

# **Workshop in Diagnostic Immunohistochemistry**

## **Aalborg University Hospital, October 2022**

**Technical aspects of immunohistochemistry & pitfalls**  
**Pre-Analytical - Analytical (I & II) - Post Analytical phase**

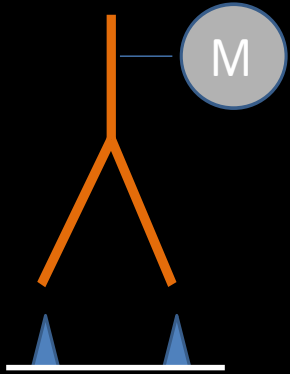
**Michael Bzorek**

**Histotechnologist**


**Department of Surgical Pathology**

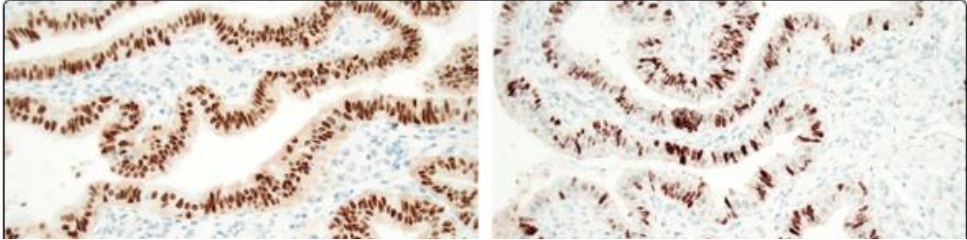
**University Hospital, Region Zealand, Denmark**


# Technical aspects of immunohistochemistry & pitfalls - Analytical phase



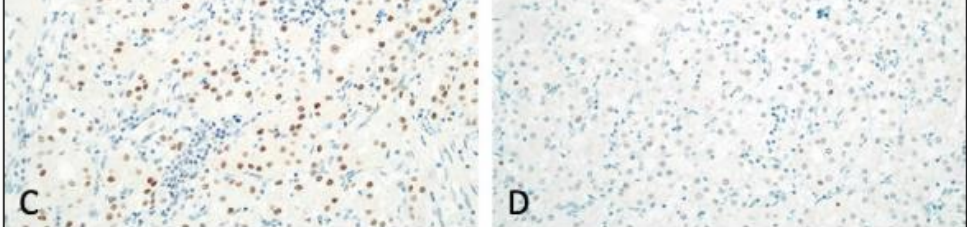
## Immunohistochemistry – A simple technique ?

[Info](#) [Modules](#) [Assessments](#) [Protocols](#) [Controls](#) [Events](#) [Login](#)



 **Events**  
[NordiQC Workshop in Diagnostic Immunohistochemistry](#)  
2-4 Oct 2019: Aalborg, DK  
[6th Academy of Diagnostic Immunohistochemistry](#)  
9-11 Oct 2019: Krakow, Poland


**External Quality Assurance programs**  
**Staining quality varies significantly between different laboratories depending on the individual selection of methods and the technical expertise**



**C**  
IHC for PAX8 in two laboratories: Lab 1 (A/C): Optimal results in Fallopian tube and Renal clear cell carcinoma. Lab 2 (B/D): Insufficient result in same tissues, especially characterized by false negative reaction in the neoplastic cells of the Renal clear cell carcinoma.

**D**

7 Sep 2019  
Slide circulation  
10 Sep 2019  
Slide return deadline  
11 Oct 2019  
Publication of results  
6 Dec 2019

 **Questions**  
Check out our [FAQ](#) (Frequently asked questions) or [contact us](#)

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

**The total test paradigm:  
Key elements in the IHC procedure**

## The analytic phase :

**Begins with dewax of the cut slides  
and is completed with the  
coverslipping of the stained slides.**

**Unlike the pre-analytic factors,  
analytic parameters can easily be  
modified and controlled within the  
immunohistological laboratory.**



### Pre-analytic phase

Pre-fixation  
Fixation  
Post-Fixation/Decalcification  
Processing  
Dehydration & clearing  
Paraffin embedding  
Sectioning  
Drying/Storage



### Analytic phase

Platform (manual/ Automated)  
Epitope retrieval  
Blocking  
Primary Antibody  
Detection system  
Chromogen  
Counterstain  
Mounting



### Post-analytic phase

Design of controls  
Critical stain indicators  
Internal/External control  
Interpretation  
Positive/Negative  
Localization  
Quantification  
Cutt-of levels  
Reporting



# Technical aspects of IHC and pitfalls – Analytical phase

## Optimization of the IHC assay – issues to be addressed

- Purpose and/or “fit-for-purpose” (assay validated for intended use)
- How to establish “best practice protocol” of the IHC test (Calibration of the IHC assay with focus clone, antigen retrieval, titer & detection system)
  - Is the IHC test reproducible/robust (pre-analytic conditions)
  - Evaluation of the analytical sensitivity and specificity
- Identification of most robust controls providing information that the established level of detection is obtained in each test performed in daily practice.

**Tissue materials are essential for all these processes (calibration, validation and controls)**



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

**Immunohistochemistry: Calibration of a biomarker/antibody depend on the type of marker and purpose of the test**

## IHC-type 1 markers (Diagnostic)

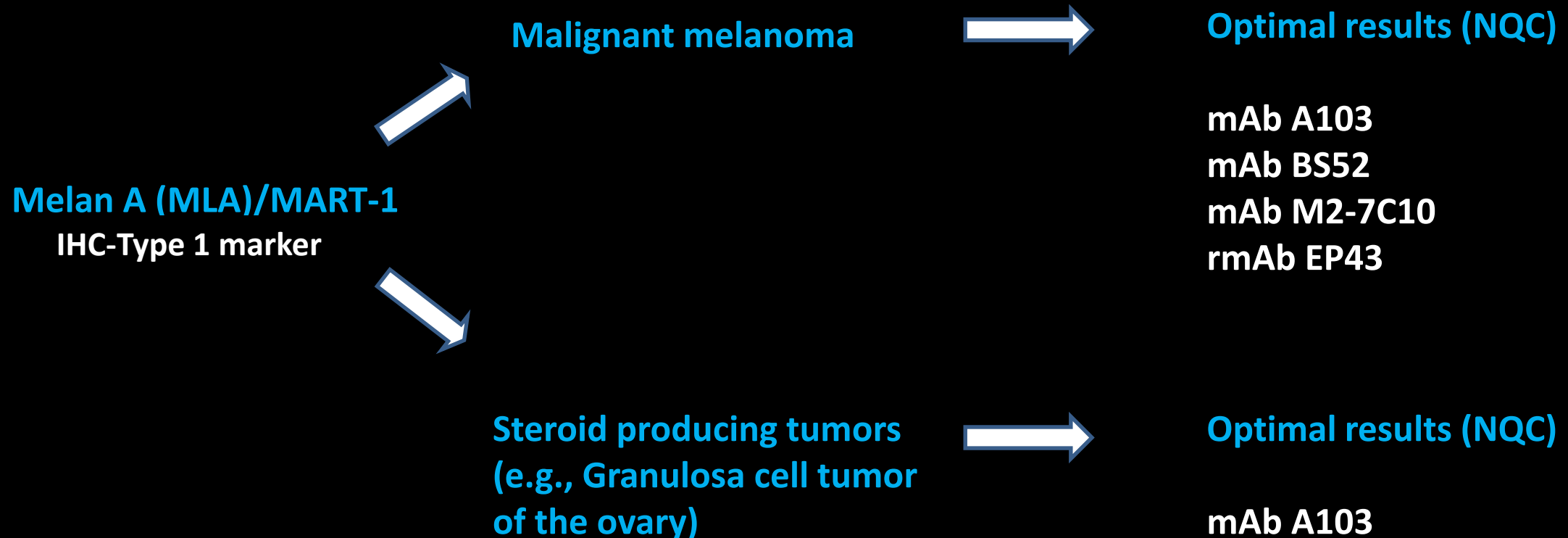
**Often calibrated to produced appropriate level of sensitivity and specificity (positive versus negative)**

## IHC-type 2 markers (Disease screening, predictive treatment & prognosis)

**Predictive markers: The assays are calibrated to provide information of which patients will or will not benefit from a specific treatment (HER-2, PD-L1 .....)**

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Purpose and Intended use



mAb clone A103, cross-react with an unknown protein in steroid hormone producing cells and tumours derived from these cells.

