Overview of Immunohistochemistry and Histochemistry (講義·英語)

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免疫組織染色と組織化学染色の原理と有用性について英語で概説する。

Histochemistry and immunohistochemistry are classic research techniques, yet they are still essential methods to identify expression of target molecules on tissue sections. By taking this class, you will learn basic principles of histochemistry and immunohistochemistry and recent advances in imaging technologies using fluorescence probes.

Program of 2023 Intensive Course in Basic Science Fundamentals & Multidisciplinary Seminars / Central Research Laboratory Special Seminars

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Overview of Immunohistochemistry and histochemistry

免疫組織化学染色と組織化学染色の原理

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Learning objectives

- 1. To learn how pathologists examine tissues and cells with different methods.
- 2. To understand the principle of histochemistry and immunohistochemistry (IHC).
- 3. To understand how to detect the antigen and antibody in IHC.
- 4. To learn how to improve the specificity and sensitivity of IHC.

Pathology

Pathology is a morphology based medical specialty that diagnoses diseases, mostly through analysis of tissue under the microscope.





Squamous cell carcinoma

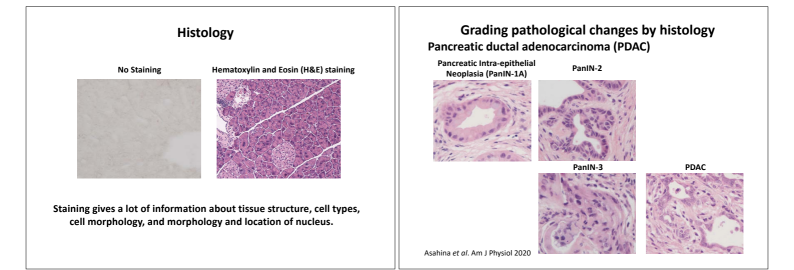


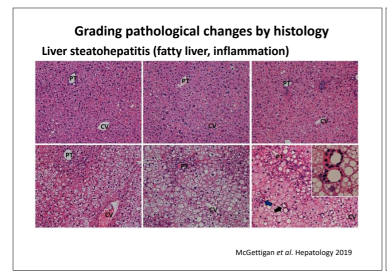
columnar shaped cells with glandular growth pattern

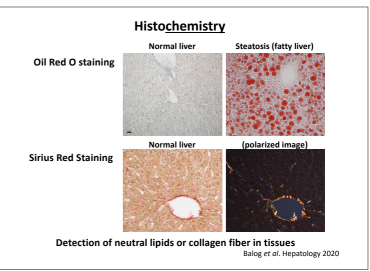
Experimental Pathology

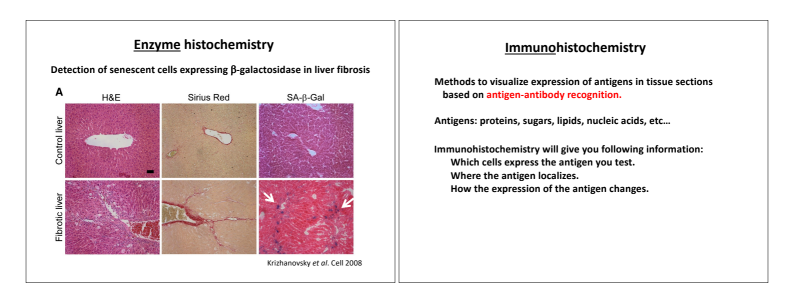
To reproduce pathogenesis of human disease using animals, such as mice, rats, or monkeys.

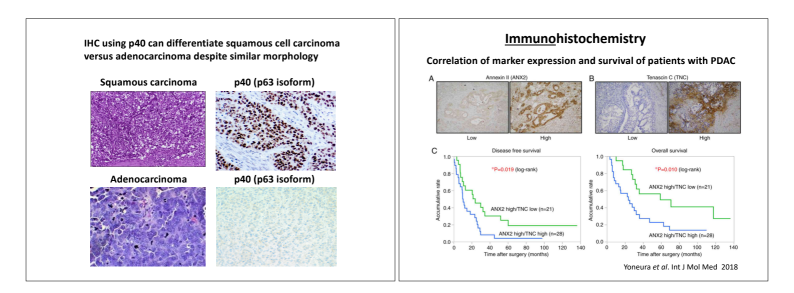
To understand cellular and molecular mechanisms underlying pathogenesis of the disease.

