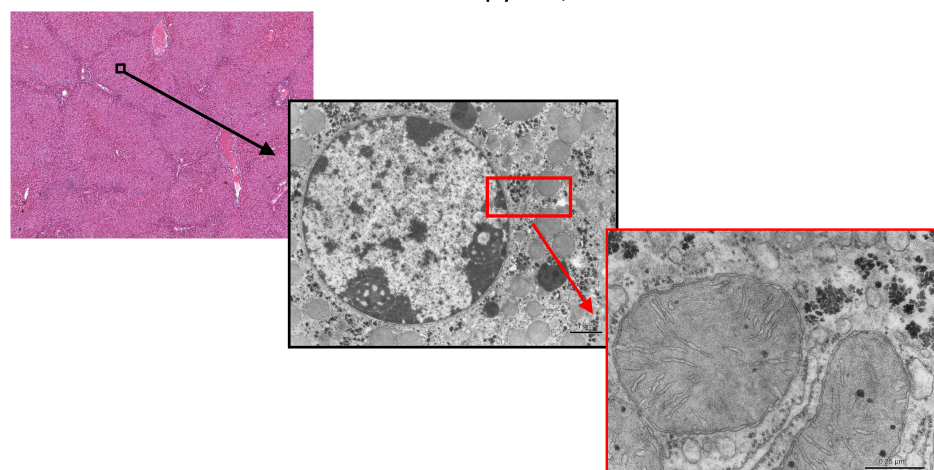
INTRODUCTION TO HISTOLOGICAL TECHNIQUES AND SAMPLE PREPARATION FOR LIGHT AND ELECTRON MICROSCOPY

Histology

- Resolution of naked eye 0,1 mm
- Resolution of light microscopy 0.5 μ m (usually)
- Resolution of electron microscopy 0,1 nm



Tissue processing for the light microscopy (LM)

(making of permanent preparations – slides)

- **SAMPLING** (obtaining of material cells, tissue pieces)
- **FIXATION** of samples (tissue blocks)
- **RINSING** (washing) of samples
- **EMBEDDING** of samples embedded blocks
- CUTTING of blocks sections
- AFFIXING of sections
- **STAINING** of sections
- MOUNTING of sections

SAMPLING

- A small piece of organ (tissue) is sampled and quickly put into the fixative medium.
- Biopsy during surgical dissection of organs in living organism
 - = excision
 - = puncture (liver or kidney parenchyma, bone marrow)
 - = curettage (uterine endometrium, adenoid vegetation)
- Necropsy from dead individual (sections); in experiments laboratory animals are used and tissue have to be sampled as soon as possible after the break of blood circulation
- The specimens shouldn't be more than $5 10 \text{ mm}^3$ thick and fixation should follow immediately.

FIXATION

- Definition: denaturation and stabilization of cell proteins with minimum artifacts)
- The reason of fixation: freshly removed tissues are chemically unstable dry, shrink, undergo hypoxia, autolysis and bacteriological changes
- To stop or prevent these changes and preserve the structure tissue samples have to be fixed. During the fixation, all tissue proteins are converted into inactive denaturized (stable) form.
- 3 main requirements on fixatives:
 - good preservation of structure
 - quick penetration into tissue block
 - no negative effects on tissue staining

- <u>Fixatives</u>: solutions of various chemicals
 - organic fixatives <u>ALDEHYDES</u> formaldehyde (*most frequently used for LM*)
 - glutaraldehyde (used for EM)
 - ALCOHOLS 96 100 % (absolute) ethylalcohol
 - ORGANIC ACIDS glacial acetic acid, picric acid,

trichloracetic acid

- inorganic fixatives INORGANIC ACIDS chromic acid, osmium tetraoxide (OsO4)
 - SALTS OF HEAVY METALS mercuric chloride HgC₁₂
- compound fixatives mixtures (two or more chemical components to offset

undesirable effects fo indiviual (simple) fixatives.

FLEMMING's fluid – with OsO₄

ZENKER's and HELLY's fluid, SUSA fluid – with HgCl₂

BOUIN's fluid - with picric acid

CARNOY's fluid – with alcohol

Performance: fixatives are carried out at room temperature, the duration varies between **12 – 24 hours**, specimen must be covered by 20 – 50 times fixative volume:

Ratio of tissue block volume to fixative volume 1 cm³: 20 – 50 cm³

RINSING and EMBEDDING

 All samples should be washed to remove the excess of fixative; the choice of rinsing medium is determined by type of fixative: running tap-water or 70-80% ethanol

 Relevance of embedding: tissues and organs are brittle and unequal in density, they must be hardened before cutting

Embedding media

- water soluble gelatine, celodal, water soluble waxes
- anhydrous paraffin, celoidin