Table 1. Antibodies and assessment marks for MLA, Run 56

Concentrated antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff. <sup>1</sup>	Suff. OPS <sup>2</sup>
mAb clone <b>A103</b>	69 19 5 1 1 1	Dako/Agilent Novocastra/Leica Cell Marque Diagnostic BioSystems Immunologic Monosan Thermo Scientific	22	43	24	8	67%	73%
mAb clone <b>BS52</b>	1	Nordic Biosite	1	0	0	0	-	-
mAb clone cocktail <b>HMB45+MC-7310 +M2-9E3+T311</b>	3	Biocare	1	2	0	0	-	-
mAb clone cocktail <b>M2-7C10+M2-9E3</b>	1	NeoMarkers	0	1	0	0	-	-
mAb clone <b>M2-7C10</b>	1	Zytomed	1	0	0	0	-	-
rmAb <b>EP43</b>	5 3 3	Nordic Biotite Cell Marque Epitomics	9	1	1	0	91%	100%
Ready-To-Use antibodies								
mAb clone <b>A103 790-2990</b>	87	Ventana/Roche	8	31	45	3	45%	66%
mAb clone <b>A103, IR633/IS633</b>	26	Dako/Agilent	10	14	2	0	92%	95%
mAb clone <b>A103, IR633/IS633<sup>3</sup></b>	29	Dako/Agilent	9	12	8	0	72%	-
mAb clone <b>A103, IR633/IS633<sup>4</sup></b>	9	Dako/Agilent	1	2	5	1	-	-
mAb clone <b>A103, PA0233</b>	9	Novocastra/Leica	1	8	0	0	-	-
mAb clone <b>A103, PA0233<sup>5</sup></b>	1	Novocastra/Leica	0	0	1	0	-	-
mAb clone <b>A103, 281M-87/281M-88</b>	3	Cell Marque	1	0	2	0	-	-
mAb clone <b>A103, API3114</b>	1	Biocare	0	1	0	0	-	-
mAb clone <b>A103, MAB-0275</b>	1	Maixin	1	0	0	0	-	-
mAb clone cocktail <b>HMB45+A103+T311, 904H-08</b>	1	CellMarque	1	0	0	0	-	-
mAb clone cocktail <b>HMB45+A103+T311, 790-4677</b>	1	Ventana/Roche	0	1	0	0	-	-
rmAb clone <b>EP43, MAD-000695QD-7/N</b>	2	Master Diagnostica	0	1	1	0	-	-
rmAb clone <b>EP43, 8319-C010</b>	2	Sakura Finetek	1	1	0	0	-	-
Total	286		67	118	89	12	-	
Proportion			24%	41%	31%	4%	65%	

1) Proportion of sufficient stains (optimal or good)

2) Proportion of sufficient stains with optimal protocol settings only, see below.

3) RTU system developed for the Dako/Agilent semi-automatic system (Dako Autostainer), but used by laboratories on the Dako fully-automatic platform (Dako Omnis)

4) RTU system developed for the Dako/Agilent semi-automatic system (Dako Autostainer), but used by laboratories on different platforms (e.g. Ventana BenchMark)

5) RTU system developed for the Leica Bond system, but used on the Ventana BenchMark system.

## Purpose and Intended use

### Material

The slide to be stained for MLA comprised:

1. Skin, 2. Kidney, 3. Adrenal gland, 4-5. Malignant melanoma, 6. Granulosa cell tumour.

All tissues were fixed in 10% neutral buffered formalin.



## Melan A/MART-1 (Run 56)

MLA, A103 (melanoma/melanocyte assessment): 62% Sufficient/ 20% Optimal

Optimal results: Efficient HIER (alkaline buffer), high conc. of the primary Ab (app. 1:10-50) and a sensitive detection system (3-step)

Melanoma/melanocyte + steroid hormone assessment): Suff. 29%

Sufficient steroid hormone related A103 (cross)reaction can be very difficult to obtain on the Leica Bond, Ventana BenchMark and Dako Omnis platforms.

## Melan A (MLA) / MART-1:

263 participants ~ 92% used clone A103 as single reagent (no cocktails)

Table 1. Antibodies and assessment marks for MLA, Run 60

Concentrated antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff. <sup>1</sup>	OR <sup>2</sup>
mAb clone <b>A103</b>	57 19 6 1 1 1 1 1 1	Dako/Agilent Novocastra/Leica Cell Marque Abcam Biocare Monosan Biogenex Zeta Corporation	21	57	9	0	90%	24%
mAb clone <b>BS52</b>	3	Nordic Biosite	3	0	0	0	-	-
mAb clone <b>M2-7C10</b>	1	Zytomed	1	0	0	0	-	-
mAb clone cocktail <b>HMB45+MC-7310 +M2-9E3+T311</b>	3	Biocare	2	1	0	0	-	-
mAb clone cocktail <b>HMB45+A103+T311</b>	1	Diagnostic Biosystems	0	0	0	1	-	-
mAb clone cocktail <b>M2-7C10+M2-9E3</b>	1	Thermo F. Scientific	1	0	0	0	-	-
rmAb clone <b>EP1442Y</b>	1	Abcam	1	0	0	0	-	-
rmAb clone <b>EP43</b>	9 9 1	Nordic Biotite Epitomics Cell Marque	18	1	0	0	100%	95%
Ready-To-Use antibodies								
mAb clone <b>A103 790-2990<sup>3</sup></b>	3	Ventana/Roche	0	0	3	0	-	-
mAb clone <b>A103 790-2990<sup>4</sup></b>	94	Ventana/Roche	6	74	11	3	85%	6%
mAb clone <b>A103, IR633/IS633<sup>3</sup></b>	14	Dako/Agilent	1	13	0	0	100%	7%
mAb clone <b>A103, IR633/IS633<sup>4</sup></b>	56	Dako/Agilent	12	36	7	1	85%	21%
mAb clone <b>A103, PA0233/PA0044<sup>3</sup></b>	7	Leica Biosystems	2	5	0	0	100%	29%
mAb clone <b>A103, PA0233/PA0044<sup>4</sup></b>	10	Leica Biosystems	6	4	0	0	100%	60%
mAb clone <b>A103, 281M-87/281M-88</b>	1	Cell Marque	1	0	0	0	-	-
mAb clone <b>A103, API3114</b>	1	Biocare	0	0	1	0	-	-
mAb clone <b>M2-7C10, 281M-97/281M-98</b>	2	Cell Marque	1	1	0	0	-	-
mAb clone cocktail <b>HMB45+A103+T311, 904H-08</b>	1	Cell Marque	0	0	1	0	-	-
mAb clone cocktail <b>HMB45+A103+T311, 790-4677</b>	1	Ventana/Roche	0	1	0	0	-	-
mAb clone cocktail <b>HMB45+MC-7310 +M2-9E3+T311, PM165</b>	1	Biocare	0	1	0	0	-	-
rmAb clone <b>BP6086, I1064</b>	1	Tuling Biotechnology	0	0	1	0	-	-
rmAb clone <b>EP43, MAD-000695QD-7/N</b>	1	Master Diagnostica	1	0	0	0	-	-
rmAb clone <b>EP43, 8319-C010</b>	3	Sakura Finetek	3	0	0	0	-	-
Total	312		80	194	33	5	-	
Proportion			26%	62%	11%	1%	88%	

1) Proportion of sufficient results (optimal or good). (≥5 assessed protocols).

2) Proportion of Optimal Results (OR).

3) Vendor Recommended Protocol Settings (VRPS) to a specific RTU product applied on the vendor recommended platform(s) (≥5 assessed protocols).

4) Laboratory Modified Protocol Settings (LMPS) to a specific RTU product applied either on the vendor recommended platform(s) or on validated semi/fully automatic systems or used manually (≥5 assessed protocols).

## Purpose and Intended use

### Material

The slide to be stained for MLA comprised:

1. Kidney, 2. Skin, 3-4. Malignant melanoma, 5. Colon Adenocarcinoma.

All tissues were fixed in 10% neutral buffered formalin.



## Melan A/MART-1 (Run 60)

MLA, A103 (melanoma/melanocyte assessment): 87% Sufficient / 18% Optimal.

Optimal results: Efficient HIER (alkaline buffer), high conc. of the primary Ab (app. 1:10-50) and a sensitive detection system (3-step).

Performance of MLA clone A103 is sensitive to the chosen platform e.g. Omnis or Benchmark.

## Melan A (MLA) / MART-1:

273 participants ~ 88% used clone A103 as single reagent (no cocktails)

Is MLA , A103 the best primary Ab for detection of melanomas ?



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

MLA Run 56, Mab clone A103

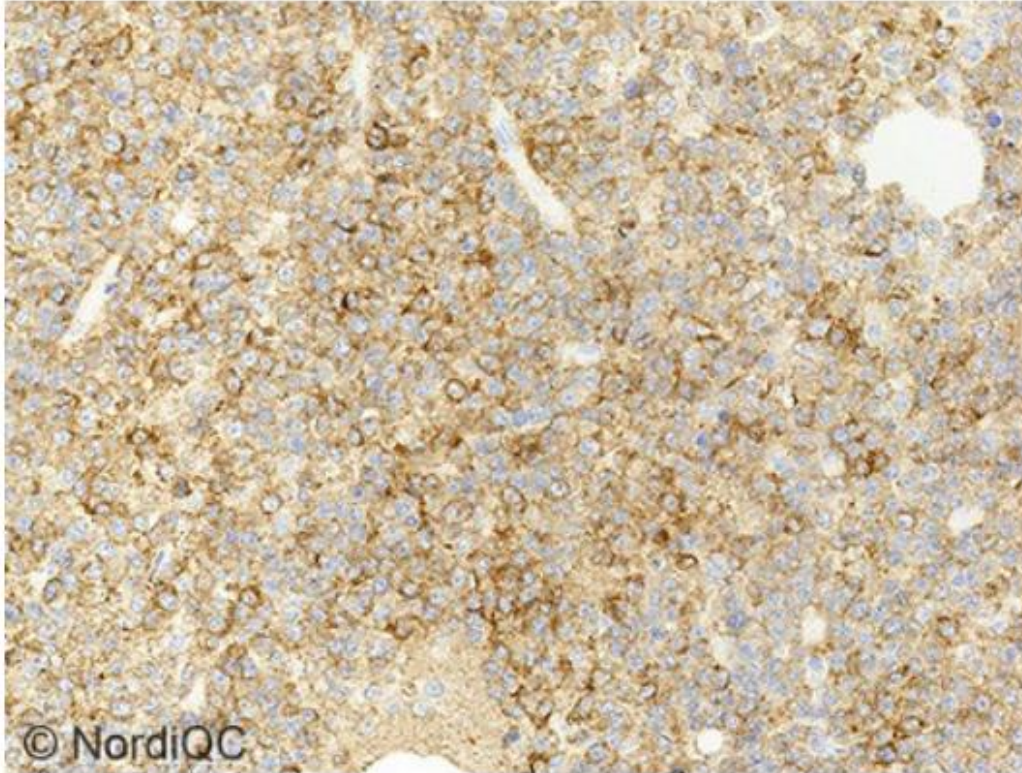


Fig. 8a  
Optimal MLA staining of the malignant melanoma tissue core no. 4 (low-level expressor) using same protocol as in Figs. 7a and 7b. The majority of neoplastic cells show a weak to moderate cytoplasmic staining reaction. Compare with Figs. 7a and 8b, same protocol. Also compare with Figs. 5a and 5b - same field.