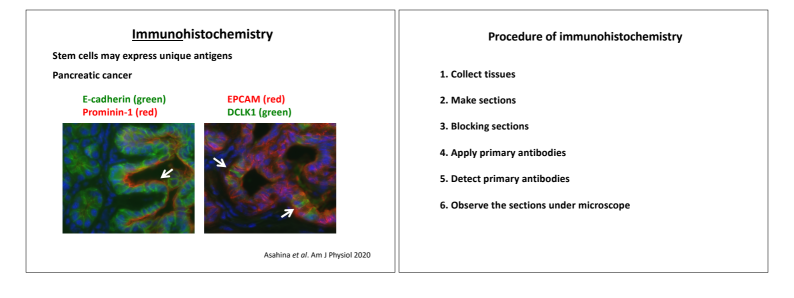


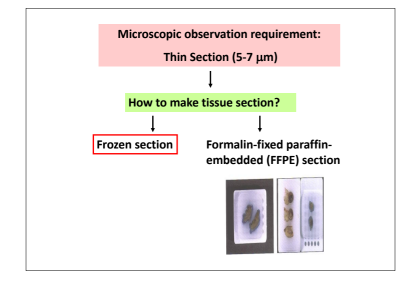












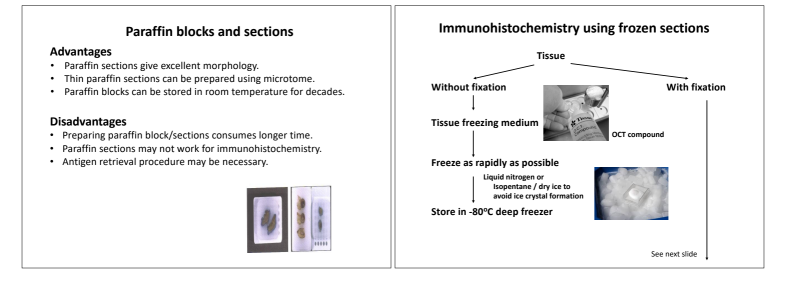
Frozen sections

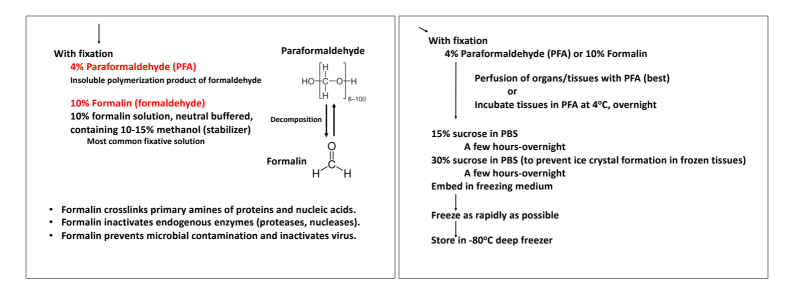
Advantages

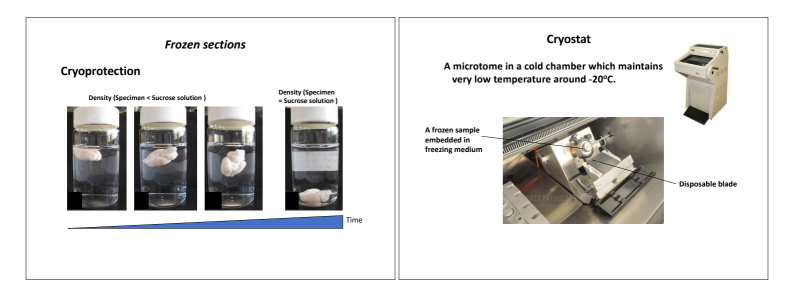
- Frozen samples can be prepared in short time with or without fixation of tissues.
- Proteins may keep their native structure in frozen sections.
- Many antibodies may work well in frozen sections.