EMBEDDING into PARAFFIN

- dehydration to remove water from fixed samples by ascending series of ethanol is used (50%, 70%, 90%, 96%. each step - 2 – 6 hours
- clearing the ethanol must be replaced with organic solvatant that dissolves paraffin – benzene or xylene
- infiltration melted paraffin wax (56°C) is used; 3 x 6 hours.
- casting (blocking out) moulds (plastic, paper or metal chambers) are used for embedding.
- The moulds are filled with melted paraffin, tissue samples are then placed inside and immediately immersed in cold water to cool paraffin quickly down.
- These paraffin blocks are ready for trimming



Leica TP 1020

Automated device for tissue dehydration

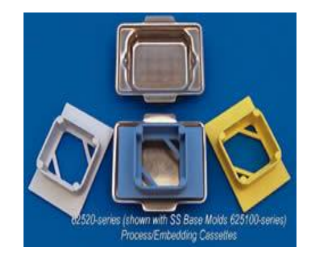


Paper chambers

- metal



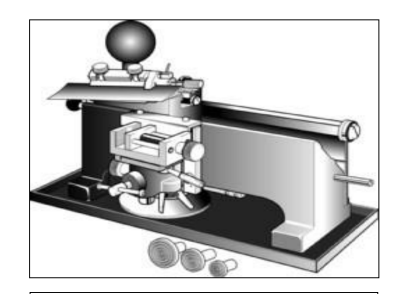




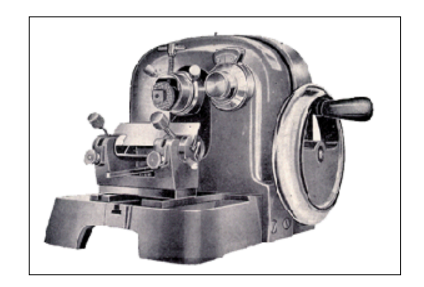


CUTTING

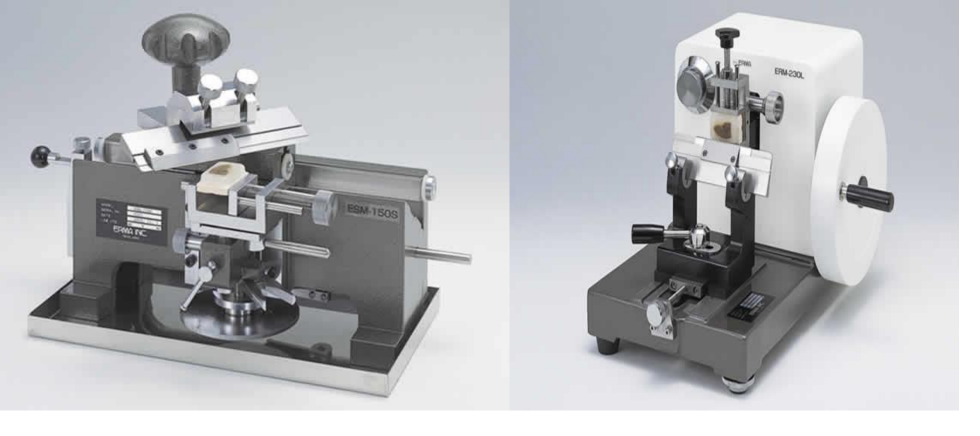
• Microtome — a machine with automatic regulation of section thickness: $5-10 \mu m$ is optimum.



sliding microtome – block is fixed in holder, knife or razor moves horizontally



rotary microtome – knife is fixed, block holder moves vertically



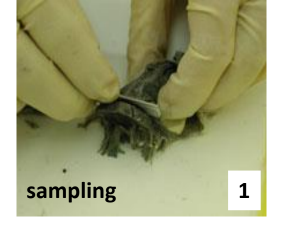
Sliding microtome

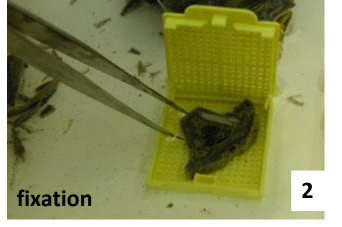
Rotary microtome

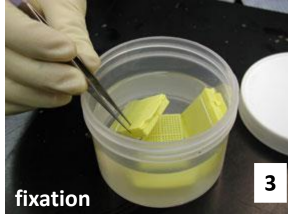
Freezing microtome (**Cryostat**)
= rotary microtome housed in freezing box
(- 60° C)

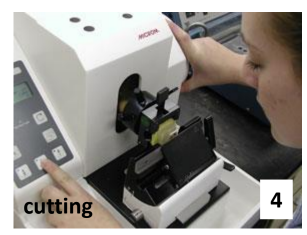
Cutting of frozen tissue without embedding