MLA Run 56, Rab clone EP43

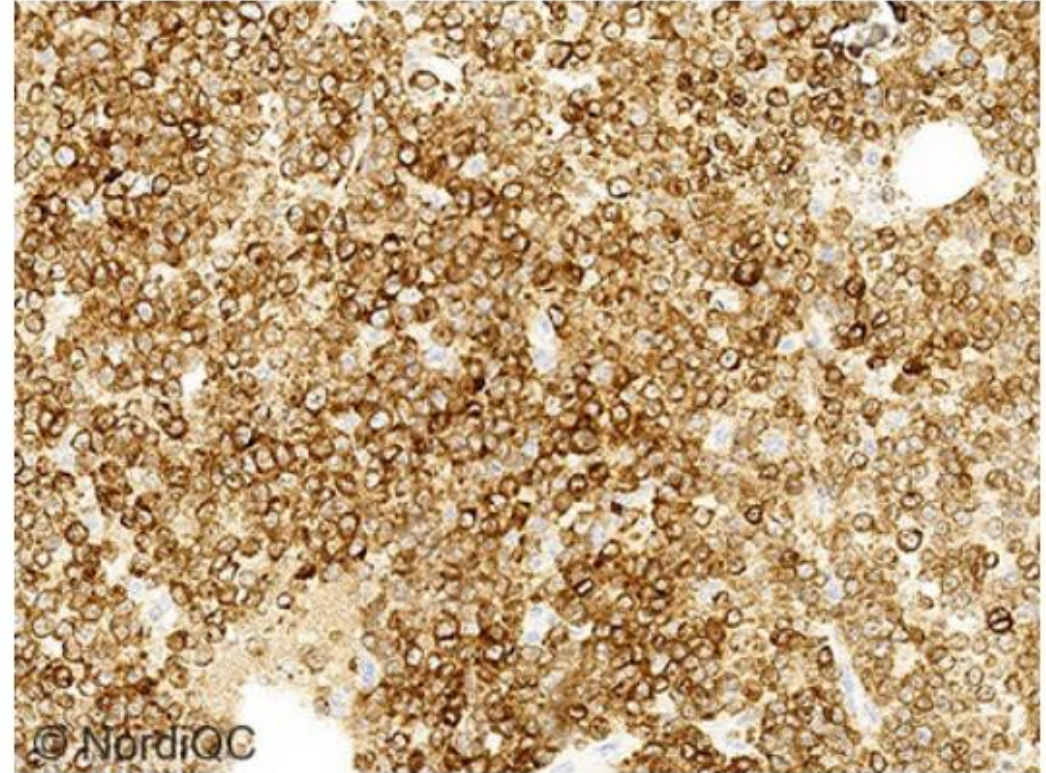


Fig. 8b  
Optimal MLA staining of the malignant melanoma tissue core no. 4 (low-level expressor) using **rmAb EP43** in similar protocol settings as in Fig. 8a (and 7a and 7b) All neoplastic cells show a moderate to strong cytoplasmic staining reaction. In melanomas, rmAb EP43 produce very strong reactions. Compare with mAb A103 in Fig. 8a - same field.



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Melan A /MART1

Melanoma (Sentinel node)

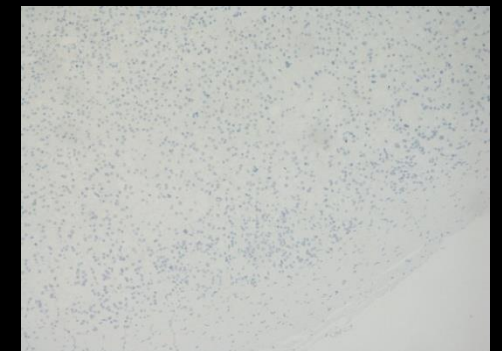
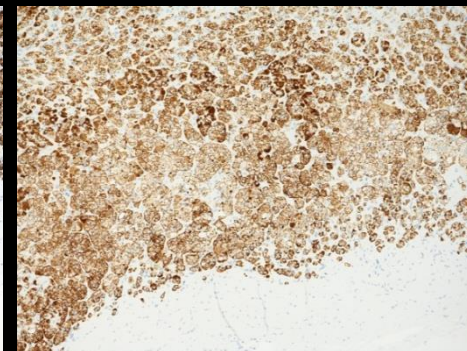
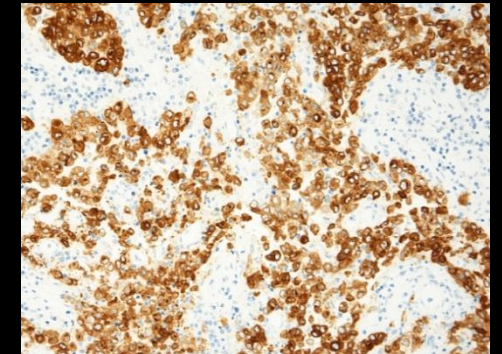
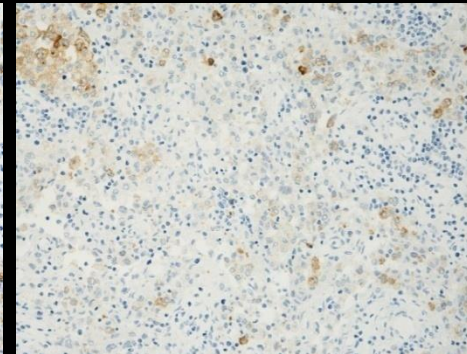
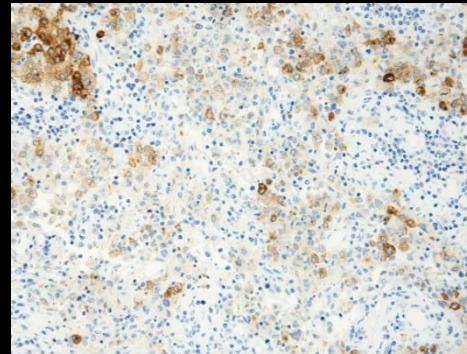
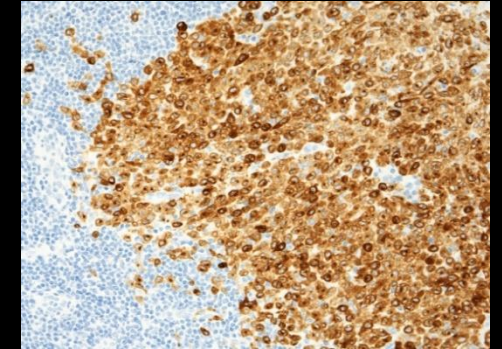
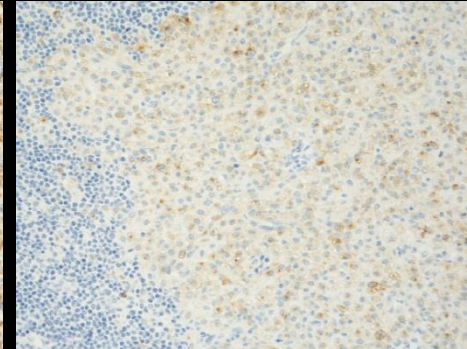
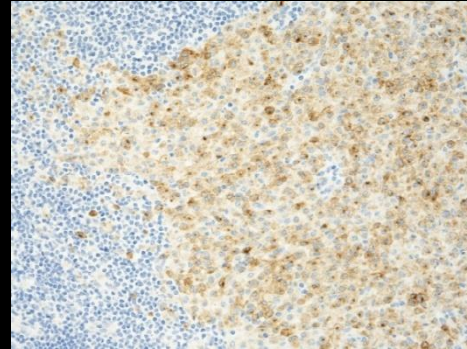
Melanoma (Lymph node)

Adrenal gland

MLA, A103 1:25  
AutoStainer

MLA, A103 1:25  
Omnis

MART1, EP43 1:30  
Omnis



TRS High pH/Flex+ protocols

# Technical aspects of IHC and pitfalls – Analytical phase

## Optimization of the IHC assay – issues to be addressed

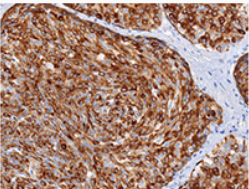
- Purpose and/or “fit-for-purpose” ( assay validated for intended use)
- How to establish “best practice protocol” of the IHC test (Calibration of the IHC assay with focus clone, antigen retrieval, titer & detection system)
  - Is the IHC test reproducible/robust (pre-analytic conditions)
  - Evaluation of the analytical sensitivity and specificity
- Identification of most robust controls providing information that the established level of detection is obtained in each test performed in daily practice.