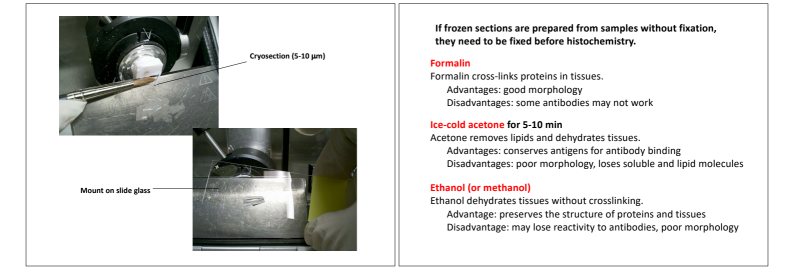
Disadvantages

- Frozen sections give poor morphology.
- Cryostat (expensive) is necessary for making frozen sections.
- Frozen samples/sections are needed to be stored in freezers.









Procedure of immunohistochemistry in frozen sections

- Make cryosections (7 µm) 1.
- 2. Dry
- 3. Fix the sections if necessary
- Remove freezing medium in PBS 4.
- Blocking the sections for 30 min (usually 5% goat or donkey serum and 5. 0.2% bovine serum albumin in PBS)
- 6. Apply primary antibody for 1 hour room temperature or over night at 4°C
- 7. Wash with PBS
- Apply secondary antibody for 30 min room temperature 8. 9.
- Wash with PBS 10. Detection of the secondary antibody
- 11. Counterstain nuclei if necessary

Primary antibody

Polyclonal antibody vs. monoclonal antibody

Polyclonal antibody

Advantages

may recognize different epitopes of the antigen Disadvantages

may vary its affinity to the antigen among a host animal may be obtained limited amount from a host animal

Monoclonal antibody

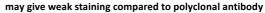
Advantages

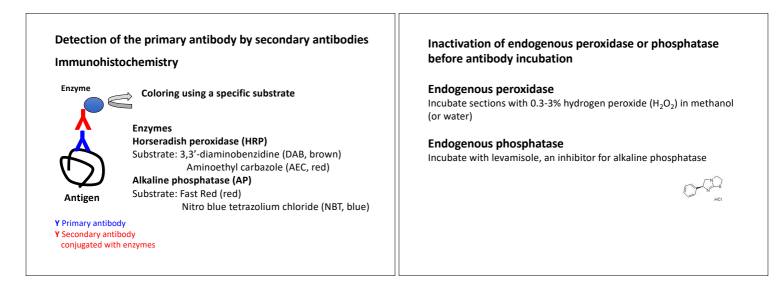
recognizes a single epitope of the antigen can be produced stable quality from a hybridoma clone

