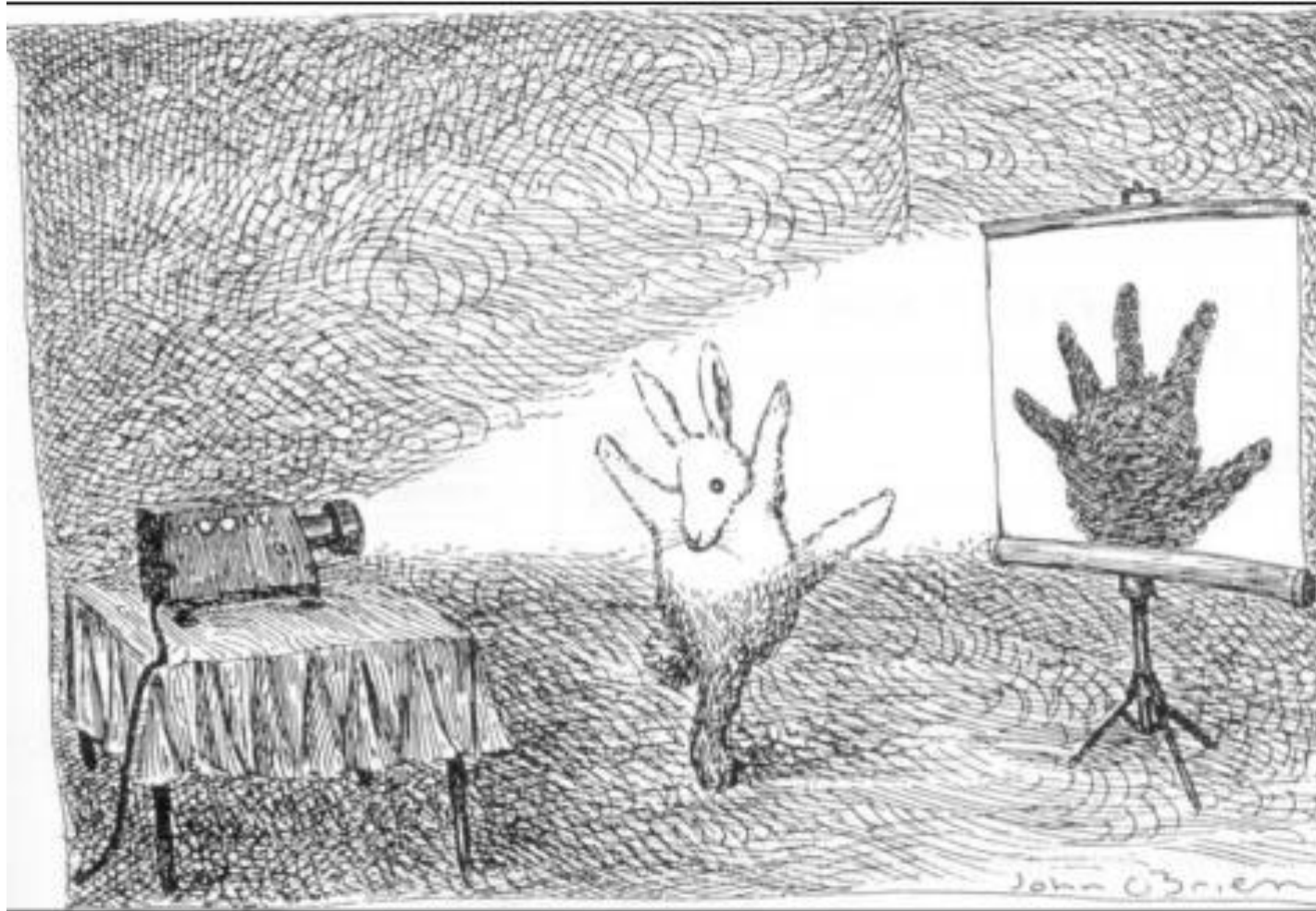


Nitrogen



Advanced EM techniques

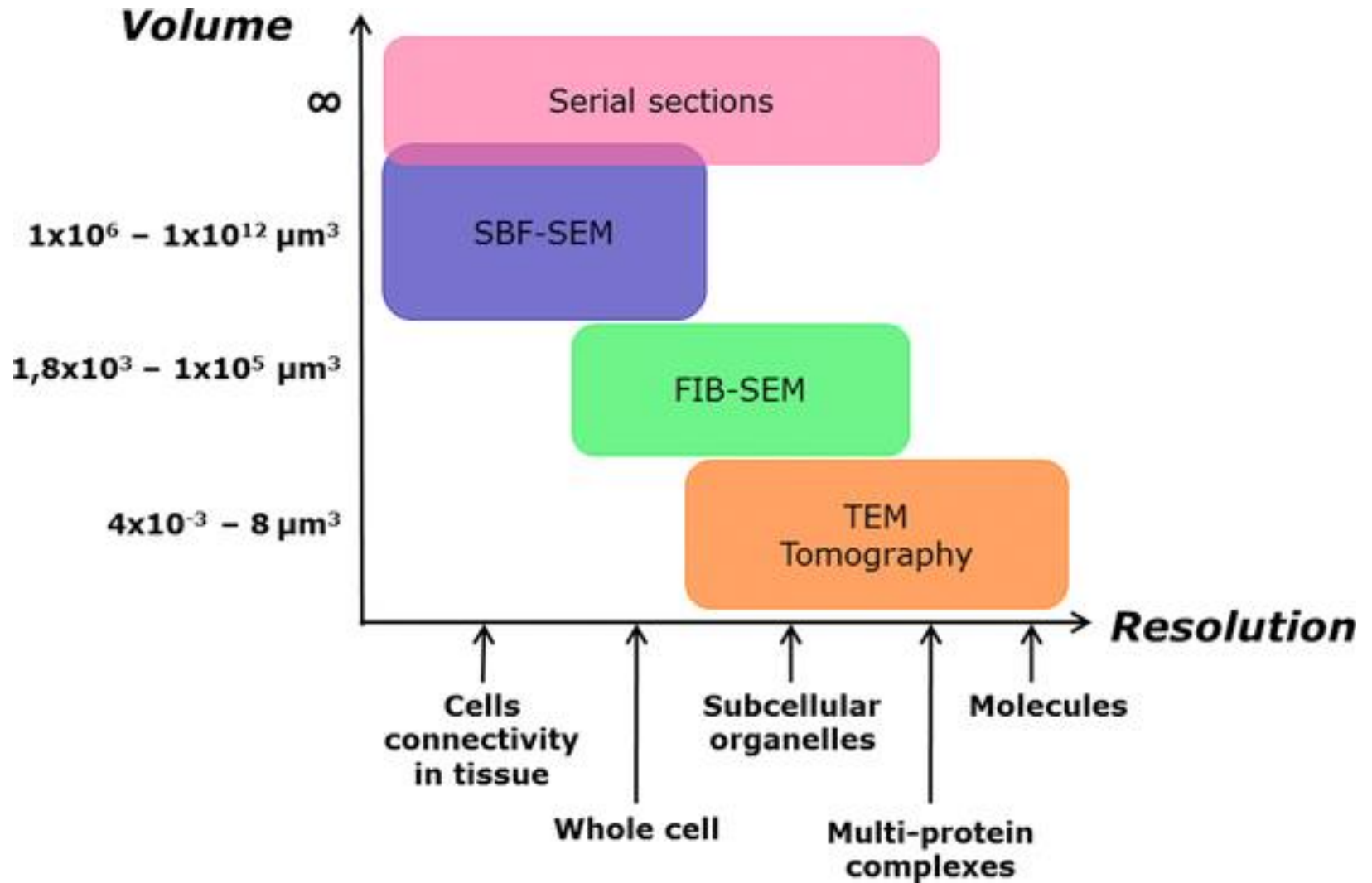
Volume electron microscopy



Drawing by John O'Brien, The New Yorker Magazine (1991)

3D/volume EM techniques

Overview



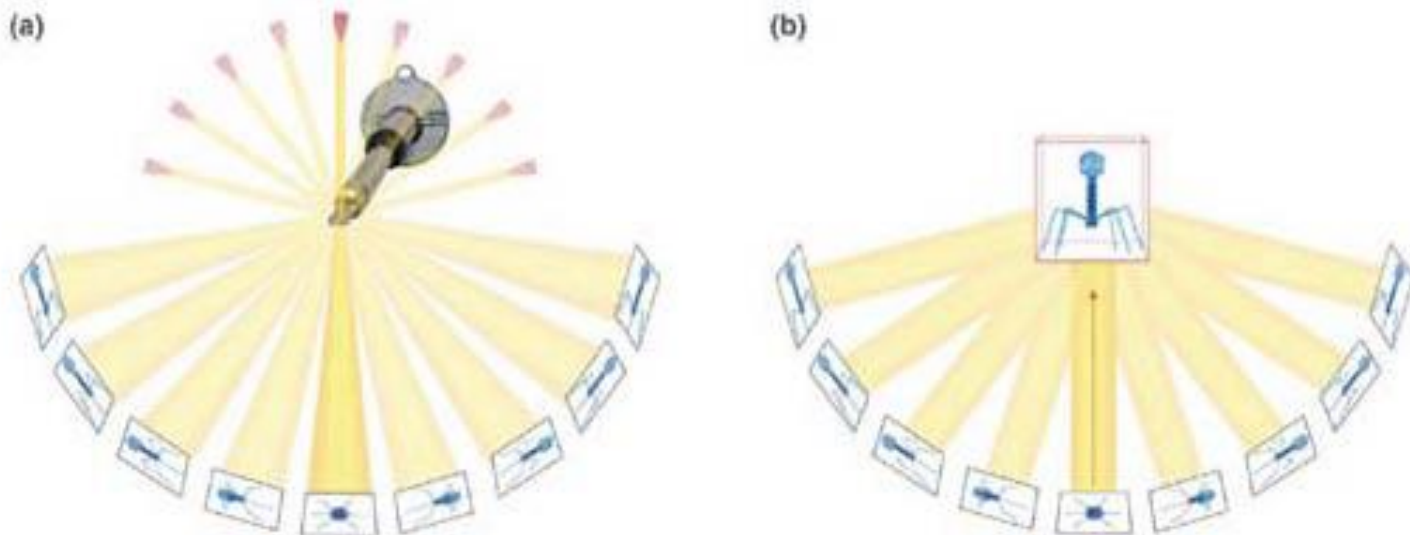
From: Kizilyaprak et al (2014) *J Microscopy*, 254(3).