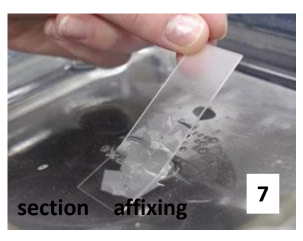


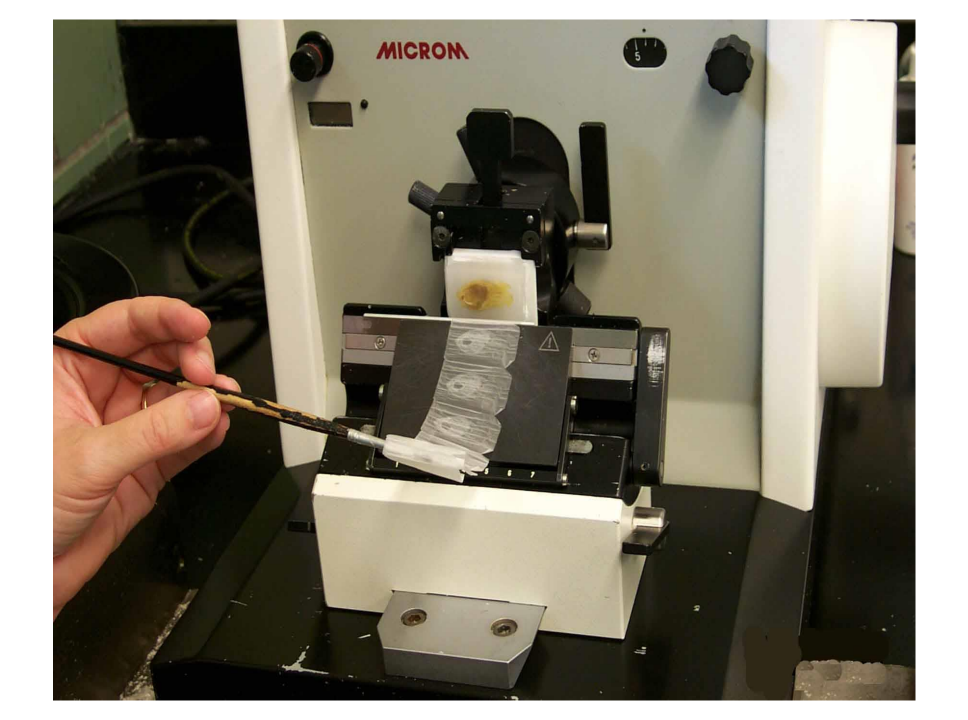


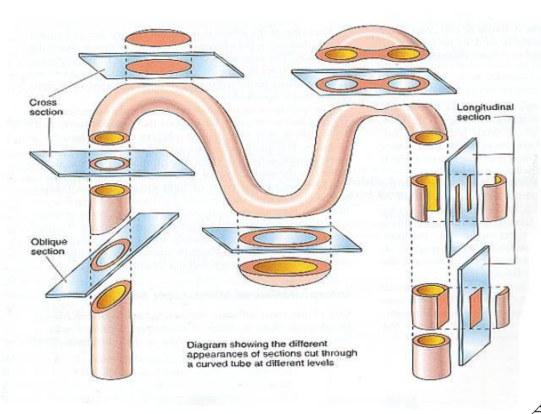




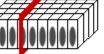








perpendicular section











oblique section

AFFIXING

- Mixture of glycerin and egg albumin or gelatin
- Section are transferred from microtome razor or knife on the level of warm water (45° C), where they are stretched; then they are put on slides coated with adhesive mixture; excess of water is drained and slides are put in incubator (thermostat, 37° C) over night to affixing of sections.



Stretching of sections on warm water



Stretching on a warm plate



STAINING

- Different cell or tissue structures are not apparent without staining.
- Cellular structures exhibit different affinity to <u>staining dyes</u>

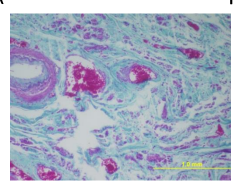
alkaline dyes (basic or nuclear) – react with anionic groups of cell and tissue components

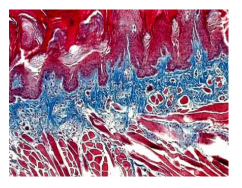
basophilia – basophilic structures in the cell

acid dyes (cytoplasmic) – react with cationic groupsacidophilia – acidophilic structures in the cellneutrophilia – no reaction

Staining methods:

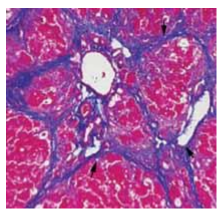
<u>routine</u> – HE, AZAN (demonstrate all components of tissue)