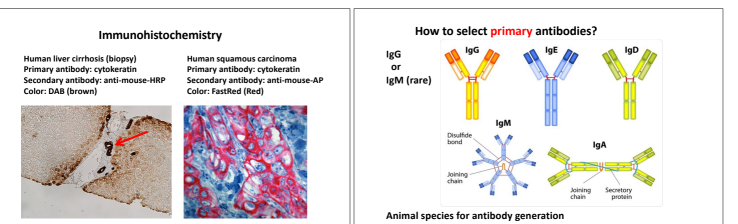
Antigen

Antigen

Disadvantages is often produced from mice

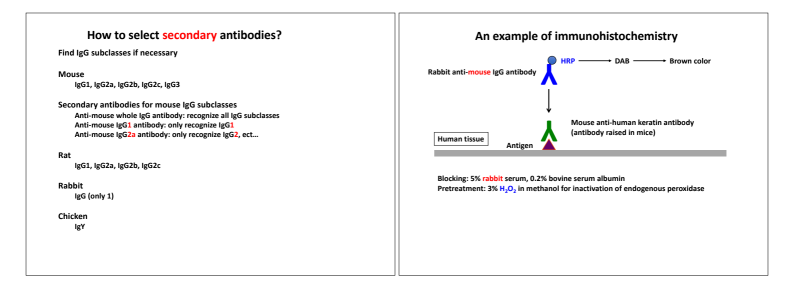


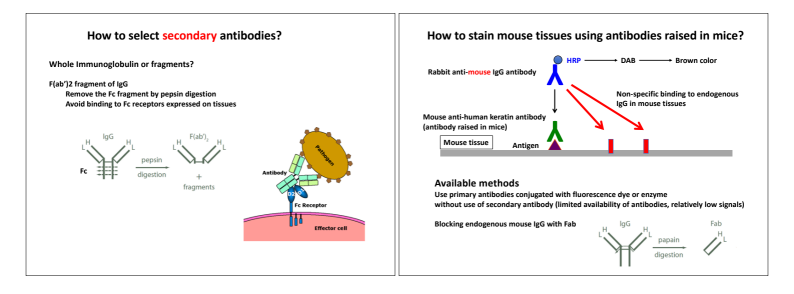


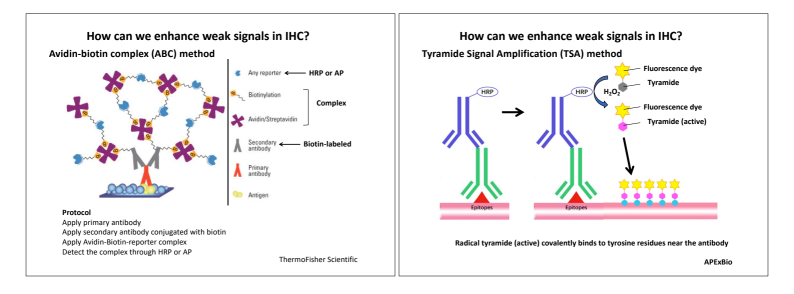


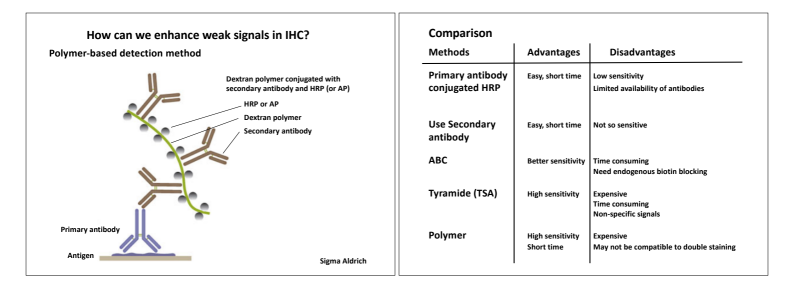
Mouse, rat, rabbit, goat, hamster, chicken, etc... Check reactivity of the antibody you use