3D EM Techniques

TEM - Electron tomography

- Thicker sections (150-300 nm) on filmed slot grids with gold fiducial markers
- Use specialised tomography holder for dual axis tilting of the specimen
- Reconstruct using modelling software

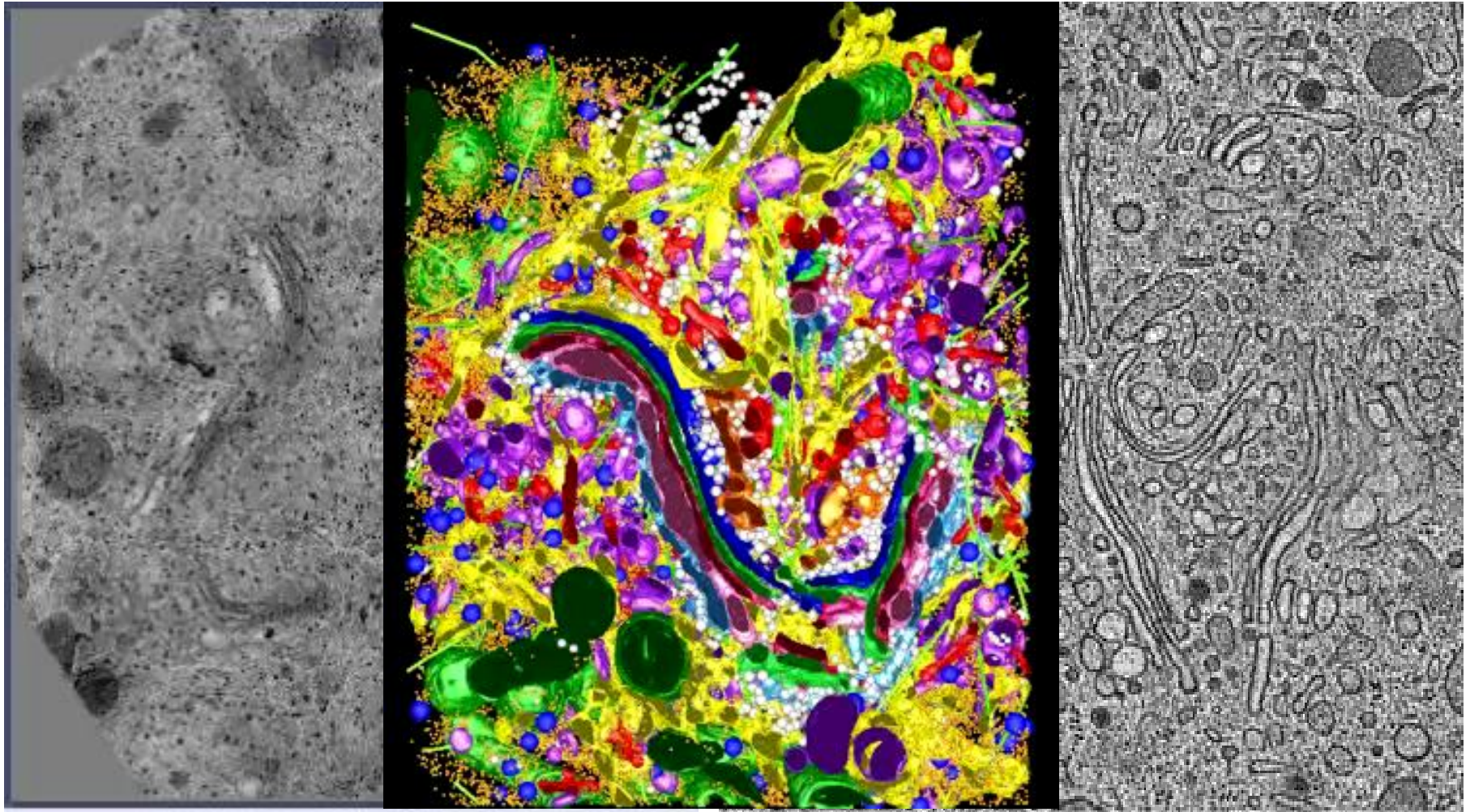
Z resolution: ~2 nm



Principles of Electron Tomography. (a) A biological specimen, in this case a bacteriophage contained in an EM sample holder, can be imaged from several orientations by tilting the holder in the electron microscope. (b) Process of computed backprojection, in which each tilted view is used to reconstruct to three-dimensional information of the original structure. [McIntosh, et al. (2005) Trends Cell Biol. 15:43-51].

3D EM Techniques

TEM - Electron tomography



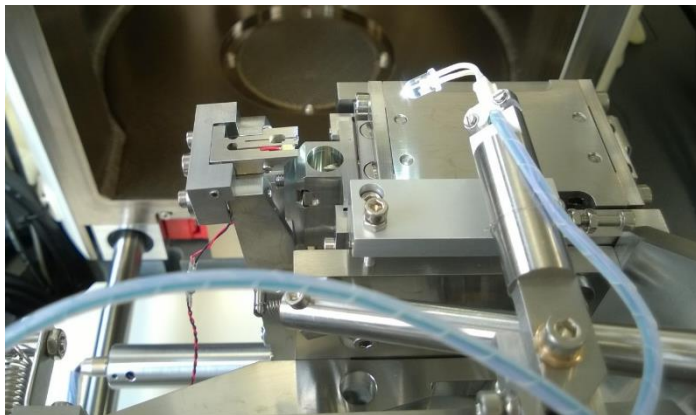
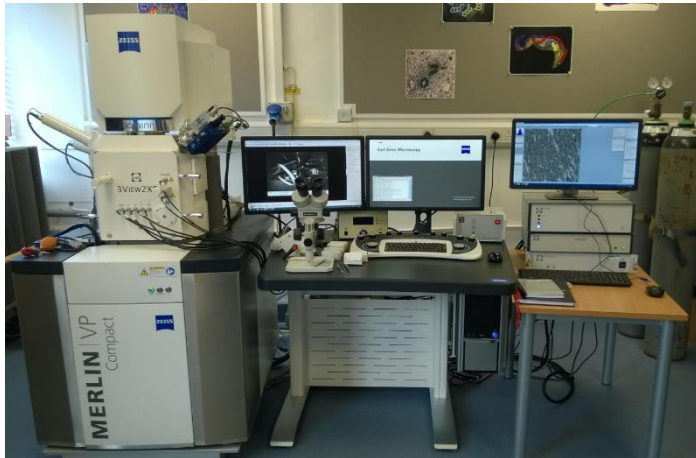
3D ultrastructure in the Golgi region of a pancreatic beta cell line.
Volume: $\sim 3.1 \times 3.2 \times 1.2 \mu\text{m}$, Marsh et al (2001) PNAS, 98.

3D EM Techniques

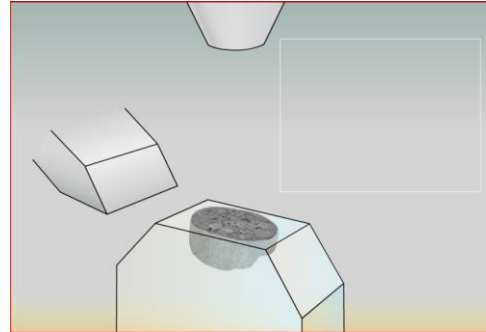
SEM - Serial Block Face Sectioning with Gatan 3View

One method for generating a 3D high resolution image stack is to use serial block face sectioning with the Gatan 3View system

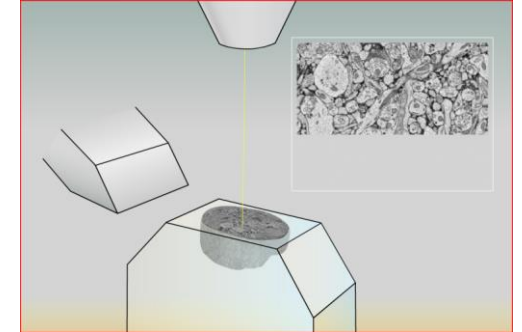
Z resolution: 30-200 nm



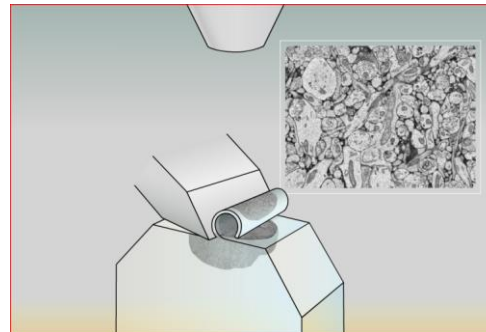
1. Start



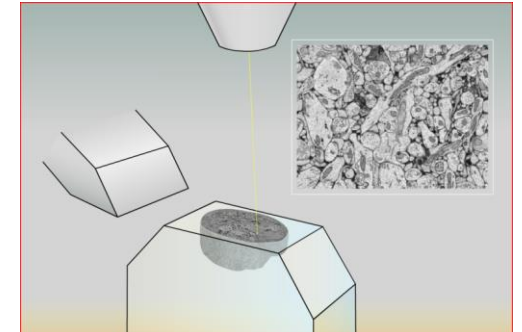
2. Block face scanned



3. Block moves up 50 nm and the diamond cuts the surface



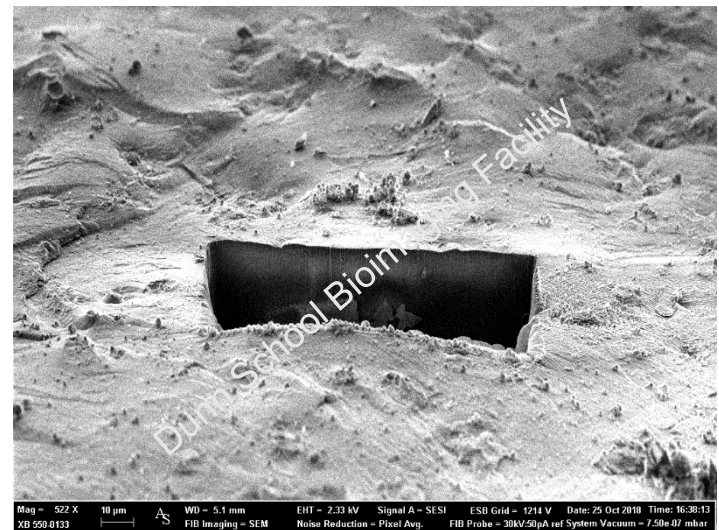
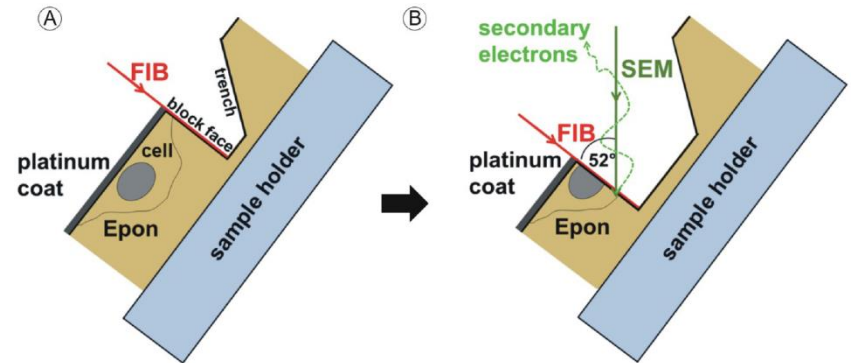
4. Newly revealed block-face is scanned