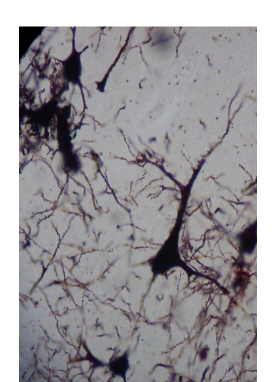
HE – the most frequent used method

special
visualizes only special structures



Lipid droplets detected by oil red

<u>impregnation</u>by silver salt for detectionof nerve or reticular fibers

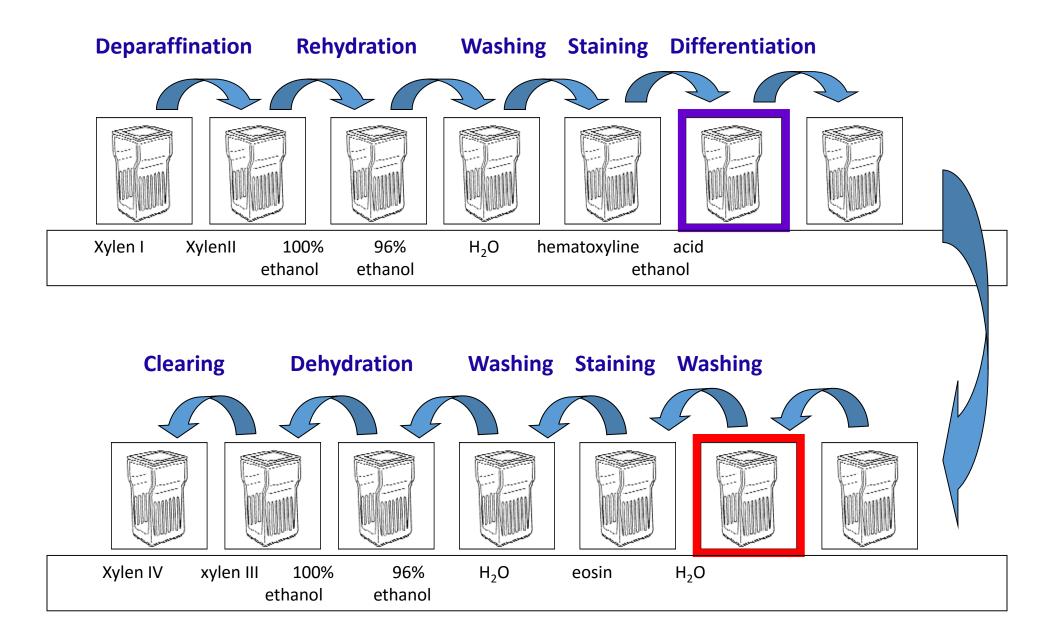


ROUTINE STAINING with HEMATOXYLINE – EOSIN (HE)

Hematoxyline – basic (nuclear) dye Eosin – acid (cytoplasmic dye

- <u>Staining procedure</u>:
- paraffin must be removed (dissolved) by xylene
- sections are rehydrated in descending series of ethanol (100% \rightarrow 96% \rightarrow 80%)
- staining with hematoxyline
- differentiation in acid ethanol and water (excess of dye is removed)
- staining with eosin
- rinsing in water (excess of dye is removed)
- dehydration in graded ethanol series (80% \rightarrow 96% \rightarrow 100%)
- clearing in xylene

HEMATOXYLINE – EOSIN (HE)



Staining results:

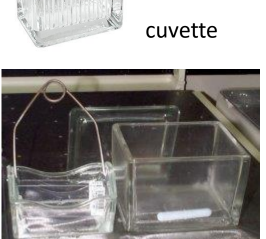
HE = Hematoxyline – Eosin
 nuclei – bright clear blue or dark violet
 cytoplasm and collagen fibers – pink
 muscle tissue – red

• <u>HES</u> = *Hematoxyline* – *Eosin* – *Safron* connective tissue – <u>yellow</u>

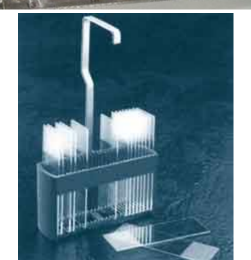
AZAN = AZocarmin - ANiline blue - orange G
 nuclei - red
 erythrocytes - orange
 muscle - red
 collagen fibers - blue

Staining tools:





flask



slides holder (basket)







Automatic slide stainer

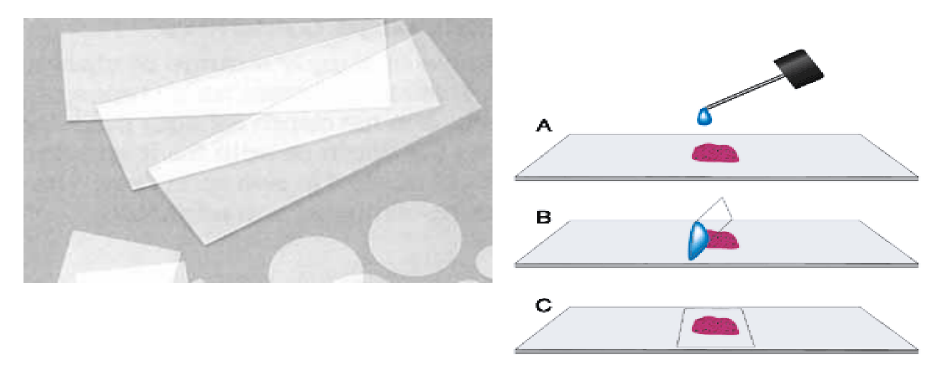


staining set of boxes with media



MOUNTING

• Finally, preparates are closed with coverslip (coverglass) to form a <u>permanent</u> <u>preparate</u>. Small amount of mounting medium must be placed between stained section and the coverslip.