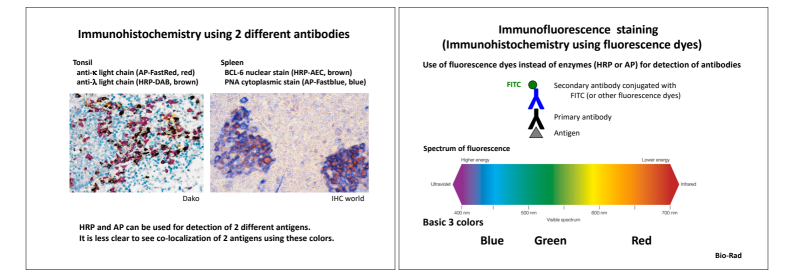
Enzo

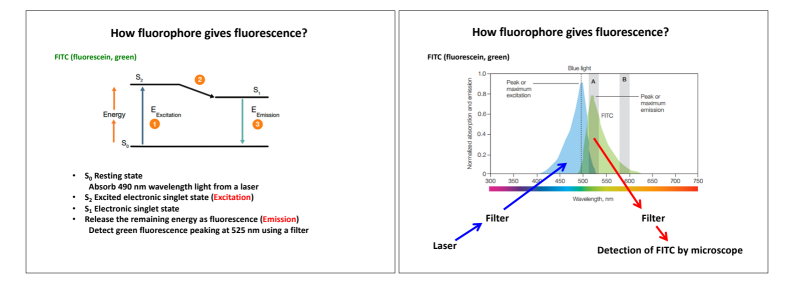


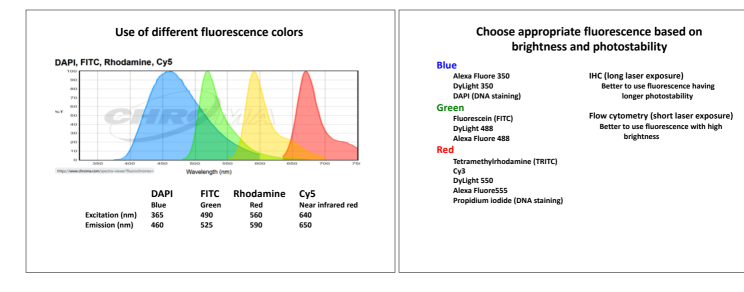






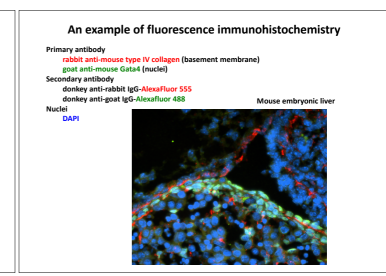


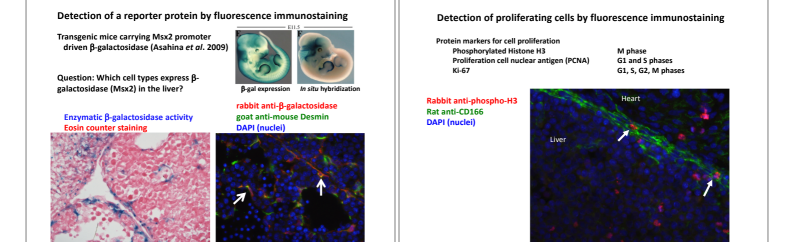


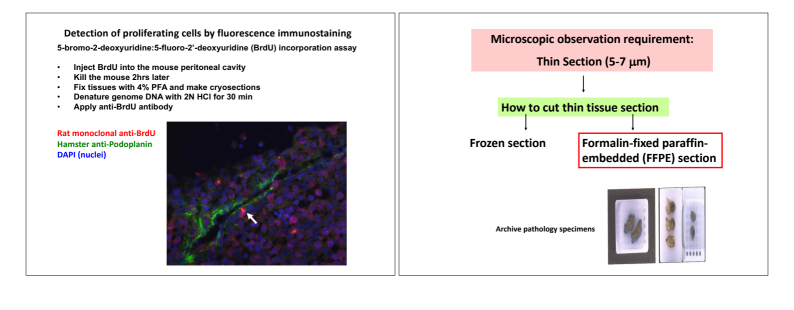


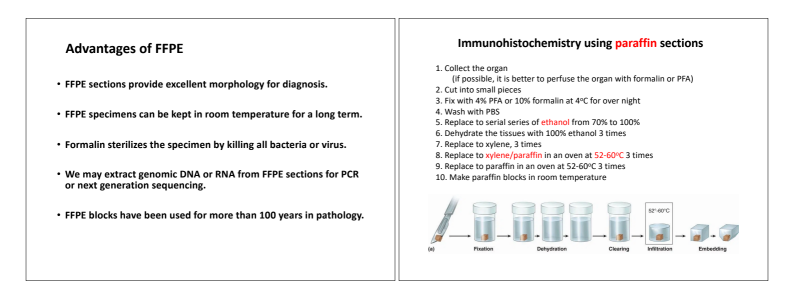
Procedure of fluorescence immunohistochemistry in frozen sections

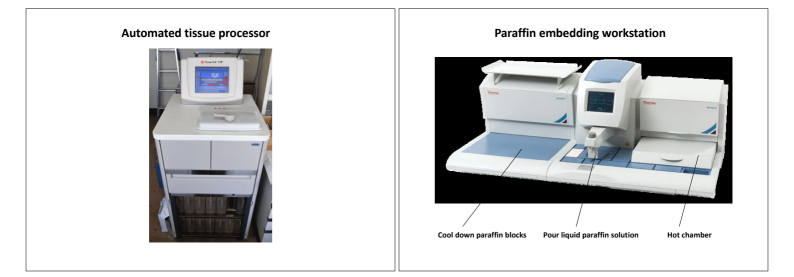
- 1. Make cryosections (7 µm)
- 2. Dry
- 3. Fix the sections if necessary
- 4. Remove frozen medium with PBS (no need to inactivate endogenous peroxidase or alkaline phosphatase)
- 5. Blocking the sections for 30 min (usually 5% goat or donkey serum and 0.2% bovine serum albumin in PBS)
- 6. Apply primary antibody for 1 hour room temperature or over night at $4^{\circ}C$
- 7. Wash with PBS
- 8. Apply secondary antibody conjugated with a fluorescent dye for 30 min room temperature
- 9. Wash with PBS
- 10. Counterstain nuclei with DAPI if necessary