**Tissue materials are essential for all these processes (calibration, validation and controls)**



# Calcitonin optimization (data sheets ?)

## anti-Calcitonin (SP17), Rabbit Monoclonal Primary Antibody



**Catalog Number:** 760-4705  
**Ordering Code:** 06586554001  
**Quantity:** 50 tests  
**Controls:** Medullary Carcinoma of Thyroid  
**Isotypes:** IgG  
**Clone Name:** SP17  
**Species:** Rabbit  
**Localization:** Cytoplasmic  
**Regulatory Status:** IVD

Roche/ Ventana/ Cell Marque

This antibody is intended for in vitro diagnostic (IVD) use. Calcitonin (SP17) antibody is intended for qualified laboratories to qualitatively identify by light microscopy the presence of associated antigens in sections of formalin-fixed, paraffin-embedded tissues using standard test methods. This antibody is used as an aid in the diagnosis of thyroid medullary carcinoma within the clinical history, and other diagnostic tests determined.

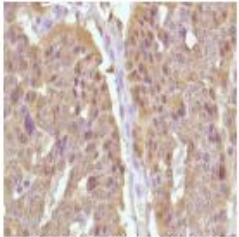
Recommended staining protocol with ultraView	
Procedure Type	Method
Deparaffinization	Selected
Cell Conditioning (Antigen Unmasking)	Cell Conditioning 1, Mild
Enzyme (Protease)	Not required
Antibody (Primary)	BenchMark ULTRA instrument: 16 minutes, 36°C
	BenchMark XT instrument: 16 minutes, 37°C
	BenchMark GX instrument: 16 minutes, 37°C

Optimizing an assay can be confusing  
(Vendor recommendations)



## Rabbit Anti-Human Calcitonin Monoclonal Antibody (Clone SP17)

Spring Cat#	Roche P/N	Product Description
M3170	05492769001	0.1 ml rabbit monoclonal antibody supplied as tissue culture supernatant in TBS/1% BSA buffer pH 7.5 with less than 0.1% sodium azide.
M3172	06970419001	0.5 ml rabbit monoclonal antibody supplied as tissue culture supernatant in TBS/1% BSA buffer pH 7.5 with less than 0.1% sodium azide.
M3174	05298725001	1.0 ml rabbit monoclonal antibody supplied as tissue culture supernatant in TBS/1% BSA buffer pH 7.5 with less than 0.1% sodium azide.
M3171	05298717001	7.0 ml pre-diluted rabbit monoclonal antibody supplied as tissue culture supernatant in TBS/1% BSA buffer pH 7.6 with less than 0.1% sodium azide. (For manual IHC only)



Human thyroid medullary carcinoma stained with anti-calcitonin antibody

<b>INTENDED USE:</b>	For Research Use Only. Not for use in diagnostic procedures.
<b>CLONE:</b>	SP17
<b>IMMUNOGEN:</b>	Synthetic human calcitonin 1-32 amino acid peptide.
<b>IG ISOTYPE:</b>	Rabbit IgG
<b>EPITOPE:</b>	Not determined
<b>MOLECULAR WEIGHT:</b>	15kDa
<b>SPECIES REACTIVITY:</b>	Human (tested). (See <a href="http://www.springbio.com">www.springbio.com</a> for information on species reactivity predicted by sequence homology.)
<b>DESCRIPTION:</b>	Calcitonin is a 32 amino acid peptide which can be demonstrated in C cells of the normal and hyperplastic thyroid. Staining for calcitonin may be used for the identification of a spectrum of C cell proliferative abnormalities ranging from C cell hyperplasia to invasive tumors. Staining for calcitonin in medullary carcinoma of the thyroid produces a fine granular pattern in the cytoplasm. Amyloid deposits within the tumor may also exhibit varying degrees of calcitonin activity.
<b>APPLICATIONS:</b>	Immunohistochemistry (IHC)
<b>IHC PROCEDURE:</b>	<b>Specimen Preparation:</b> Formalin-fixed, paraffin-embedded tissues are suitable for use with this primary antibody. <b>Deparaffinization:</b> Deparaffinize slides using xylene or xylene alternative and graded alcohols. <b>Antibody Dilution:</b> If using the concentrate format of this product, dilute the antibody 1:100 in Antibody Diluent (Cat# ADS-125). The dilutions are estimates; actual results may differ because of variability in methods and protocols. <b>Antigen Retrieval:</b> None <b>Primary Antibody Incubation:</b> Incubate for 30 minutes at room temperature. <b>Slide Washing:</b> Slides must be washed in between steps. Rinse slides with PBS/0.05% Tween. <b>Detection:</b> Detect the antibody as instructed by the instructions provided with the detection system.
<b>POSITIVE CONTROL:</b>	Thyroid medullary carcinoma
<b>CELLULAR LOCALIZATION:</b>	Cytoplasm

Can we use the recommendations provided by the manufactures spec sheets?



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Optimizing IHC test (IHC-type 1) - parameters to consider

Use a “Test battery approach” (pre-treatment and dilution range)

Test more than one antibody clone against an antigen of interest before implementation in the routine

Test with robust, specific & sensitive detection system

Test/validate on normal and tumor tissue material with broad spectrum of antigen densities (specificity/sensitivity)

Compare results with external quality assurance programs, literature or colleagues

No antibody should be acquired without the basic knowledge of its performance characteristics and expected expression pattern

Hadi Yaziji and Todd Barry – Adv Anat Pathol • Vol. 13, Number 5, September 2006

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Laboratory Developed Assays : Dept. of Surgical Pathology, Region Zealand, Denmark  
Omnis + Autostainer platforms

## Antibody Performance Testing (“Test Battery approach”)

	Dil. 1	Dil.2	Dil.3
A	None	None	None
B	Enzyme (1) 5 min.	Enzyme (1) 5 min.	Enzyme (1) 5 min.
C	HIER TRS Low pH 6.1 (30`)	HIER TRS Low pH 6.1 (30`)	HIER TRS Low pH 6.1 (30`)
D	HIER TRS High pH 9.0 (24`)	HIER TRS High pH 9.0 (24`)	HIER TRS High pH 9.0 (24`)
<hr/>			
E	TRS Low (20`) + Pep (12`)	TRS Low (20`) + Pep (12`)	TRS Low (20`) + Pep (12`)
F	HIER TRS High pH 9.0 (30`)	HIER TRS High pH 9.0 (30`)	HIER TRS High pH 9.0 (30`)
G	Pep 6 & 10 min + TRS High *	Pep 6 & 10 min + TRS High	Pep 6 & 10 min + TRS High
H	Pepsin 20 min.	Pepsin 20 min	Pepsin 20 min

## Omnis

Protocol A: 0.5 %  
Protocol B: 2.0 %  
Protocol C: 10.0 %  
Protocol D: 83.5 %

Protocol E: 1.0 %  
Protocol F: 3.0 %

Protocol G: 0 %  
Protocol H: 0 %

\* Off board enzymatic pre-treatment

Identify the protocol that discriminate between the desired (specific) positive staining and any unwanted (non-specific) background staining

# Test “battery” approach (BMU/Ventana)

## IHC – Biomarker controls



### Concentrated antibodies – VMS ULTRA

	1:25	1:100	1:400
A	None	None	None
B	Enzyme P1, 4 min	Enzyme P1, 4 min	Enzyme P1, 4 min
C	HIER CC1 pH 8.5*	HIER CC1 pH 8.5	HIER CC1 pH 8.5
D	HIER CC2 pH 6.0**	HIER CC2 pH 6.0	HIER CC2 pH 6.0
<hr/>			
(E)	CC1 + Enzyme P3, 8 min	CC1 + Enzyme P3, 8 min	CC1 + Enzyme P3, 8 min
(F)	Enzyme P3, 8 min + CC1	Enzyme P3, 8 min + CC1	Enzyme P3, 8 min + CC1

\*HIER time 48 min. at 99°C, \*\* HIER time 32 min. at 99°C  
32 min in primary Ab, OptiView DAB, Ventana BenchMark Ultra

Protocol A: 2 %

Protocol B: 3 %

Protocol C: 90 %

Protocol E: 3 %

Protocol F: 1 %

Others : 2 % (E.g. prolonged HIER, prolonged proteolysis, amp. Kit....)

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

TMA Normal Tissue			
Tonsil 6h	Tonsil 24-72h	Tonsil 96-168h	Liver
Colon 6h	Colon 24-72h	Colon 96-168	Placenta
Breast	Pancreas	Salivary Gl.	Skin
Kidney	Adrenal Gl.	Prostate	Brain
Cervix	Testis	Skeletal Muscle	Colon/Tonsil 24h decalc. EDTA

Protocol set-up: Evaluate analytic sensitivity and specificity

Normal tissue including fixation and decalcification controls

Identification of the best practice protocol (clone, titer, retrieval etc.)