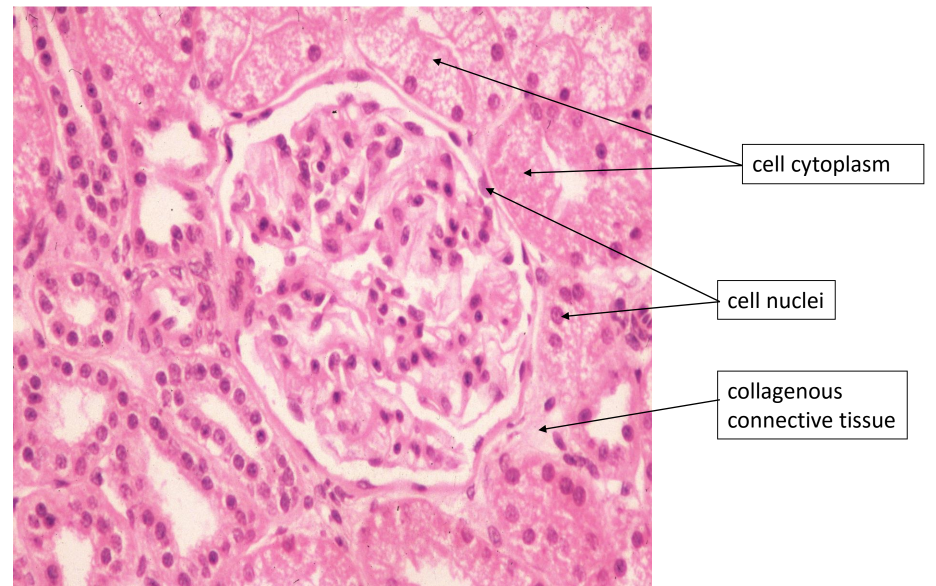


• Mounting media: soluble in xylene – canada balsam soluble in water – glycerin-gelatine, arabic gum

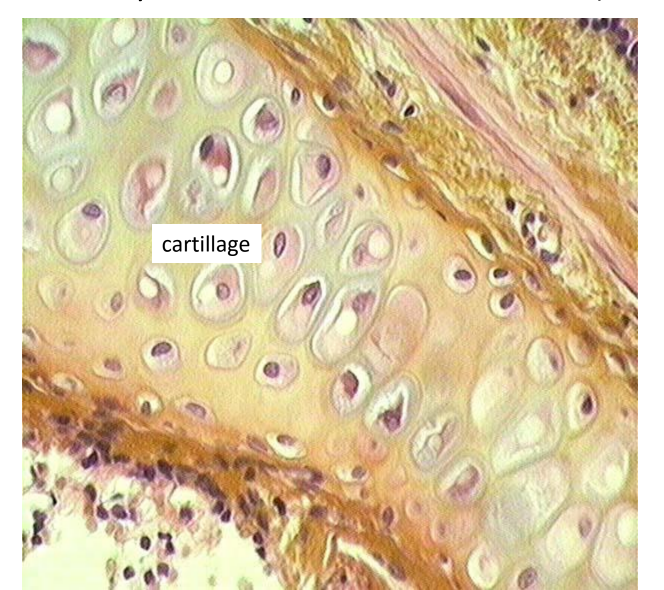


Permanent histological slides for study in the light microscope

Hematoxyline and eosin (HE)

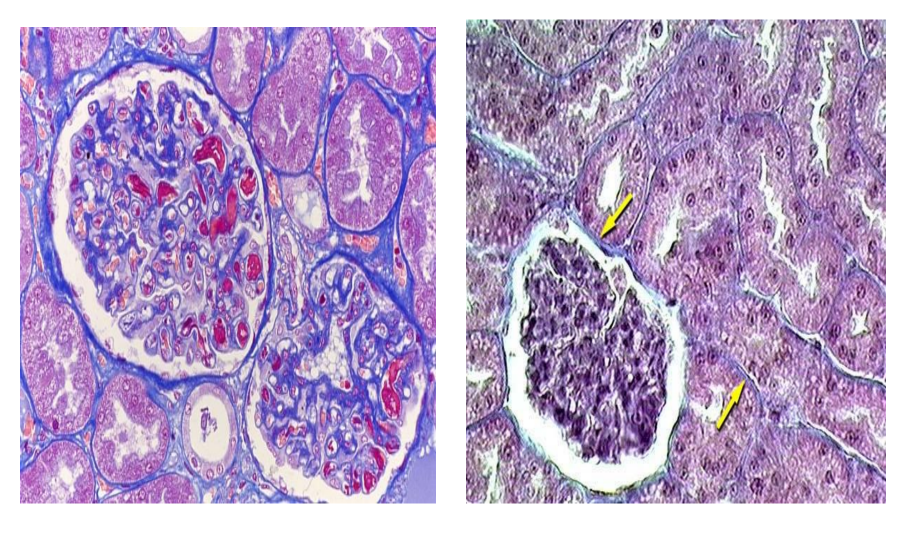


Hematoxyline, eosin and saffron (HES)