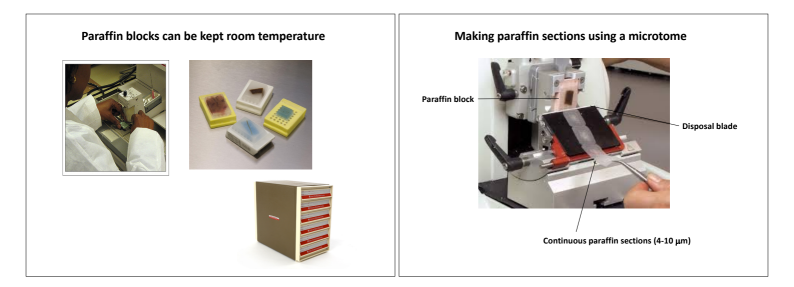


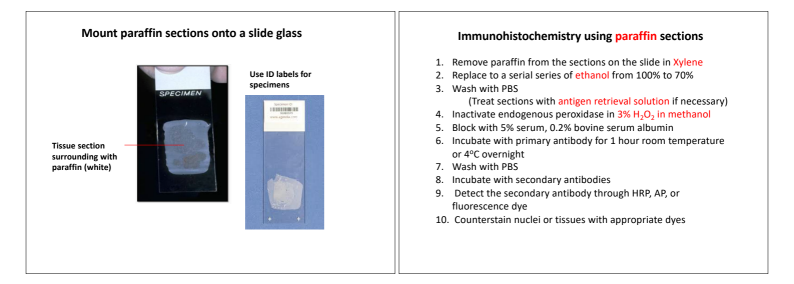


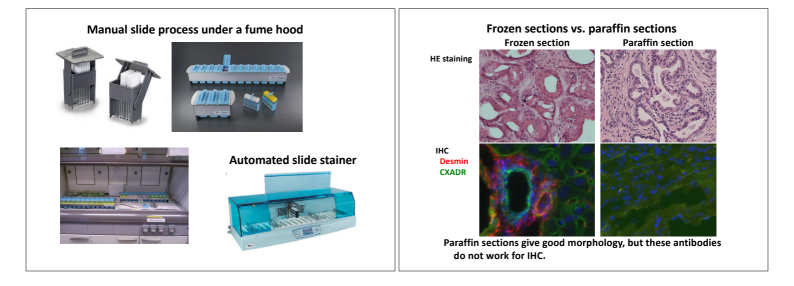












Why some antibodies do not work for paraffin sections?

Antigens in the section may lose antigenicity against antibodies

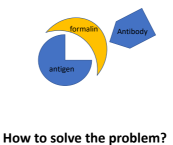
- 1. Proteins are denatured by heating during paraffin processing.
- 2. Proteins are denatured by organic solvents.
- 3. Cross-linking by formalin masks antigens.

There are several methods to overcome these issues.

- A. Use enzymes for partial digestion of sections.
- A. Use an antigen retrieval solution for breaking crosslink in sections.

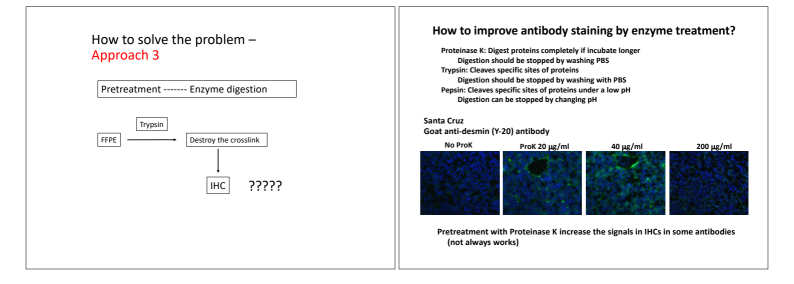
Drawback of FFPE

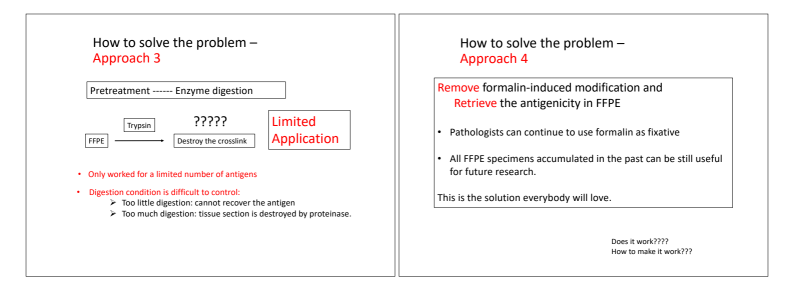
Formalin induces chemical modification to "mask" antigens, so most antigens on the tissue sections become inaccessible for immunohistochemistry (IHC).