3D EM techniques

SEM - Serial Block Face Sectioning with FIB-SEM

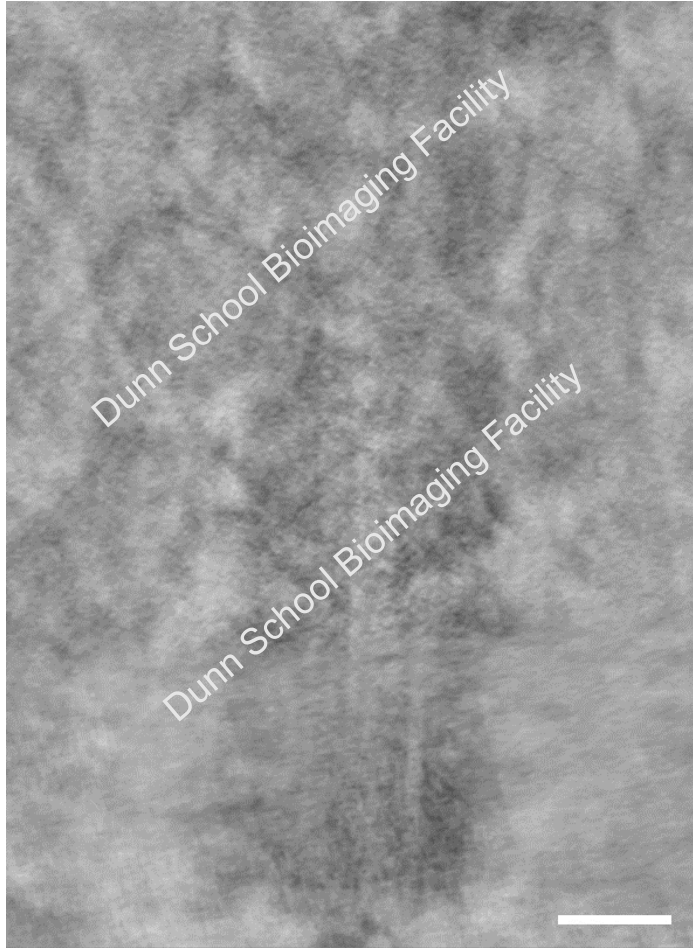
Z resolution: 5-100 nm



Cells and tissues

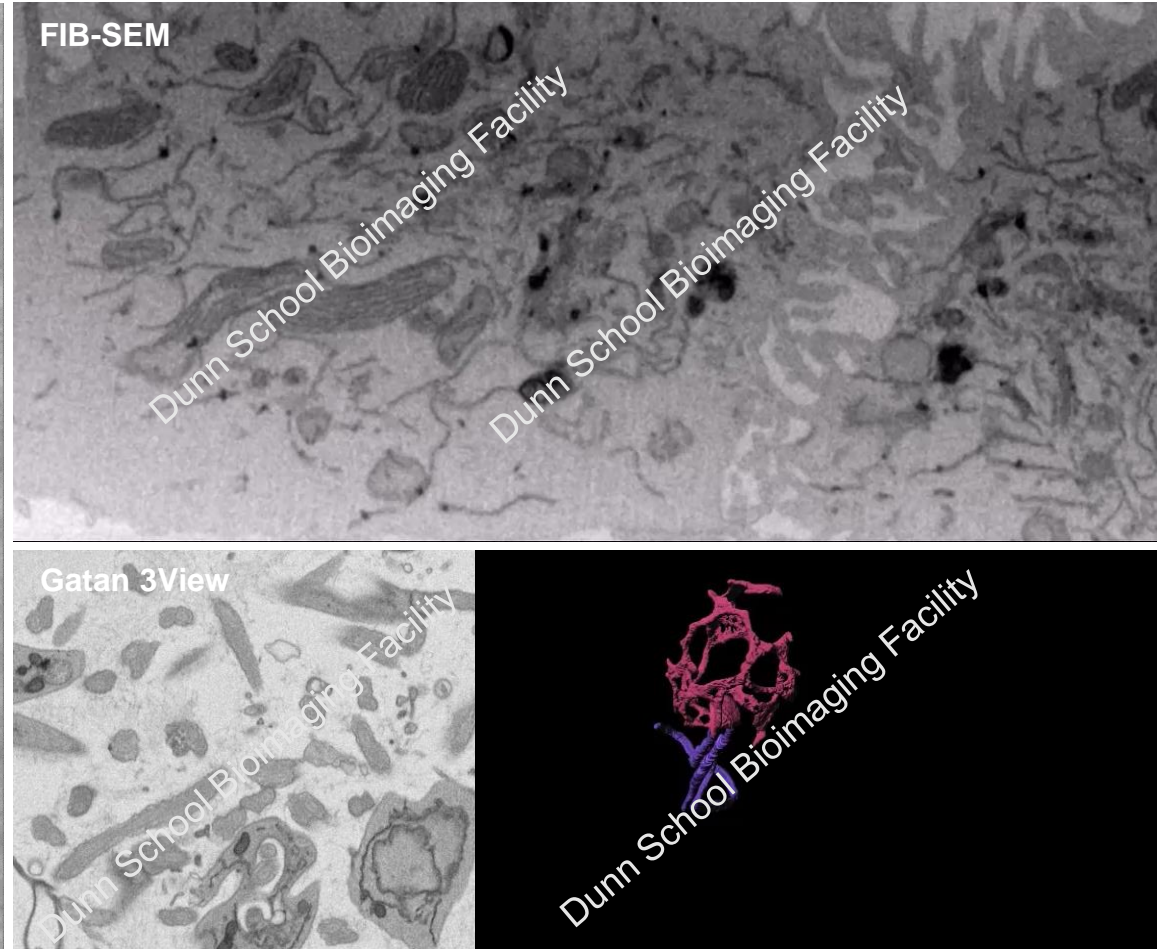
3D EM techniques

Electron tomography



Drosophila primary spermatocyte centrioles,
H Roque (Dunn School)

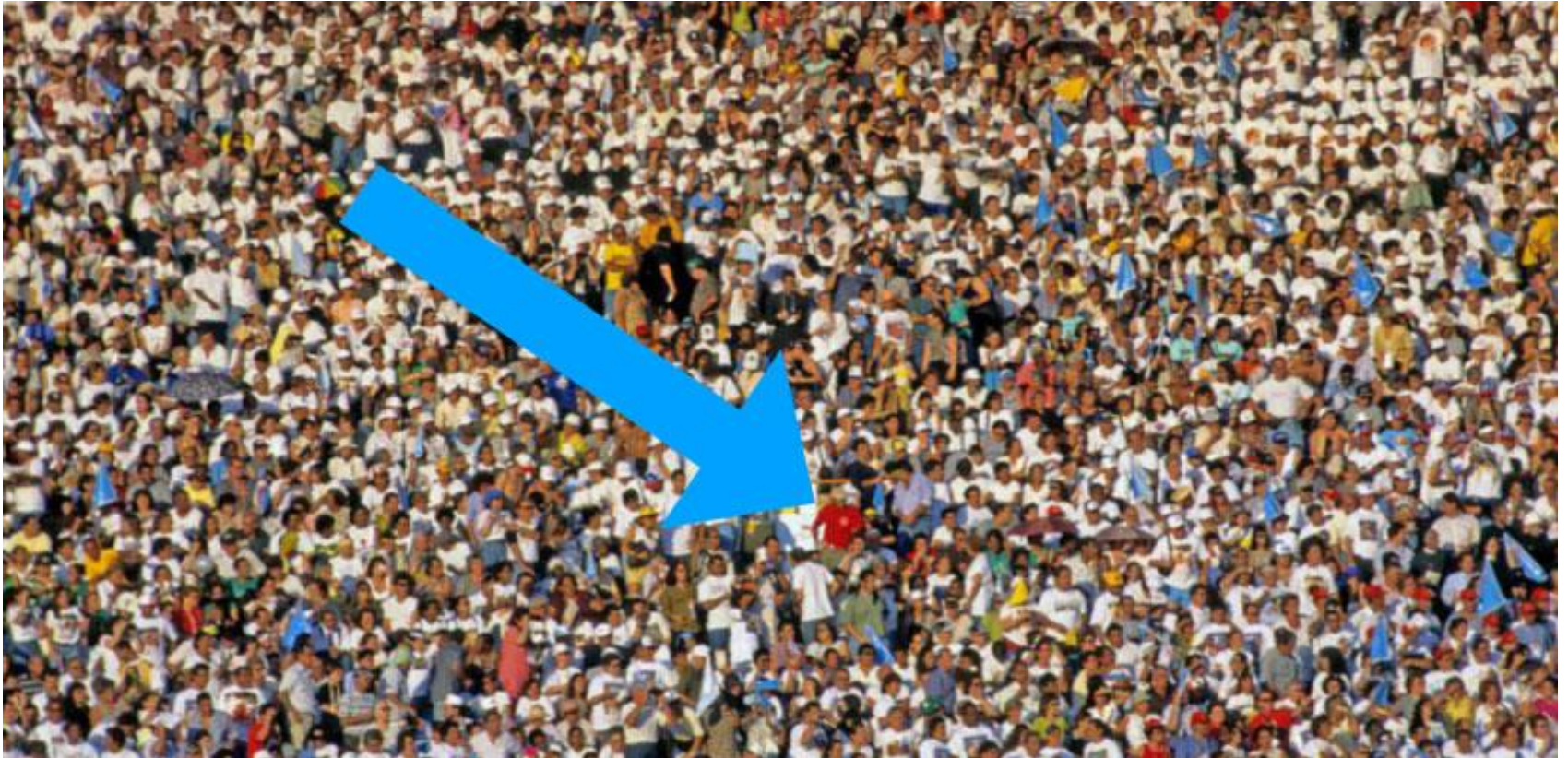
Serial Block Face Sectioning SEM



Top: FIB-SEM of epithelial cells infected with *N. cinerea*, voxel size = 5x5x15 nm³, Tang lab/E Johnson
Bottom: 3View of *L. mexicana*, volume: 9.8 μm x 12.2 μm x 16.1 μm, J Valli & E Johnson