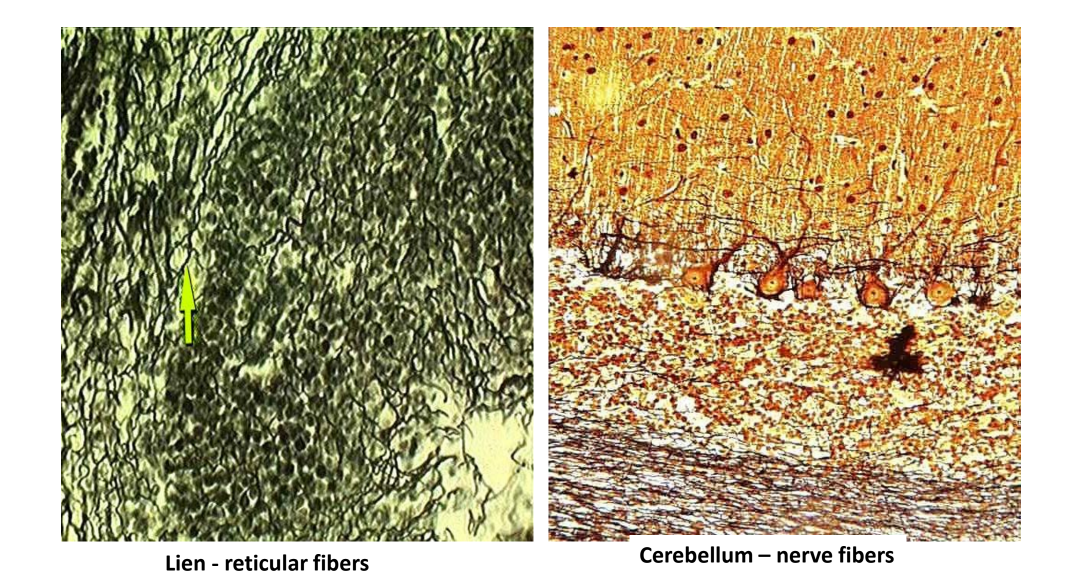
Collagenous fibers of connective tissue are yellow after staining with saffron

Azocarmine and aniline blue (AZAN)



Kidney – collagen connective tissue

Impregnation of tissue with silver

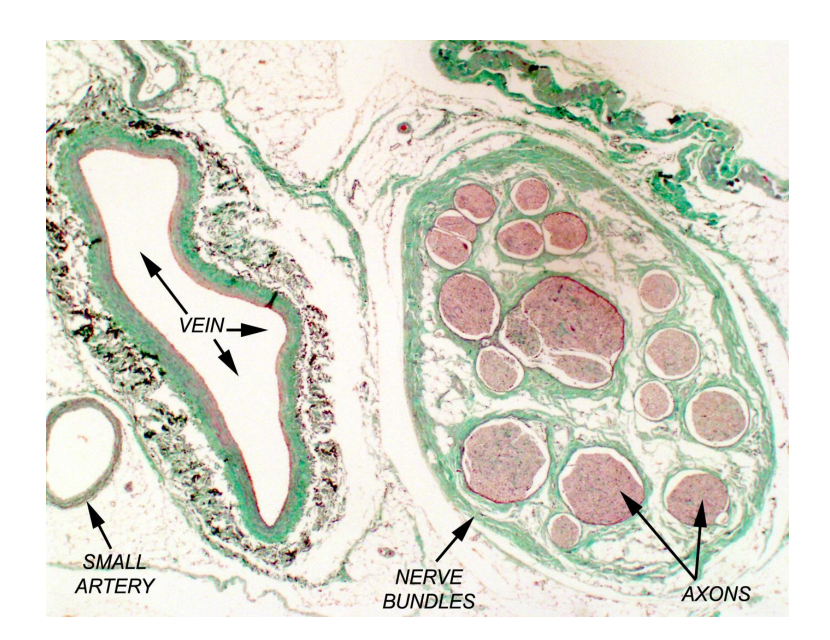


Iron hematoxyline



Skeletal muscle cells (fibers)