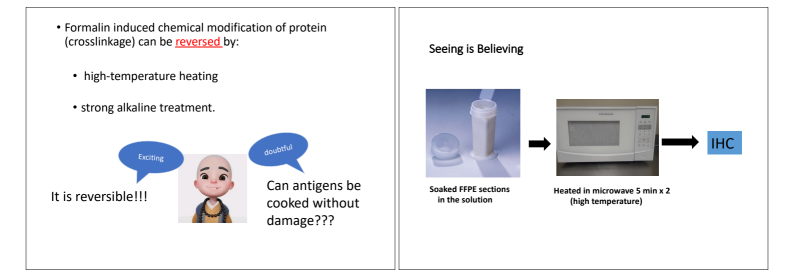


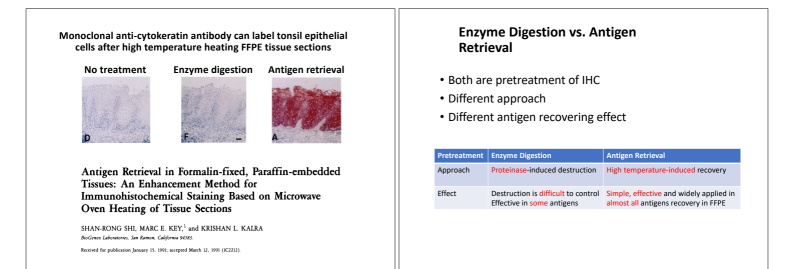


How to solve the problem-Approach 1 Replace formalin with non-formalin fixatives like ethanol, or acetone. Pathologists hate to use a new fixative. It changes the morphology and they cannot make the diagnosis. Researchers hate to use a new fixative. FFPE specimens in pathology accumulated in the past 100 years become useless.









Electron microscopy and immunohistochemistry Identification of coronavirus by immunoelectron microscopy Scanning electron microscopy Immunogold labeling Immunogold labeling Transmission electron microscopy Synaptophisin in the synaptic vesicles Dea and Garzon. J Vet Diagn Invest 1991 Immunoelectron microscopy 2010

Choice of an antibody for IHC

- On September 9th, 2021, Antibodypedia.com is referencing 4,480,670 antibodies from 95 providers against 19,109 gene products... It means about 234 antibodies are available by a gene product...
- Moser et al. (J Neurochem 2007;102:479-492) 13 working anti-nAChR antibodies were tested on nAChR-KO mice. Same immunoreactivity was observed in wildtype and KO mice. None of the antibodies was suitable for IHC!
- Egelhofer et al. (Nat. Struct. Mol. Biol. 2011;18:91-93.) 246 histone antibodies were tested: 25% were non-specific. 22% cannot precipitate chromatin. Some works but recognized the wrong target histone.

Control and assessment of the antibody specificity

All histochemical and immunohistochemical methods are prone to artefact and unspecific staining.

Control and assessment of the specificity.

For IHC, validation of the antibody is of critical importance.

About antibody validation

Magic peptides, magic antibodies: guidelines for appropriate controls for immunohistochemistry Saper CB, Sawchenko PE. J Comp Neurol 2003;465:161-163. PMID: 12949777.

An open letter to our readers on the use of antibodies. Saper CB. J Comp Neurol 2005;493:477-478. PMID: 16304632.

Antibodies as valuable neuroscience research tools versus reagents of mass distraction. Rhodes KJ, Trimmer JS. J Neurosci 2006;26:8017-8020. PMID: 16885215.

Antibody validation. Bordeaux J, Welsh A, Agarwal S, Killiam E, Baquero M, Hanna J, Anagnostou V, Rimm D. Biotechniques 2010;48:197-209. PMID: 20359301

Controls for immunohistochemistry: the Histochemical Society's standards of practice for validation of immunohistochemical assays. Hewitt SM, Baskin DG, Frevert CW, Stahl WL, Rosa-Molinar E. J Histochem Cytochem 2014;62:693-697. PMID: 25023613.

On the Necessity of Validating Antibodies in the Immunohistochemistry Literature. Gautron L Front Neuroanat 2019:13:46. PMID: 31080409

Sectioning service rendered by Central Research Laboratory

Paraffin sections

(Tissue embedded in paraffin block) Plastic sections

(Tissue embedded in plastic)

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