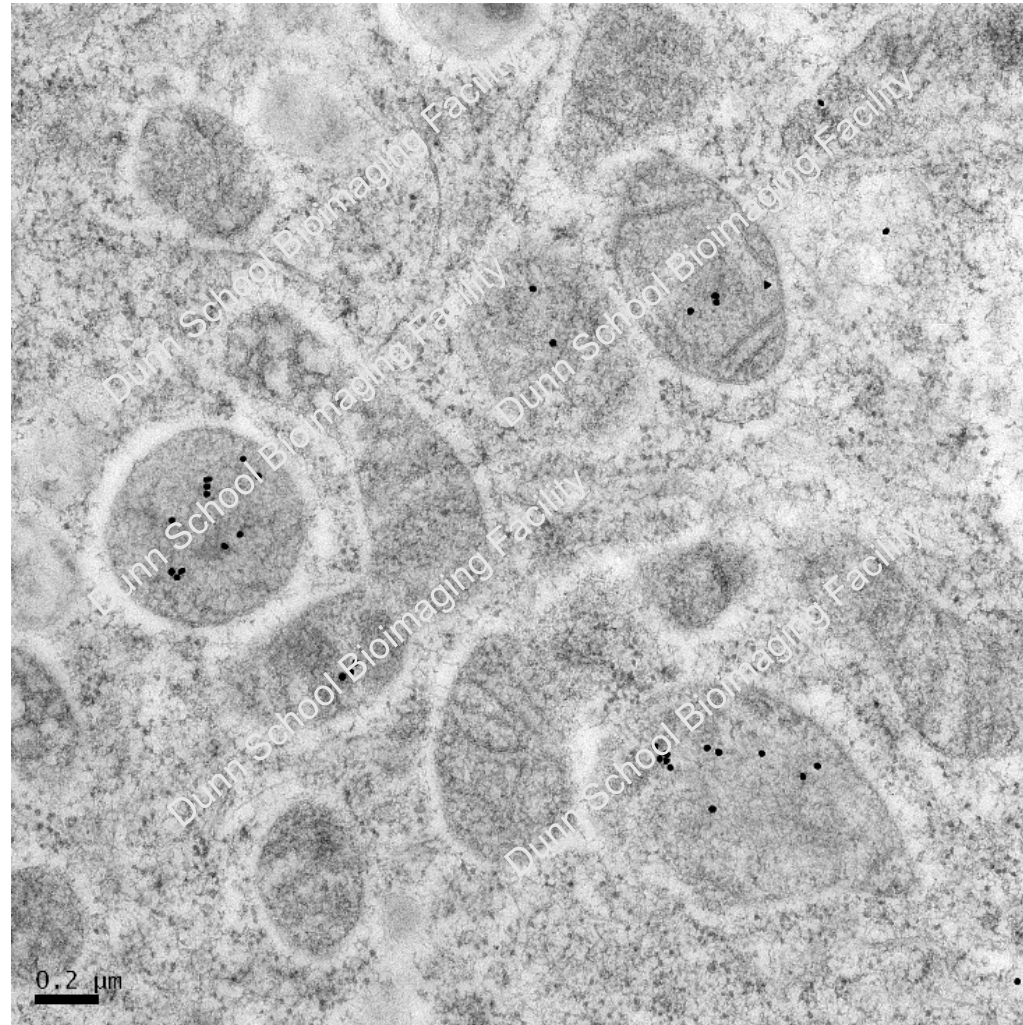
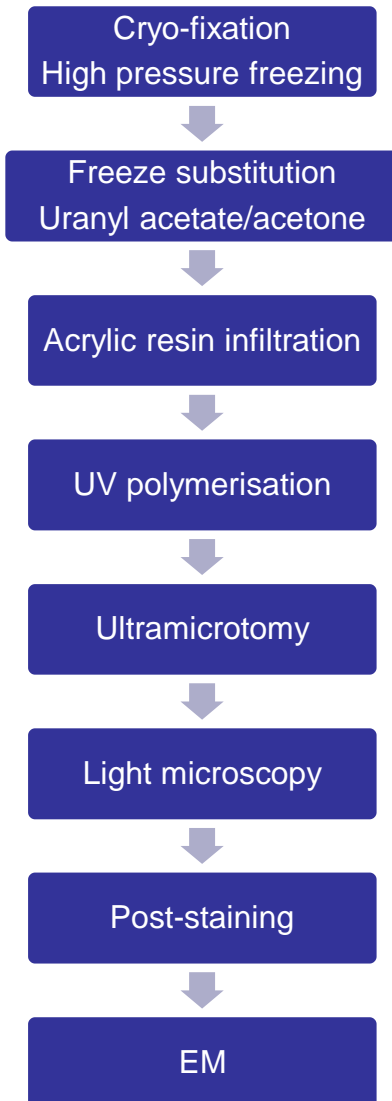
Advanced EM techniques

Protein localisation in cells and tissues



Protein localisation

Immunogold labelling – Cells & Tissues

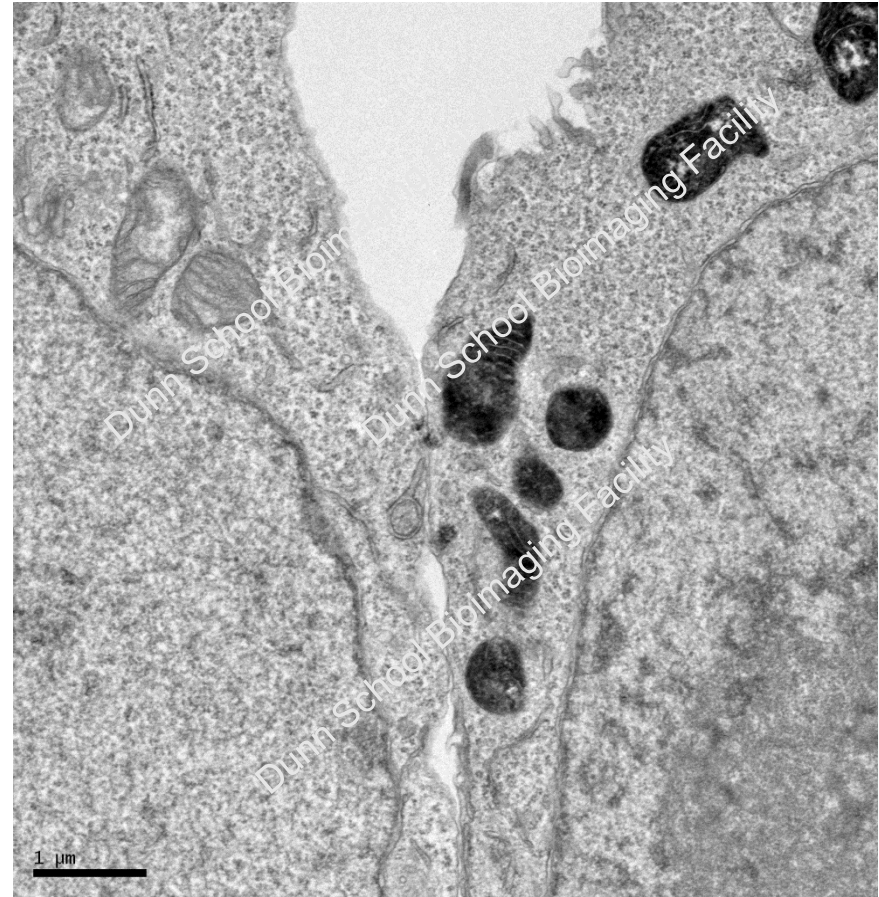


*Immunogold labelled mitochondria in a mammalian cell
(A Dhir/EJohnson)*

Protein localisation

EM genetic tags

- Several new genetically encoded tags are now available as alternatives to using immunogold labelling for identifying proteins of interest at the EM level whilst using a standard TEM prep
- APEX (Martell et al, Nature Biotech 30, 2012)
 - 28kDa peroxidase that catalyses with DAB (with H_2O_2) to produce a localised osmophilic precipitate
- miniSOG (Shu et al PLOS Biology 9, 2011)
 - Small fluorescent flavoprotein that can be photo-oxidised to react with DAB to produce a localised osmophilic precipitate - CLEM



Chemically fixed HEK cells transfected with APEX tagged to a mitochondrial matrix protein (J Long/E Johnson)