Green trichrome

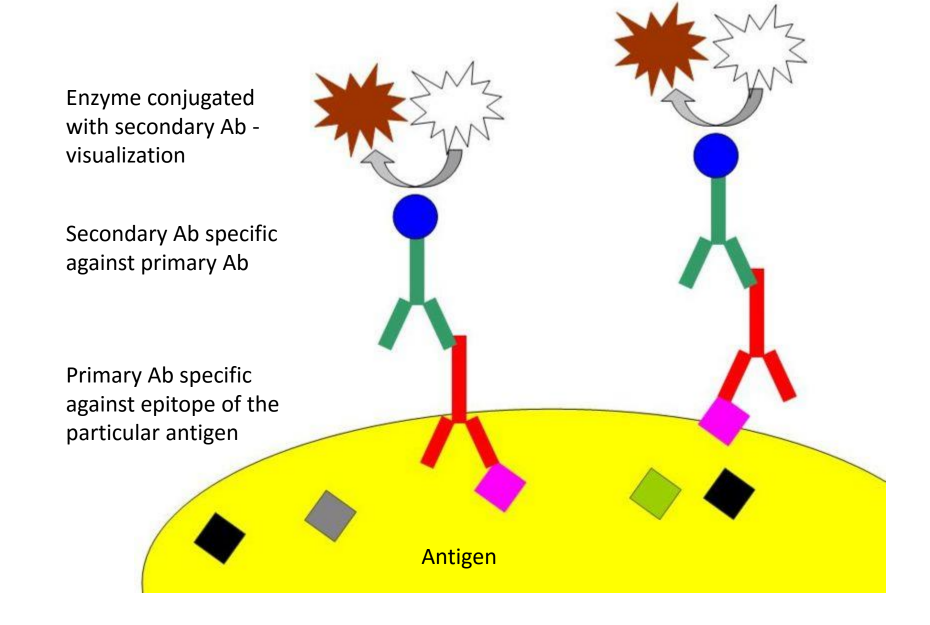


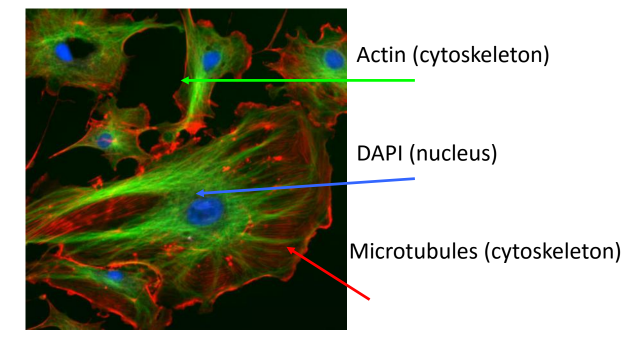
Histochemistry and Immunohistochemistry

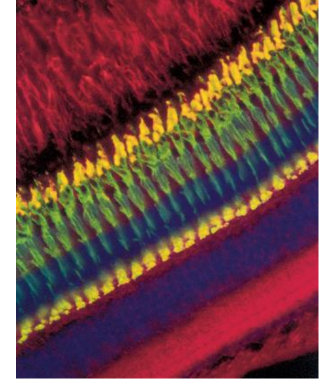
• Relevance:

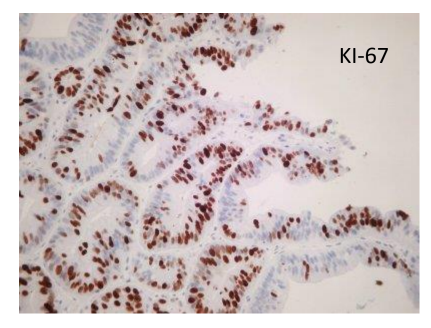
various chemical compounds detected "in situ" (proteins, AA, NA, saccharides, lipids, enzymes, pigments, inorganic substances – Fe, Ca, Zn)

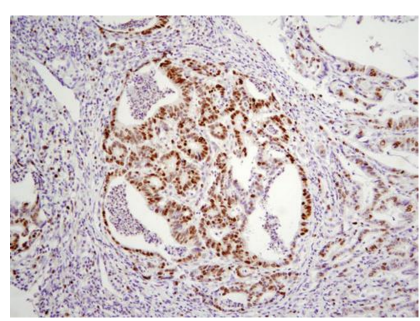
Various epitopes detected by immunotechniques











Tissue processing for the EM

pH of all solutions (media) must be buffered on 7.2 – 7.4
 Cacodylate or phosphate buffer is frequently used.

Absolutely dustfree environment

Solutions (media) have to be precise (artifacts)

Tissue processing for the EM

- SAMPLING immediately after arresting of blood circulation, tissue block sized no more than 1mm³
- FIXATION glutaraldehyde (binds amine groups) + OsO₄ (binds lipids)
 are used as double fixation
- **RINSING** distilled water
- DEHYDRATION ethanol
- **EMBEDDING** gelatin capsule or plastic forms are filled with some medium (which can be polymerized from liquid to solid form) and pieces of fixed tissue are placed into this medium. Epoxyd resins (Epon, Durcupan, Araldite) are usually used as in water insoluble media.
- CUTTING ultrathin sections (in ultramictomes)
- CONTRASTING ≈ staining

EMBEDDING

gelatin (1) or plastic (2) capsules

capsule holder (3)

embedding plates (4, 5)