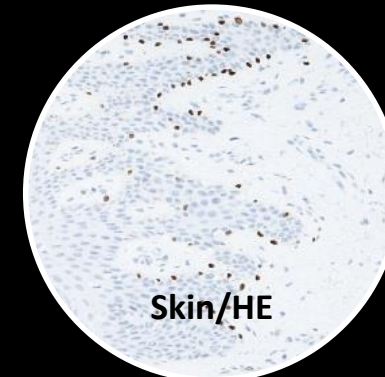
**SOX10, BS7**; HIER High pH 24` ; 1:350 RR; Flex+Mouse linker

Establishing robustness of the IHC assay / pre-analytic parameter's ?

**SOX10, BS7**; Robust to both fixation time in NBF and decalcification

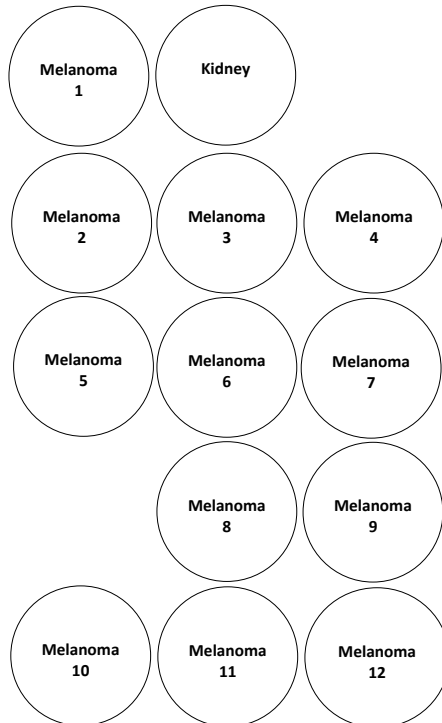
Identification of robust controls

**SOX10, BS7**; High, Low & Non-expressors ?

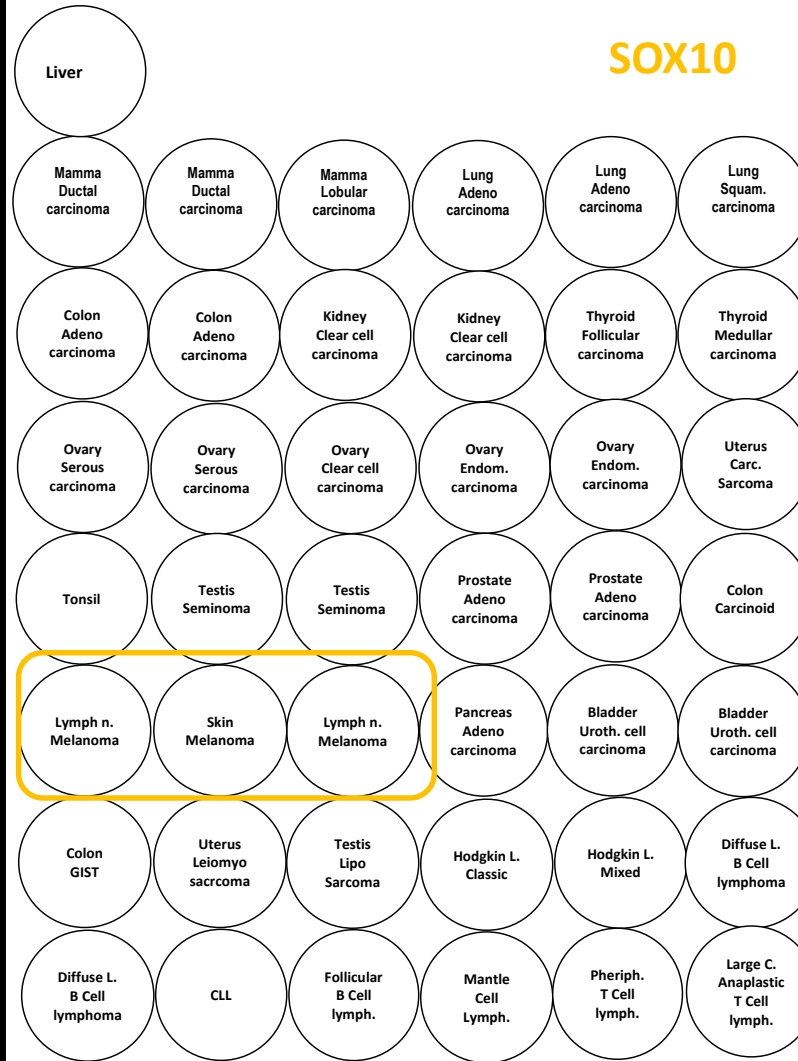




**TMA**  
Malignant Melanomas



**TMA**  
Mixed tumors



## Diagnostic potential

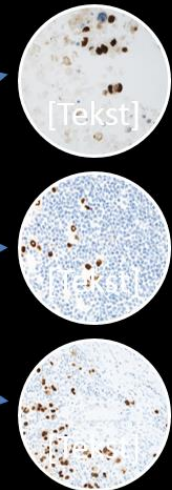
## Analytical validation



**SOX10, BS7:**

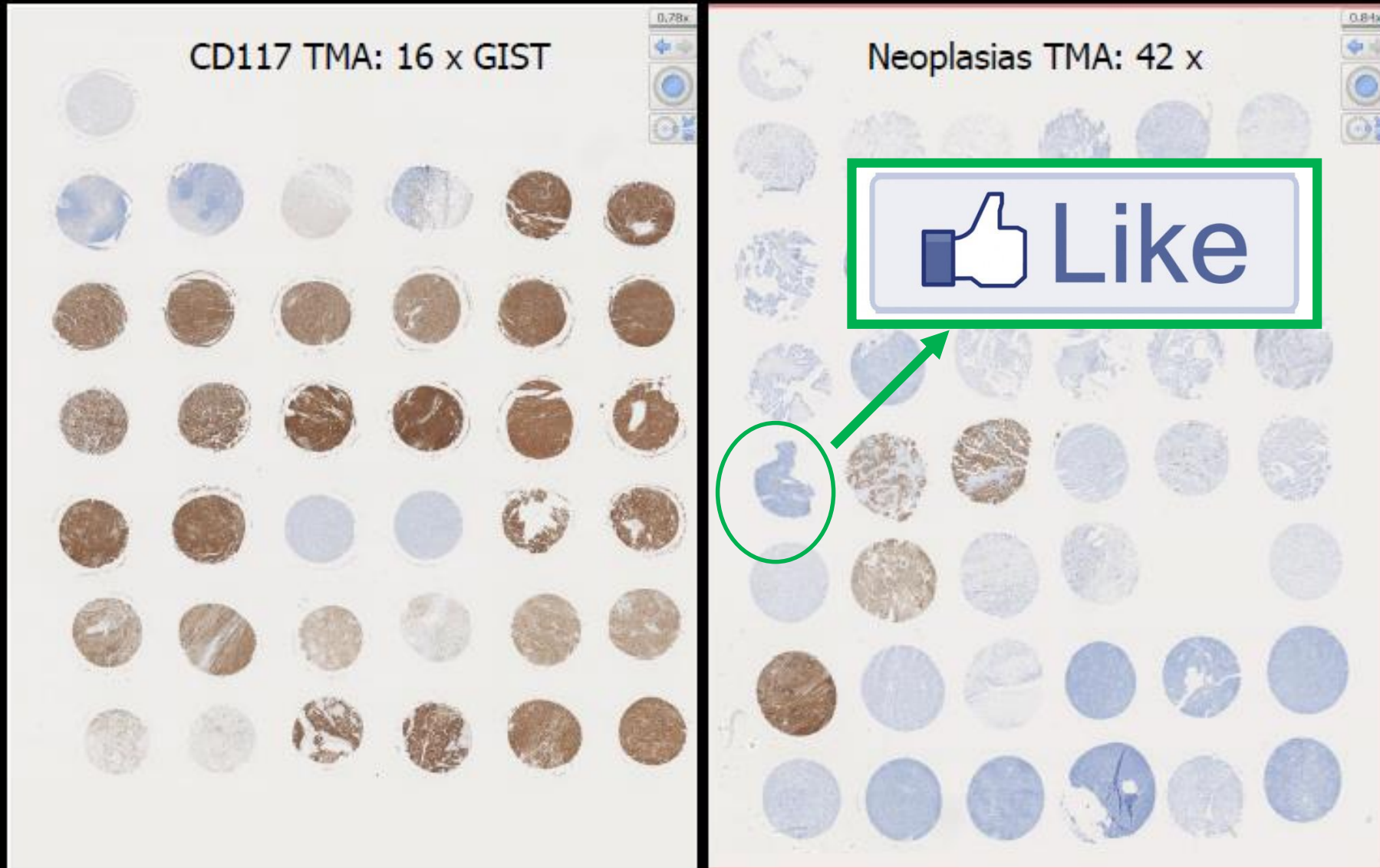
15/15 Melanomas were positive

37/37 other neoplasm's were negative



Other SOX10+ tumours: Schwannoma's, neurofibroma's, myoepithelial carcinomas, triple negative breast cancer.....?