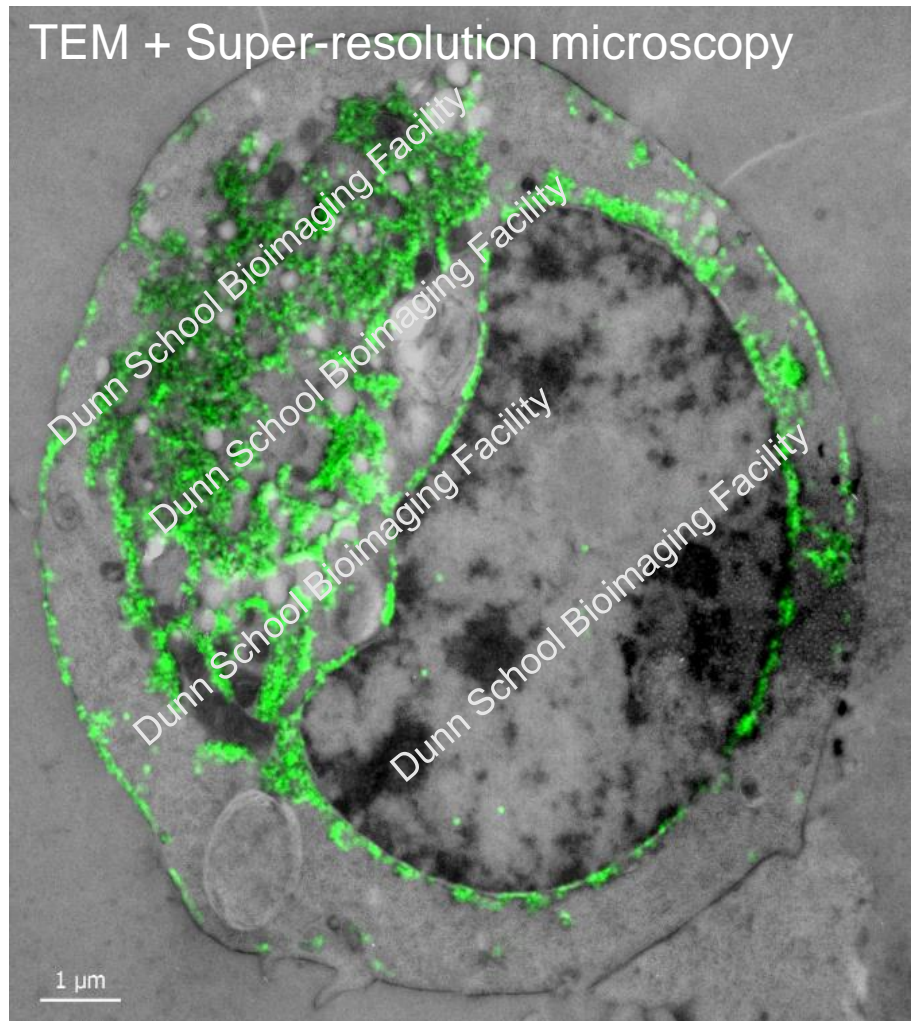
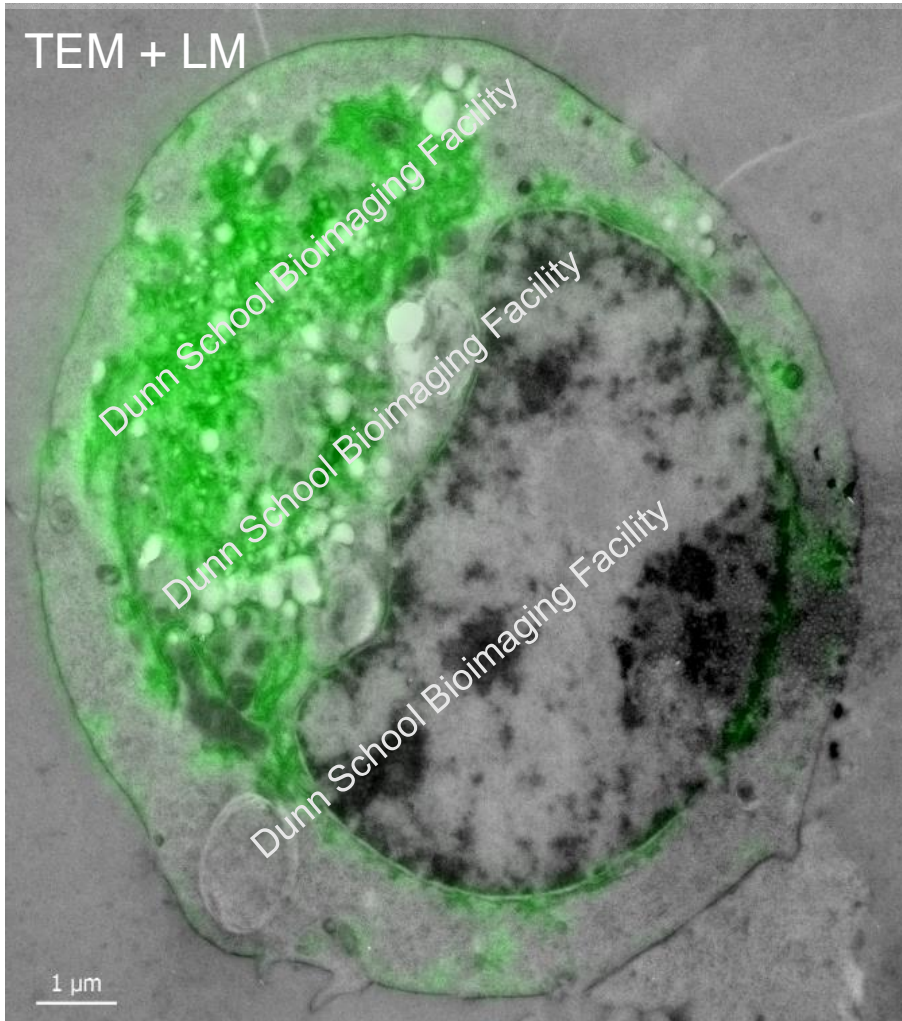
Advanced EM techniques

Correlative microscopy



Cells and tissue

Protein localisation – correlative microscopy

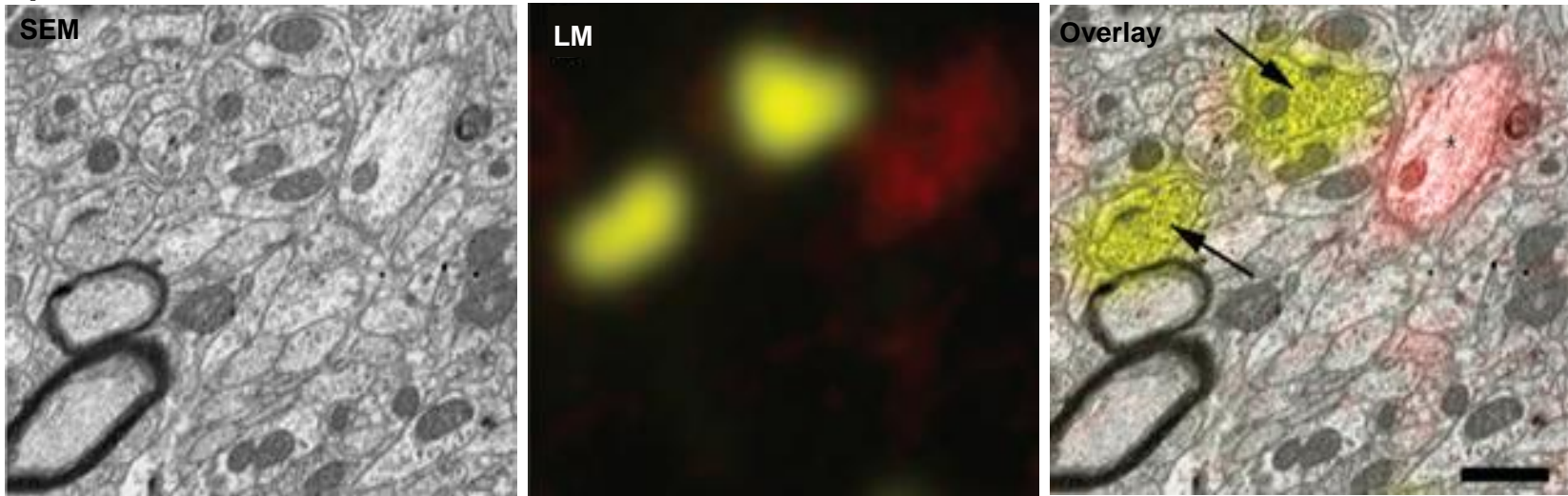


CLEM of HEK cells expressing EphA2-mVenus (E Johnson & R Kaufmann)

Correlative light and electron microscopy (CLEM)

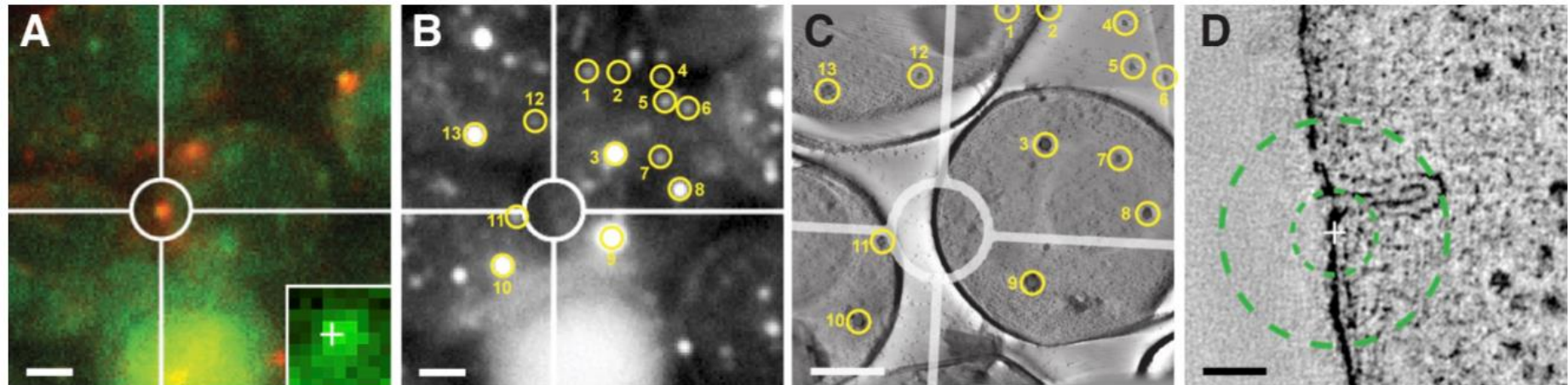
Cell/process identification

Specific cells



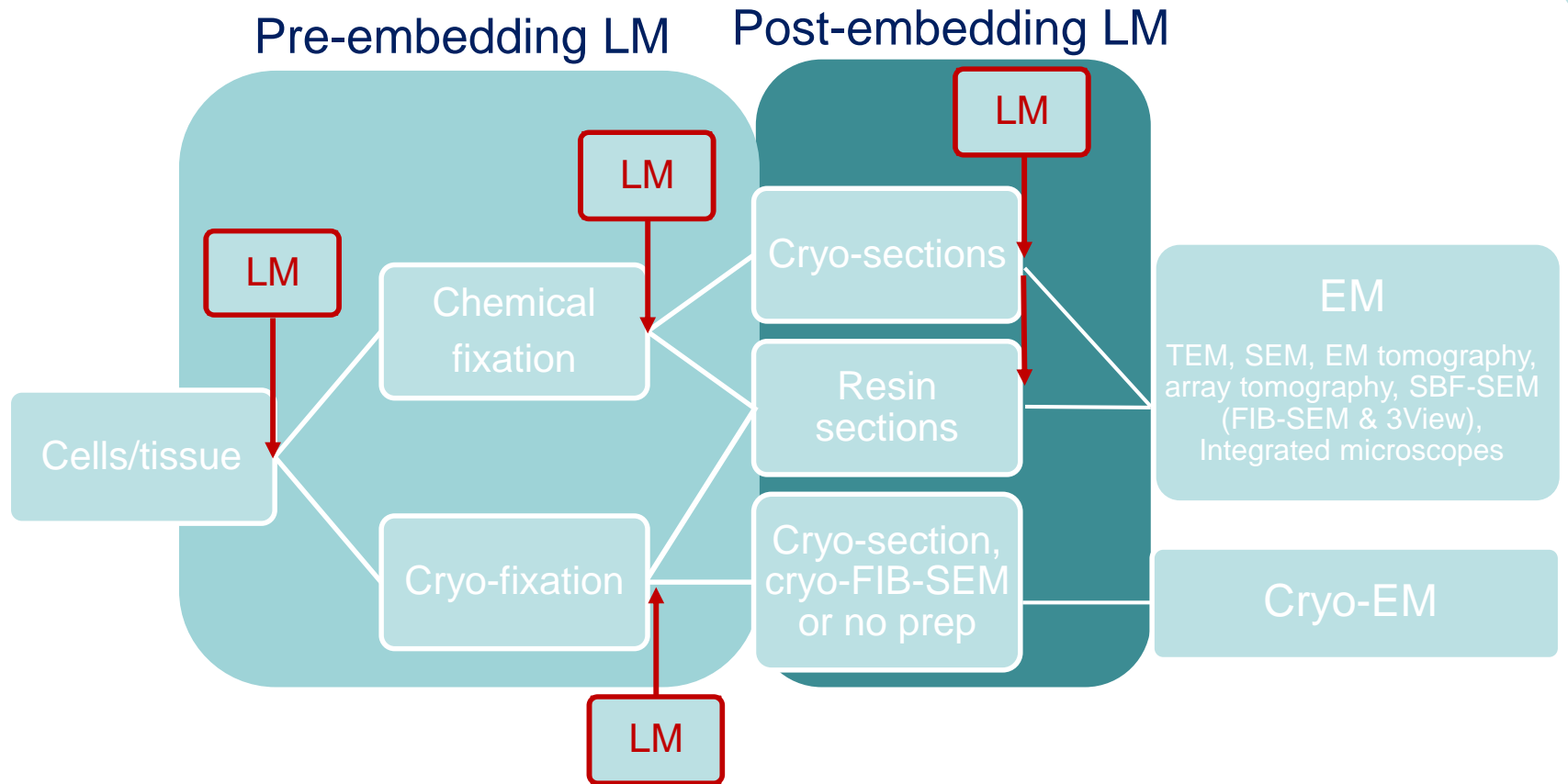
CLEM of two different projection neuron populations in Zebra Finch brain tissue (Oberti et al. 2011, *Front Neurosci*)