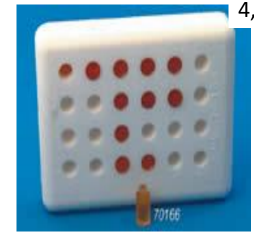






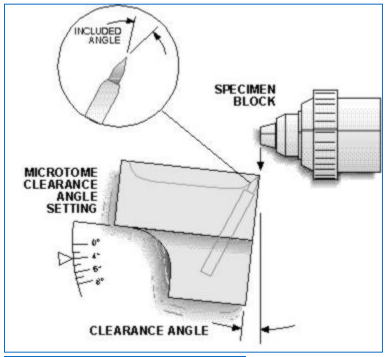


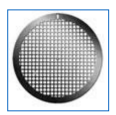
Embedded blocks prepared for cutting

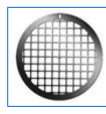




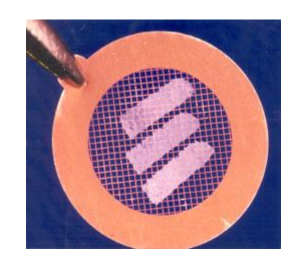
CUTTING



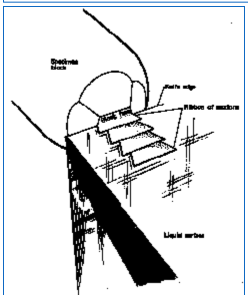


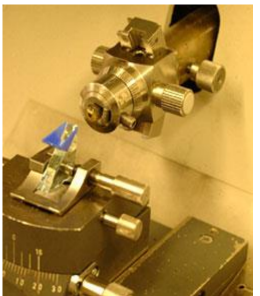


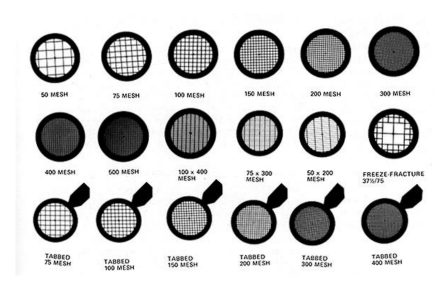












CUTTING

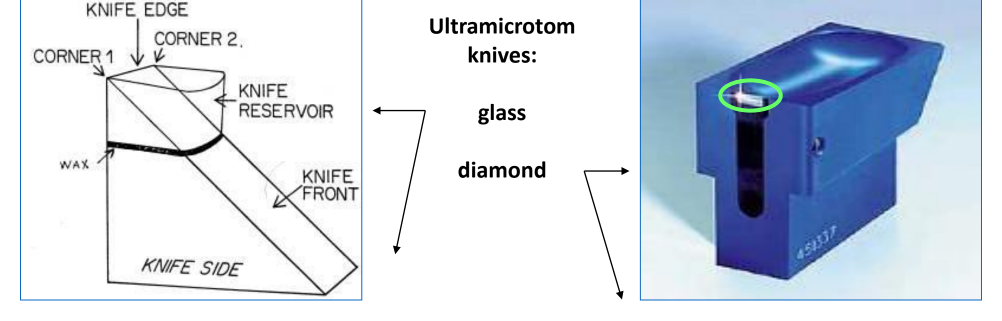
Ultrathin sections (70 – 100 nm) - ultramicrotomes.

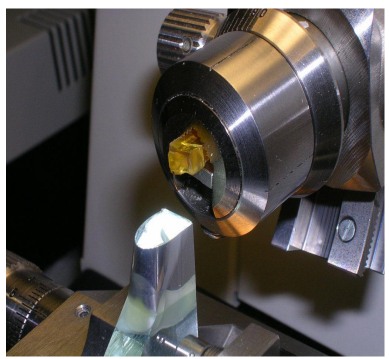
Glass or diamond (b) knives with water reservoir are used

Sections slide flow on water in small container attached to the knive

Supporting grids



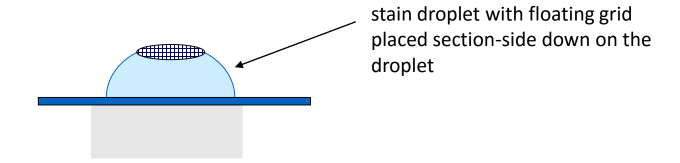






CONTRASTING (=STAINING)

 principle of differentiation of structures – different dispersion of beam of electrons depending on atomic weight of elements.
 "electron dyes" are thus mixtures of heavy metals: uranylacetate or lead citrate



Differences between LM and EM

LM EM

 $< 1 \text{ mm}^3$

seconds

glutaraldehyde

1 - 3 hours

Ultramicrotomes

50 - 100 nm

heavy metals

(uranylacetate,lead citrate)

Sampling < 1 cm³ minutes

Fixation formaldehyde 12 – 24 hours

Embedding paraffin epoxid resins (Durcupan)

 $\begin{array}{c} \text{Cutting} & \text{microtome} \\ \text{Thickness of sections} & 5-10 \ \mu\text{m} \end{array}$

Staining (LM) dyes contrasting (EM) (hematoxyline – eosin)

Mounting (only LM)

Result histological slide (preparate) photograph of ultrathin section