# IHC – The Technical Test Approach





## Evolution of Quality Assurance for Clinical Immunohistochemistry in the Era of Precision Medicine: Part 1: Fit-for-Purpose Approach to Classification of Clinical Immunohistochemistry Biomarkers

*Carol C. Cheung, MD, PhD, JD,\*† Corrado D'Arrigo, MB, ChB, PhD, FRCPath,‡§||  
Manfred Dietel, MD, PhD,¶ Glenn D. Francis, MBBS, FRCPA, MBA, FFSc (RCPA),##\*\*††  
C. Blake Gilks, MD,‡‡ Jacqueline A. Hall, PhD,§§|| Jason L. Hornick, MD, PhD,¶¶  
Merdol Ibrahim, PhD,### Antonio Marchetti, MD, PhD,\*\*\* Keith Miller, FIBMS,##  
J. Han van Krieken, MD, PhD,††† Soren Nielsen, BMS,‡‡‡§§§ Paul E. Swanson, MD,||||  
Clive R. Taylor, MD,¶¶¶ Mogens Vyberg, MD,‡‡‡§§§ Xiaoge Zhou, MD,####\*\*\*\*  
and Emina E. Torlakovic, MD, PhD,\*††††††††*

*From the International Society for Immunohistochemistry and Molecular Morphology (ISIMM)  
and International Quality Network for Pathology (IQN Path)*

original purpose for which an IHC test is developed and its

## Evolution of Quality Assurance for Clinical Immunohistochemistry in the Era of Precision Medicine – Part 2: Immunohistochemistry Test Performance Characteristics

*Emina E. Torlakovic, MD, PhD,\*†‡ Carol C. Cheung, MD, PhD, JD,\*§  
Corrado D'Arrigo, MB, ChB, PhD, FRCPath,||¶## Manfred Dietel, MD, PhD,\*\*  
Glenn D. Francis, MBBS, FRCPA, MBA, FFSc (RCPA),††‡‡§§ C. Blake Gilks, MD,||  
Jacqueline A. Hall, PhD,¶¶ Jason L. Hornick, MD, PhD,### Merdol Ibrahim, PhD,\*\*\*  
Antonio Marchetti, MD, PhD,††† Keith Miller, FIBMS,\*\*\* J. Han van Krieken, MD, PhD,†††  
Soren Nielsen, BMS,§§§|||| Paul E. Swanson, MD,¶¶¶ Mogens Vyberg, MD,§§§||||  
Xiaoge Zhou, MD,####\*\*\*\* Clive R. Taylor, MD,†††† and  
From the International Society for Immunohistochemistry and Molecular Morphology (ISIMM)  
and International Quality Network for Pathology (IQN Path)*

## Evolution of Quality Assurance for Clinical Immunohistochemistry in the Era of Precision Medicine. Part 3: Technical Validation of Immunohistochemistry (IHC) Assays in Clinical IHC Laboratories

*Emina E. Torlakovic, MD, PhD,\*†‡ Carol C. Cheung, MD, PhD, JD,\*§ Corrado D'Arrigo, MB,  
ChB, PhD, FRCPath,||¶## Manfred Dietel, MD, PhD,\*\* Glenn D. Francis, MBBS, FRCPA, MBA,  
FFSc (RCPA),††‡‡§§ C. Blake Gilks, MD,|| Jacqueline A. Hall, PhD,¶¶  
Jason L. Hornick, MD, PhD,### Merdol Ibrahim, PhD,\*\*\* Antonio Marchetti, MD, PhD,†††  
Keith Miller, FIBMS,\*\*\* J. Han van Krieken, MD, PhD,‡‡‡ Soren Nielsen, BMS,§§§||||  
Paul E. Swanson, MD,¶¶¶ Mogens Vyberg, MD,§§§|||| Xiaoge Zhou, MD,####\*\*\*\*  
and Clive R. Taylor, MD,††††*

*From the International Society for Immunohistochemistry and Molecular Morphology (ISIMM)  
and International Quality Network for Pathology (IQN Path)*

## Evolution of Quality Assurance for Clinical Immunohistochemistry in the Era of Precision Medicine: Part 4: Tissue Tools for Quality Assurance in Immunohistochemistry

*Carol C. Cheung, MD, PhD, JD,\*† Corrado D'Arrigo, MB, ChB, PhD, FRCPath,‡§||  
Manfred Dietel, MD, PhD,¶ Glenn D. Francis, MBBS, FRCPA, MBA, FFSc (RCPA),##\*\*††  
Regan Fulton, MD, PhD,‡‡ C. Blake Gilks, MD,§§ Jacqueline A. Hall, PhD,||¶¶  
Jason L. Hornick, MD, PhD,### Merdol Ibrahim, PhD,\*\*\* Antonio Marchetti, MD, PhD,†††††  
Keith Miller, FIBMS,\*\*\* J. Han van Krieken, MD, PhD,§§§ Soren Nielsen, BMS,||||¶¶¶  
Paul E. Swanson, MD,#### Clive R. Taylor, MD,\*\*\*\* Mogens Vyberg, MD,||||¶¶¶  
Xiaoge Zhou, MD,†††††††† and Emina E. Torlakovic, MD, PhD,\*§§§§||||||  
From the International Society for Immunohistochemistry and Molecular Morphology (ISIMM)  
and International Quality Network for Pathology (IQN Path)*

Article sequence (part 1-4) published in Appl Immunohistochem Mod Morphol (2017) systematically describing/defining all aspects of the IHC test - what is required for a full technical validation and implementation of a new test.

Full technical validation

## Proficiency testing in immunohistochemistry—experiences from Nordic Immunohistochemical Quality Control (NordiQC)

Mogens Vyberg<sup>1,2</sup> · Søren Nielsen<sup>1</sup>

**Table 3** Major causes of insufficient staining reactions

1. Less successful antibodies (17 %)
  - a. Poor antibodies<sup>a</sup>
  - b. Less robust antibodies<sup>b</sup>
  - c. Poorly calibrated RTUs
  - d. Stainer platform dependent antibodies
2. Insufficiently calibrated antibody dilutions (20 %)
3. Insufficient or erroneous epitope retrieval (27 %)
4. Error-prone or less sensitive visualization systems<sup>c</sup> (19 %)
- 5 Other (17 %)
  - a. Heat-induced impaired morphology
  - b. Proteolysis induced impaired morphology
  - c. Drying out phenomena
  - d. Stainer platform-dependant protocol issues
  - e. Excessive counterstaining impairing interpretation

<sup>a</sup> Consistently gives false negative or false positive staining or a poor signal-to-noise ratio in one or more assessment runs

<sup>b</sup> Frequently giving inferior staining results, e.g., due to mouse-anti-Golgi reactions or sensitive to standard operations as blocking of endogenous peroxidase

<sup>c</sup> Biotin-based detection kit for cytoplasmic epitopes, use of detection kits providing a too low sensitivity, or use of detection kits and chromogens giving imprecise localization of the staining signals complicating the interpretation

### Main causes of insufficient staining reactions are related to:

- The choice of antigen retrieval method
- The choice of primary antibody (Concentrate or RTU)
  - a) Calibration of the antibodies (dilutions)
  - b) Stainer platform dependent antibodies
- The choice of detection system

**83 % of insufficient results**

89 markers assessed during the period 2003-2015 and several markers have been assessed several times.  
Seven runs for HER2 ISH (more than 30000 slides assessed)





## Proficiency testing in immunohistochemistry—experiences from Nordic Immunohistochemical Quality Control (NordiQC)

Mogens Vyberg<sup>1,2</sup> · Søren Nielsen<sup>1</sup>

### Problems related to the choice of antigen retrieval methods:

- Use of non-alkaline HIER buffer (low pH buffer)
- Use of inefficient / too short HIER period
- Use of no / enzymatic pre-treatment instead of HIER
- Use of excessive retrieval procedure → impaired morphology

**False positive or false negative results**

Table 3 Major causes of insufficient staining reactions

1. Less successful antibodies (17 %)
  - a. Poor antibodies<sup>a</sup>
  - b. Less robust antibodies<sup>b</sup>
  - c. Poorly calibrated RTUs
  - d. Stainer platform dependent antibodies
2. Insufficiently calibrated antibody dilutions (20 %)
3. Insufficient or erroneous epitope retrieval (27 %)
4. Error-prone or less sensitive visualization systems<sup>c</sup> (19 %)
- 5 Other (17 %)
  - a. Heat-induced impaired morphology
  - b. Proteolysis induced impaired morphology
  - c. Drying out phenomena
  - d. Stainer platform-dependant protocol issues
  - e. Excessive counterstaining impairing interpretation

**27% insuff.**

<sup>a</sup> Consistently gives false negative or false positive staining or a poor signal-to-noise ratio in one or more assessment runs

<sup>b</sup> Frequently giving inferior staining results, e.g., due to mouse-anti-Golgi reactions or sensitive to standard operations as blocking of endogenous peroxidase

<sup>c</sup> Biotin-based detection kit for cytoplasmic epitopes, use of detection kits providing a too low sensitivity, or use of detection kits and chromogens giving imprecise localization of the staining signals complicating the interpretation

## Technical aspects of immunohistochemistry & pitfalls - Analytical phase

The purpose of antigen retrieval is to unmask antigenic determinants (epitopes) and recover immuno-reactivity

Antigen retrieval procedures for formalin fixed tissue:

- ☐ Heat Induced Epitope Retrieval (HIER)
- ☐ Tissue digestion using proteolytic enzymes
- ☐ Combined pre-treatment (HIER with proteolytic digestion)

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

0022-1554/91/\$3.30  
The Journal of Histochemistry and Cytochemistry  
Copyright © 1991 by The Histochemical Society, Inc.