Rare events



CLEM of in-resin fluorescing endocytic patches in yeast cells (Kukulski et al. 2011, *JCB*, 192)

Simplified overview of CLEM workflows



Advanced EM techniques

Cryo-Electron Microscopy

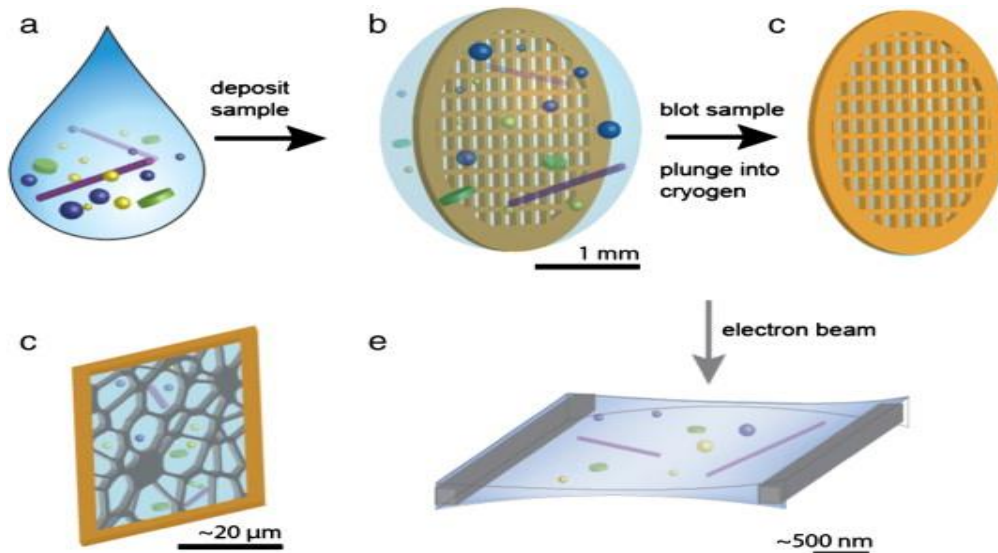


FEI Talos Arctica Cryo-TEM @ COSMIC

Cryo-TEM

Sample preparation

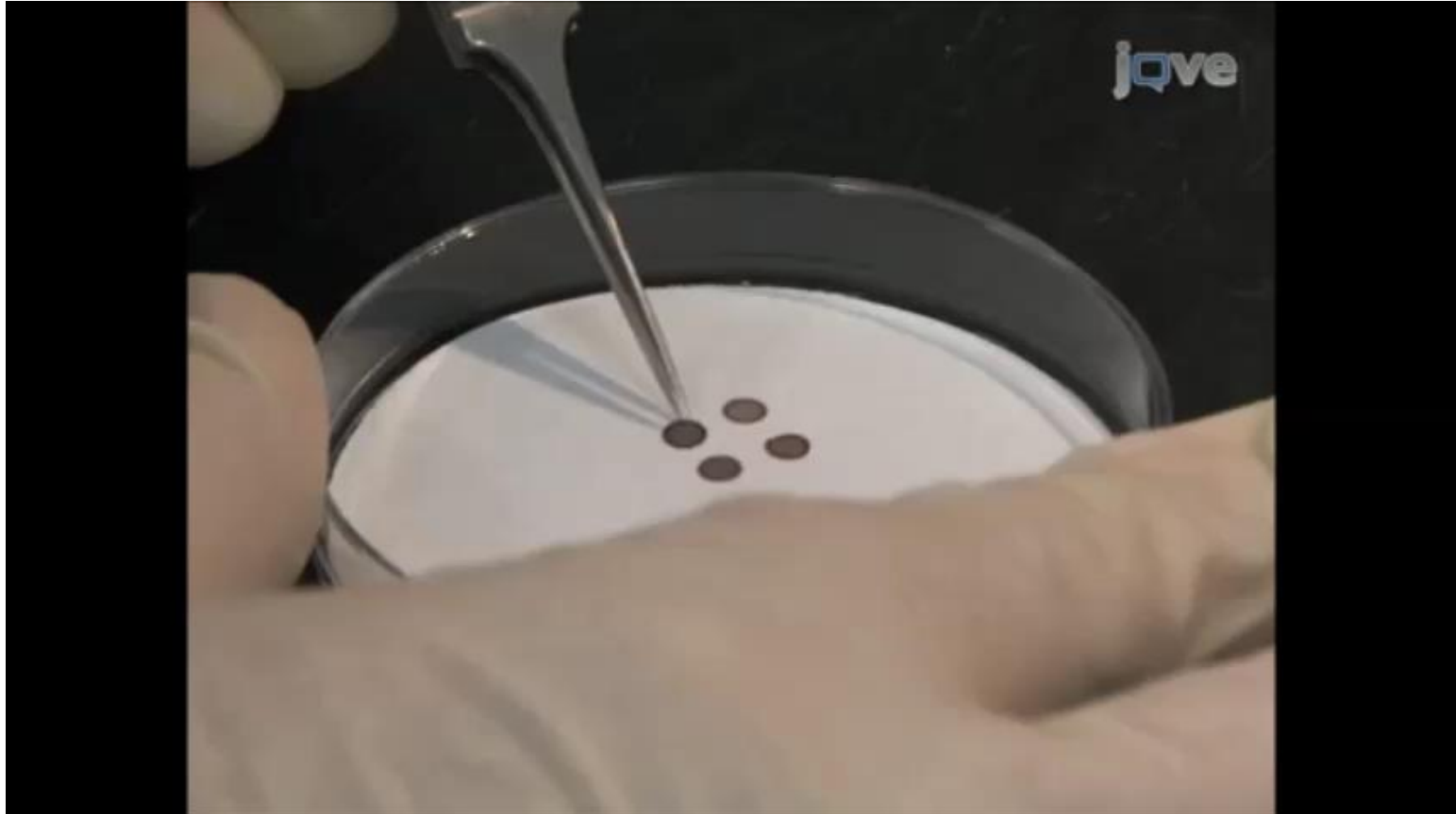
- You can freeze particulate samples and image under cryo conditions, which allows you to view them as close as possible to their native state.
 - Coat grids with plastic film and carbon
 - Apply the particulate specimen
 - Vitrify by plunge freezing into a cryogen (eg: ethane or propane)
 - Transfer to cryo-TEM under liquid nitrogen and image frozen