Vol. 39, No. 6, pp. 741-748, 1991  
Printed in U.S.A.

## Rapid Communication

### Antigen Retrieval in Formalin-fixed, Paraffin-embedded Tissues: An Enhancement Method for Immunohistochemical Staining Based on Microwave Oven Heating of Tissue Sections

SHAN-RONG SHI, MARC E. KEY,<sup>1</sup> and KRISHAN L. KALRA

BioGenex Laboratories

Received for publication

We describe a new method for antigen retrieval in formalin-fixed, paraffin-embedded tissue sections using microwave oven heating.

# HIER

#### Shi et al. demonstrated that :

- Enzyme pre-digestion of tissue could be omitted.
- Incubation time with primary antibodies could be reduced, or dilutions of primary antibodies could be increased.
- Staining could be achieved on long-term formalin fixed that failed to stain with conventional methods.
- Certain antibodies which were typically unreactive with formalin-fixed tissue gave excellent staining.

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

**The mechanism of HIER is not completely understood, but several hypothesizes has been proposed:**

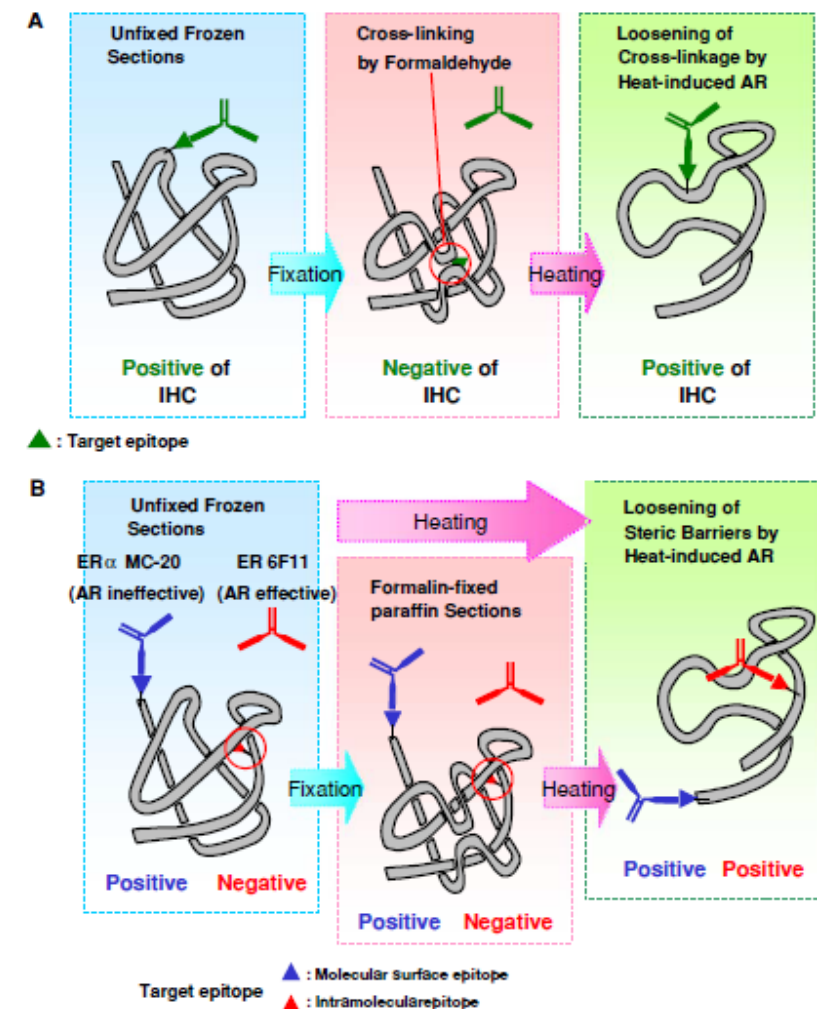
**Heating tissue sections in an appropriate buffer may unmask epitopes by :**

- ☐ **Hydrolysis/disruption of methylene cross-links formed by formalin fixation**
- ☐ **Extraction of diffusible blocking proteins**
- ☐ **Precipitation/denaturation of proteins**
- ☐ **Rehydration of the tissue section allowing better penetration of the antibody**
- ☐ **Removal of tissue-bound calcium ions by chelating substances**
- ☐ **Other mechanism's ?**

# Hypothesis for the mechanism for heat-induced antigen retrieval occurring on fresh frozen sections without formalin-fixation in immunohistochemistry

Kochi Kakimoto · Susumu Takekoshi · Katsuhiko Miyajima · R. Yoshiyuki Osamura

**Fig. 7** Conventional hypothesis (A). Formaldehyde fixation can alter the three-dimensional structure of the epitope cross-linkages; these can be reversed by high-temperature heating. Our suggested mechanism for AR in IHC (B): Antibodies recognizing molecular surface epitopes, such as ER $\alpha$  MC-20, do not show increases in detection levels with or without heating whereas antibodies recognizing intramolecular epitopes, such as ER 6F11, show significantly increased detection levels because the three-dimensional structure is likely to be altered by heat denaturation



The unfixed frozen sections, which did not show immunostaining with nine antibodies, were clearly stained after heating the sections

These results indicate that other mechanisms of breaking formalin-induced cross-linkages may be present.

The authors proposed that :

One of the other mechanisms for heat-induced AR is that accessibility to the target epitopes of antigenic proteins is limited by natural steric barriers even in the fresh state caused by the antigenic protein itself.

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Efficient HIER depends on:

- ☐ pH of the HIER buffer
- ☐ Temperature
- ☐ Time
- ☐ Elementary nature of the HIER buffer (e.g., Citrate; TRIS; EDTA; TE)

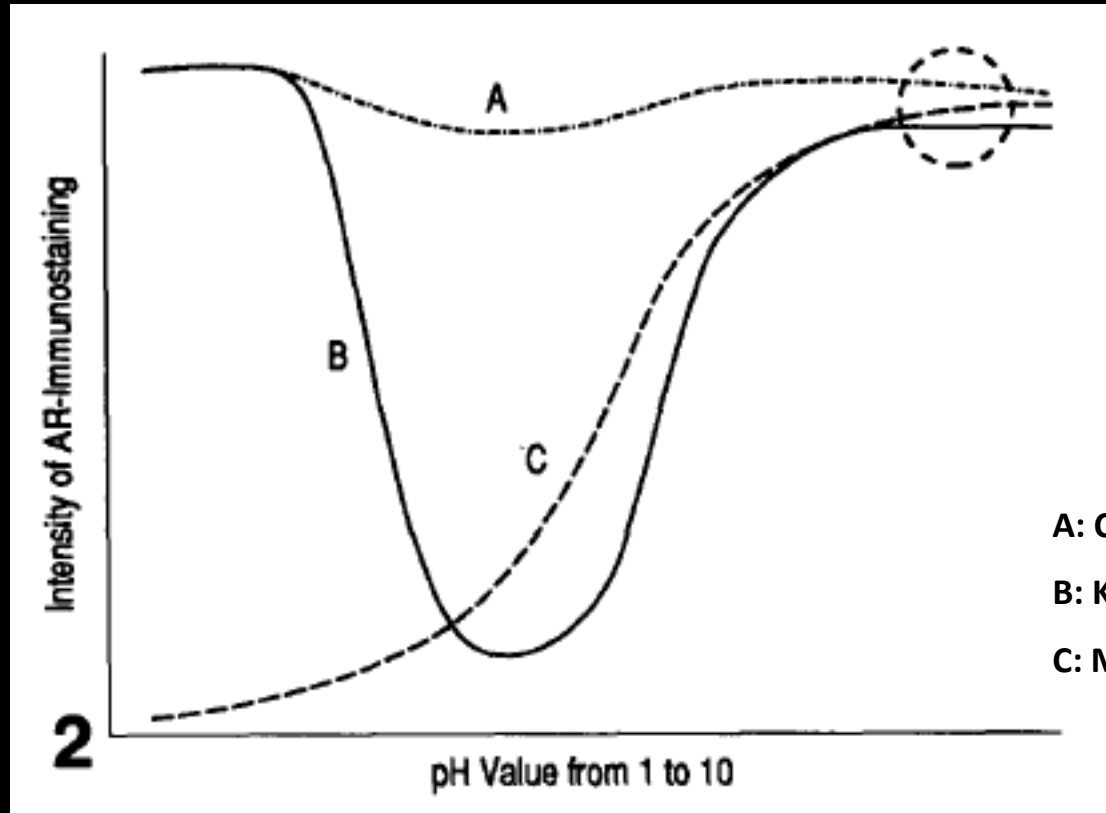
Less sensitive to routinely fixed tissue (formalin) compared to enzymatic pre-treatment

> 95% of all commonly used antibodies require HIER



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Shi SR et al. J Histochem Cytochem 1995 43:193-201



A: CD20 (clone L26)

B: Ki-67 (clone MIB1)

C: MSA (clone HMB45)

## Efficient HIER - Influence of pH

Demonstrated that the performance of monoclonal antibodies were highly influenced by pH of the Antigen Retrieval buffer (AR).