FEI Vitrobot for automated plunge freezing of grids

Cryo-TEM

Sample preparation

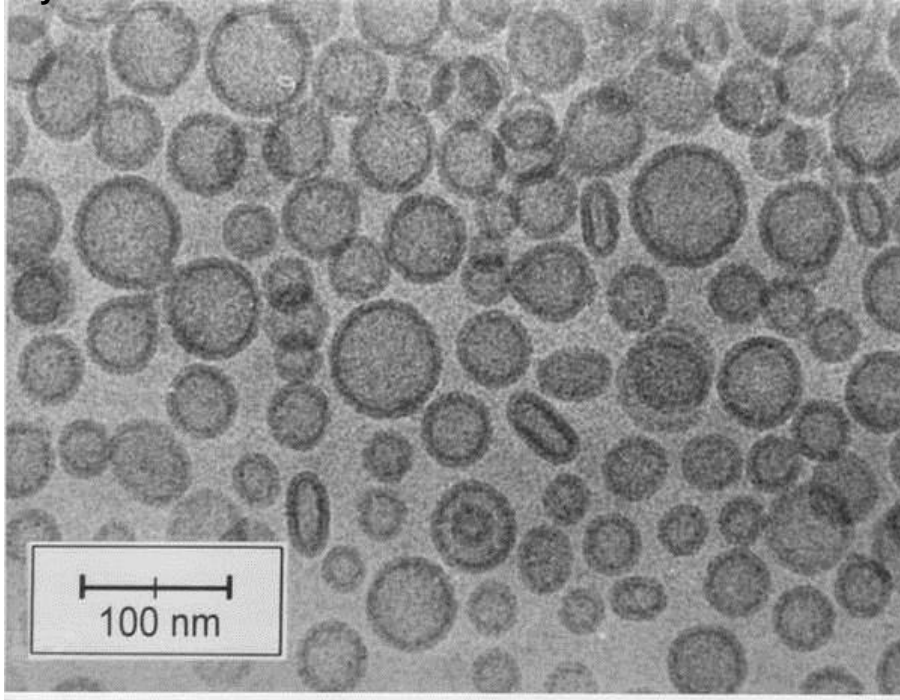


From: Chen et al (2010) J Vis Exp, 39: 1943

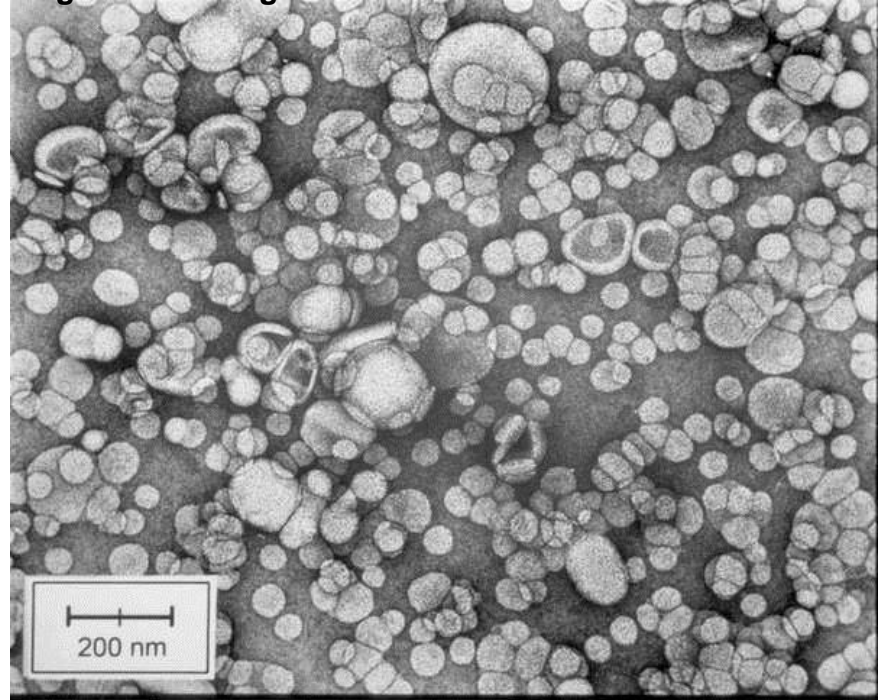
Cryo-TEM

Comparison to negative staining

Cryo-TEM



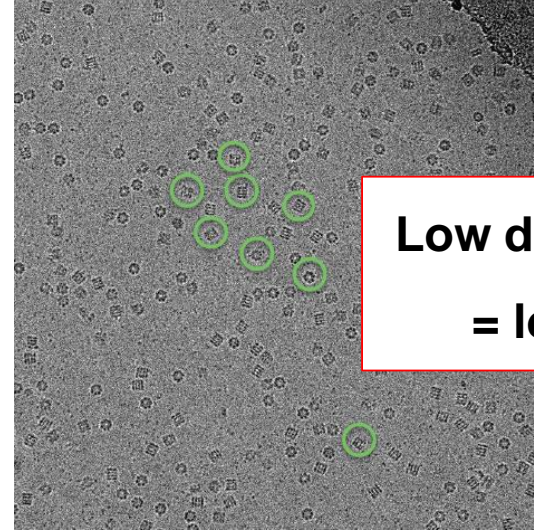
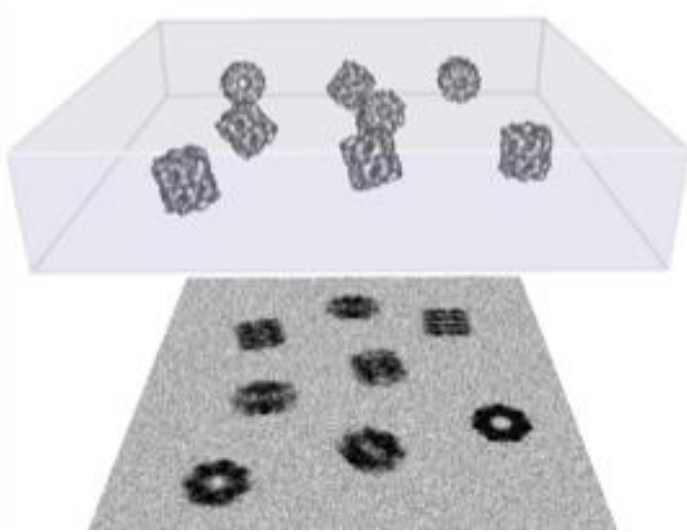
Negative staining



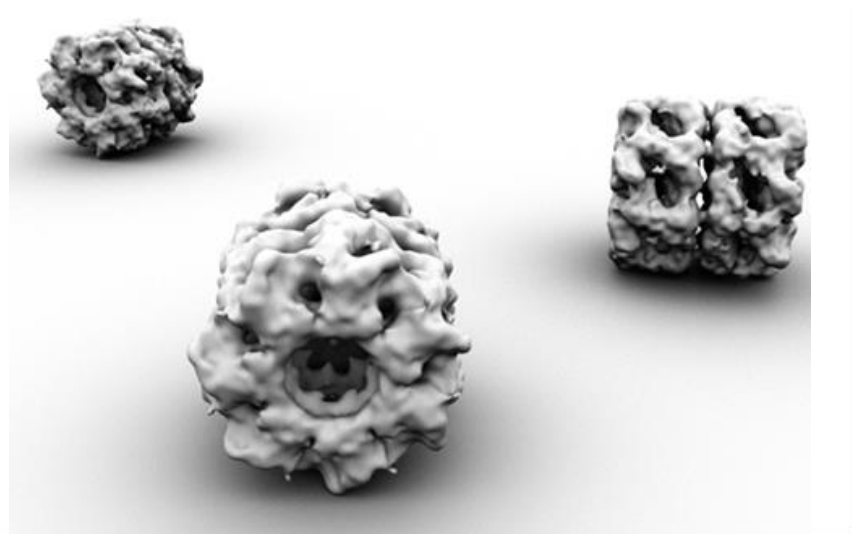
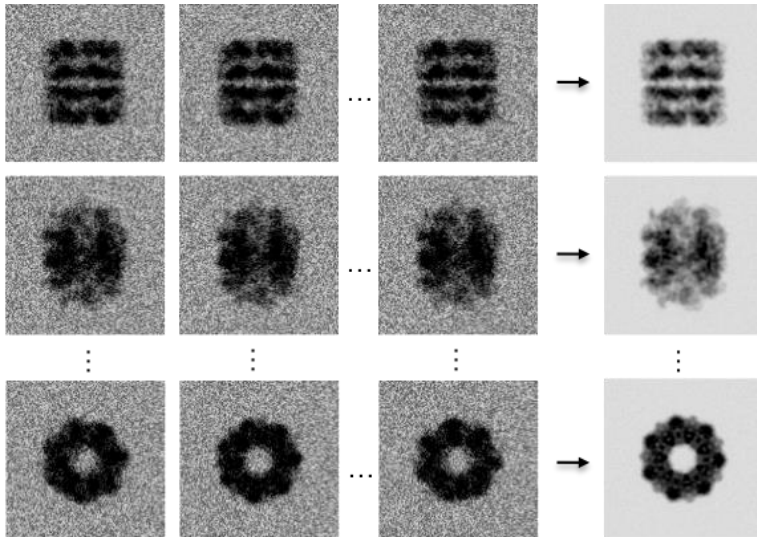
Dispersion of Egg-PC liposomes imaged frozen with Cryo-TEM (left) or negatively stained with uranyl acetate (right)
From: Laboratory for Soft Matter Electron Microscopy, University of Bayreuth

Cryo-TEM

Single particle imaging & reconstruction



**Low dose conditions
= low contrast**

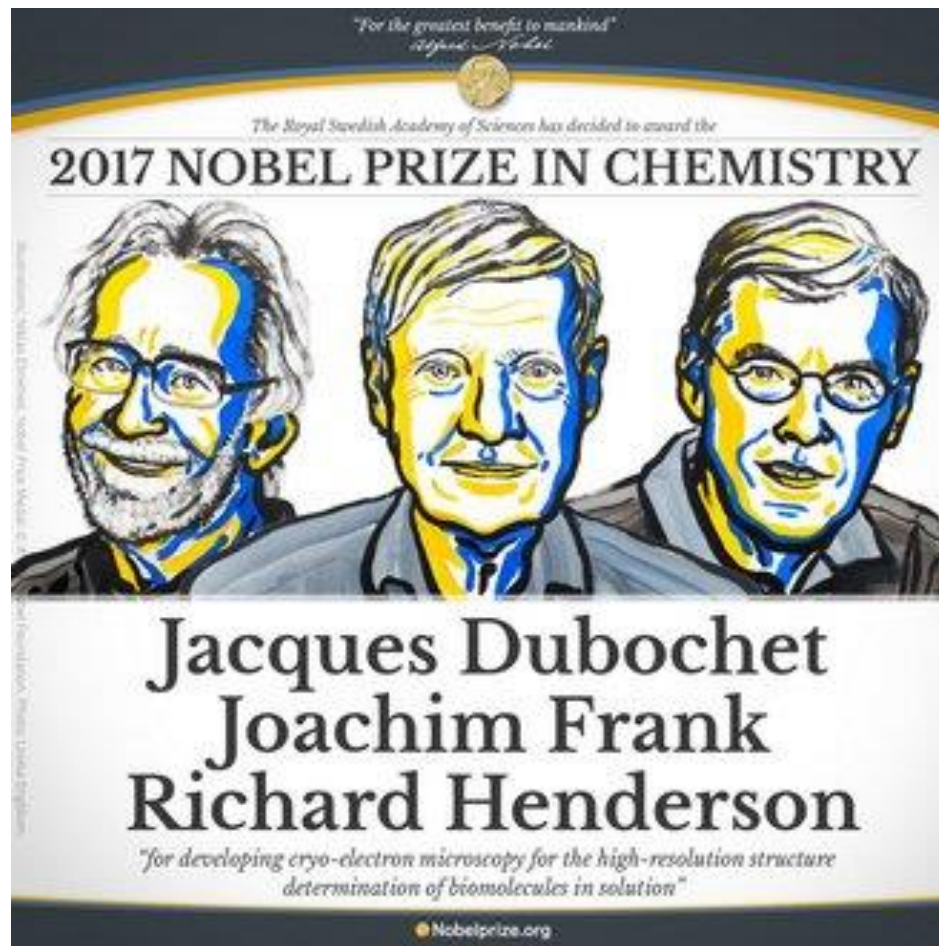


Single particle imaging and reconstruction of the GroEL chaperonin: purified complexes were applied to a grid and vitrified, then imaged with cryo-TEM. Thousands of images are collected and the same orientations are clustered together, averaged and back projected to render the complexes in 3D to 1 nm resolution. From: <http://people.csail.mit.edu/gdp/cryoem.html>

Cryo-Electron microscopy

Advances for structural biology

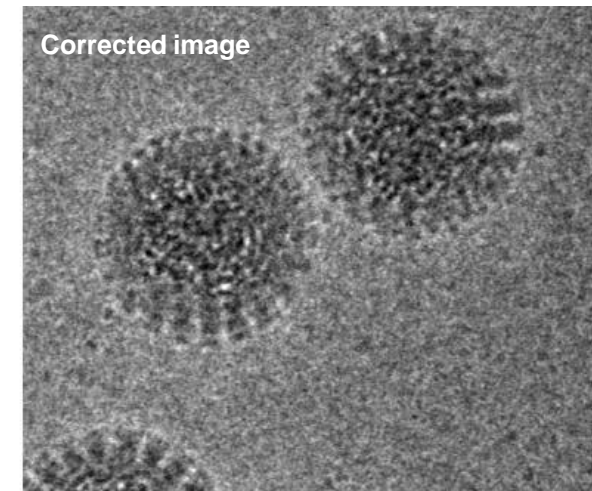
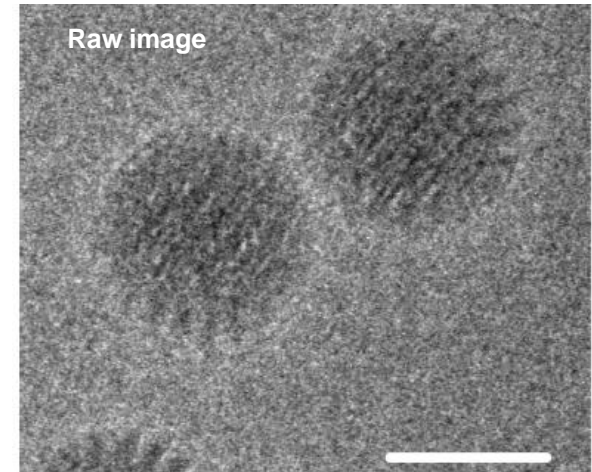
- The recent explosion of cryo-EM as a viable structural biology tool is due to several recent advances which have bought decades of work together



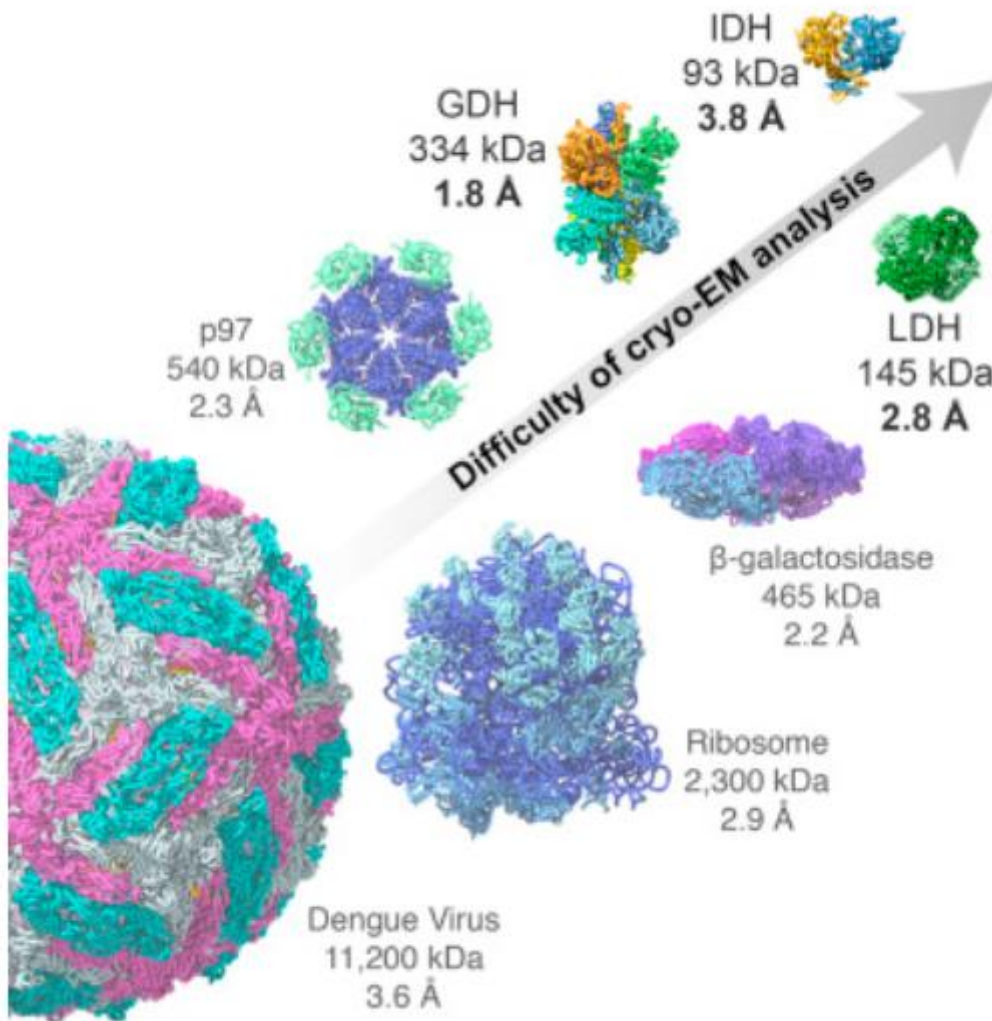
Cryo-Electron microscopy

Advances for structural biology

- Greater microscope stability
 - Titan Krios has constant power lenses for greater beam stability, a cube to reduce thermal drift
- Automated data acquisition
 - Allows data to be collected automatically over several days
- Direct electron detectors
 - Unlike CCDs, electrons are directly converted to electric signals.
 - Very sensitive and extremely fast, collecting 40 frames/sec.
 - This allows for dose fractionation and motion correction, which gives ~10x better resolution
- Computational and data processing advances



Cryo-TEM *Challenges*



- Proteins <200kDa are problematic due to low contrast
- Sample preparation optimisation: concentration, purity and stability of protein
- Vitrification optimisation: correct ice thickness, multiple orientations of protein, even distribution in ice

Degree of difficulty

Prep: High

Microscopy: High

Data analysis: High

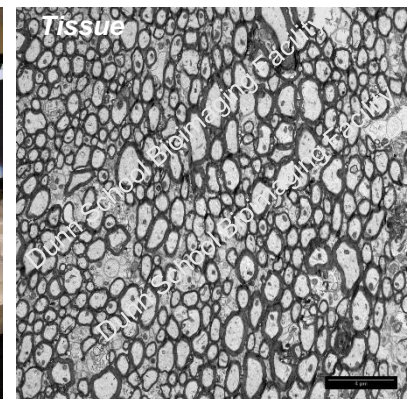
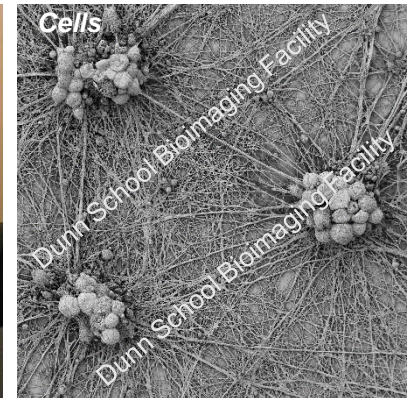
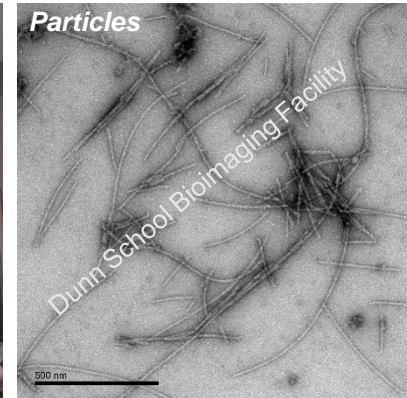
The Dunn School EM Facility & Central Oxford Single Molecule Imaging Center (COSMIC)

■ Staff:

- Errin Johnson (EM Facilities Manager)
- Raman Dhaliwal (EM Research Assistant)
- Adam Costin (Cryo-EM Research Assistant)

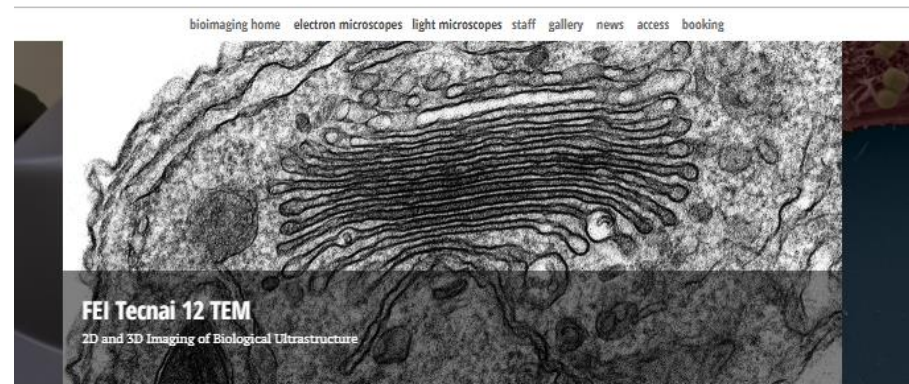
■ Instruments:

- FEI Tecnai12 120 kV TEM
- Zeiss Sigma 300 FEG-SEM
- Zeiss Merlin compact FEG-SEM + 3View
(sited at & shared with Oxford Brookes)
- FEI Talos 200c cryo-TEM
- FEI Arctica cryo-TEM with F3 detector
- FEI Titan Krios cryo-TEM with F3 and K2 GIF
- Biological EM specimen preparation lab



The Dunn School EM Facility & Central Oxford Single Molecule Imaging Center (COSMIC)

- For more information about our microscopes and the techniques mentioned in this talk, plus recommended papers/books to read and links to instructional videos, please see our website:
- <http://web.path.ox.ac.uk/~bioimaging/bioimaginghome.html>



TRANSMISSION ELECTRON MICROSCOPE (TEM)

The FEI Tecnai 12 Transmission Electron Microscope (TEM) enables both 2D and 3D imaging of specimen ultrastructure at up to 300,000x magnification. Images are acquired digitally using a bottom-mounted high resolution CMOS camera. A specialised specimen holder, together with SerialEM software, enables automated tilt series acquisition for 3D electron tomography.

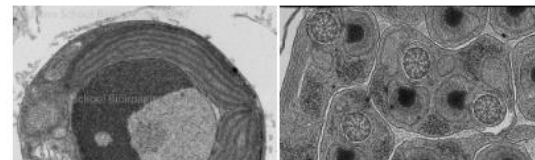
Specifications:

- Accelerating voltage: up to 120 kV
- Lanthanum hexaboride (LaB6) electron source
- Resolution: 0.49 nm (point)
- Magnification: 20x to 300,000x
- Single tilt specimen holder for standard imaging
- Fischione 2040 dual axis tomography holder
- Gatan Digital Micrograph 3.0 and SerialEM image acquisition platforms
- 16 Megapixel Gatan OneView™ CMOS camera

Applications:

- Screening negatively stained particulate samples (eg: viruses, bacteria, protein and liposomes)
- Characterisation of cellular ultrastructure (eg: mitochondria, centrosomes etc)
- Subcellular localisation of proteins using immunogold labelling
- Correlative light and electron microscopy (CLEM) to place fluorescent proteins/markers in ultrastructural context
- Electron tomography for high resolution 3D reconstructions of organelles and particulate samples

Example images:



EM LECTURES

[Micron EM Lecture 2016](#)

RESOURCES

[EMScope TEM virtual training](#)

[FEI website](#)

ELECTRON MICROSCOPY



ELECTRON MICROSCOPY