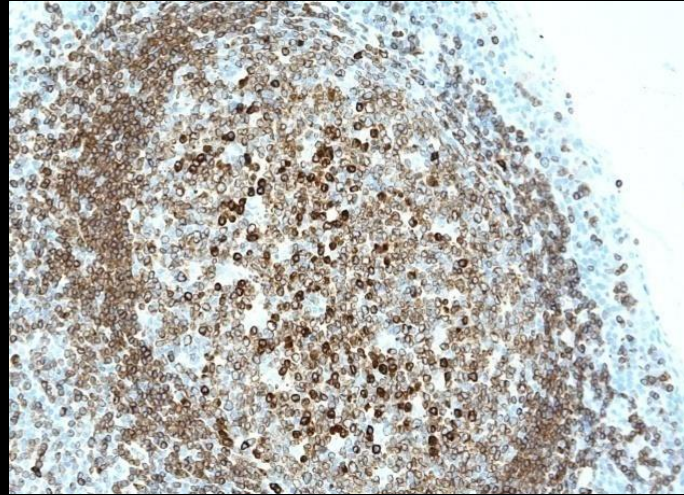
Also, the results indicate the advantage of using an AR solution of higher pH value (8-9).

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

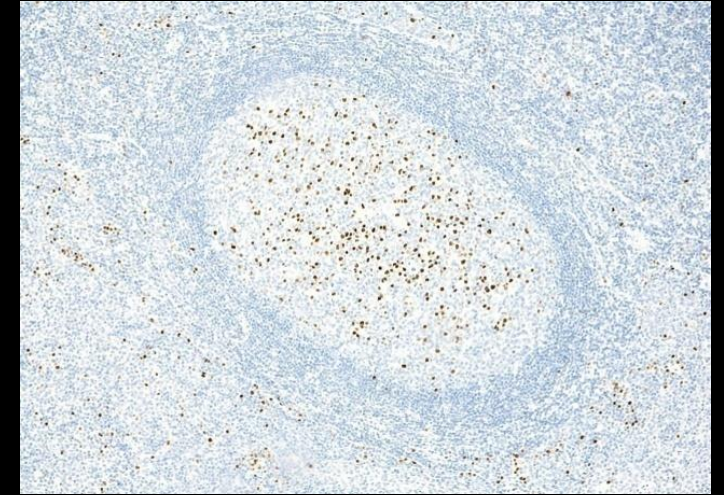
## Efficient HIER - Influence of pH

HIER in TRS pH 6.1  
(20 min at 97°C)

CD79, JCB117 (1:300)

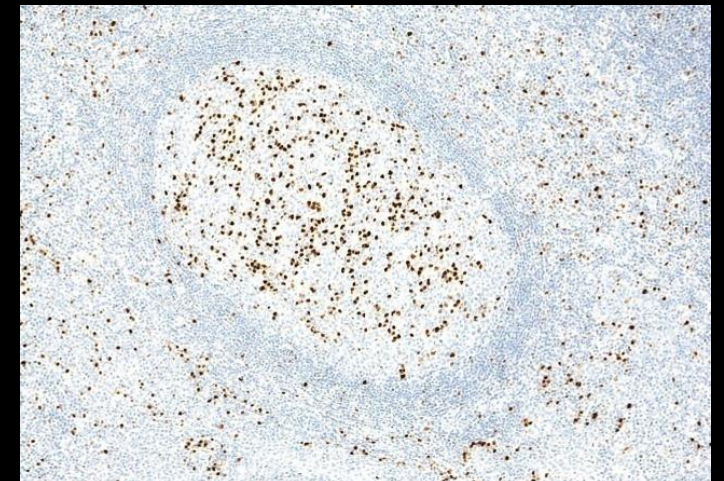


MUM-1, MUM1p (1:400)



Autostainer: Flex+ as the detection system

HIER in TRS pH 9  
(20 min at 97°C)



Tonsillar tissue fixed in 10% NBF (48h).

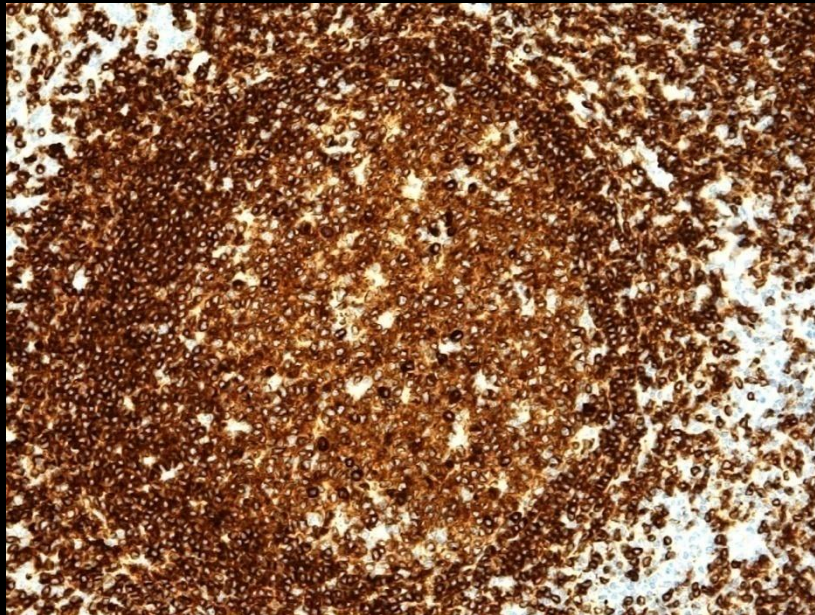


# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Efficient HIER - Influence of pH

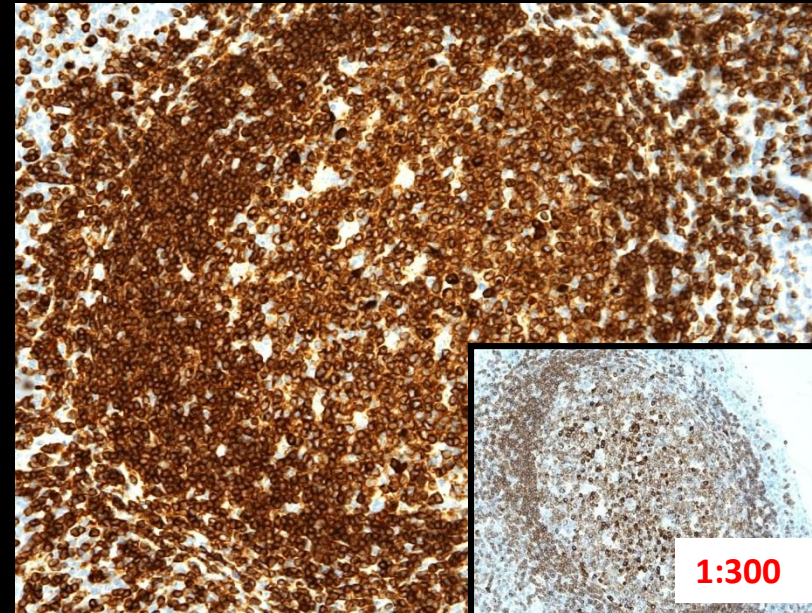
HIER in TRS pH 9

CD79, JCB117 (1:300)



HIER in TRS pH 6.1

CD79, JCB117 (1:50)



Tonsillar tissue fixed in 10% NBF (48h).  
Flex+ as detection system

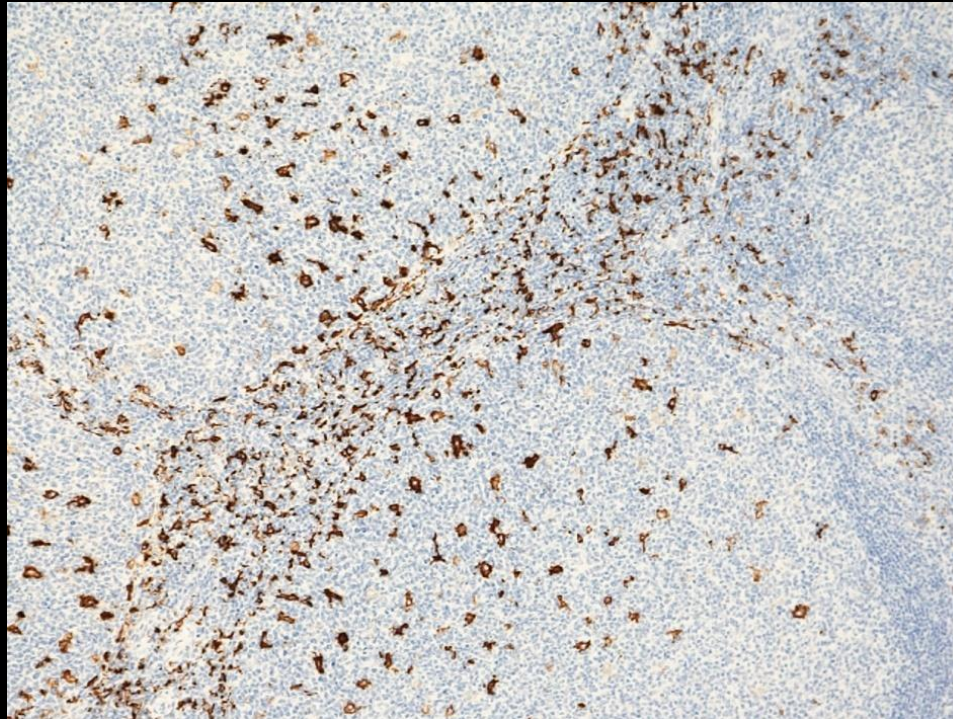


# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Efficient HIER - Influence of pH

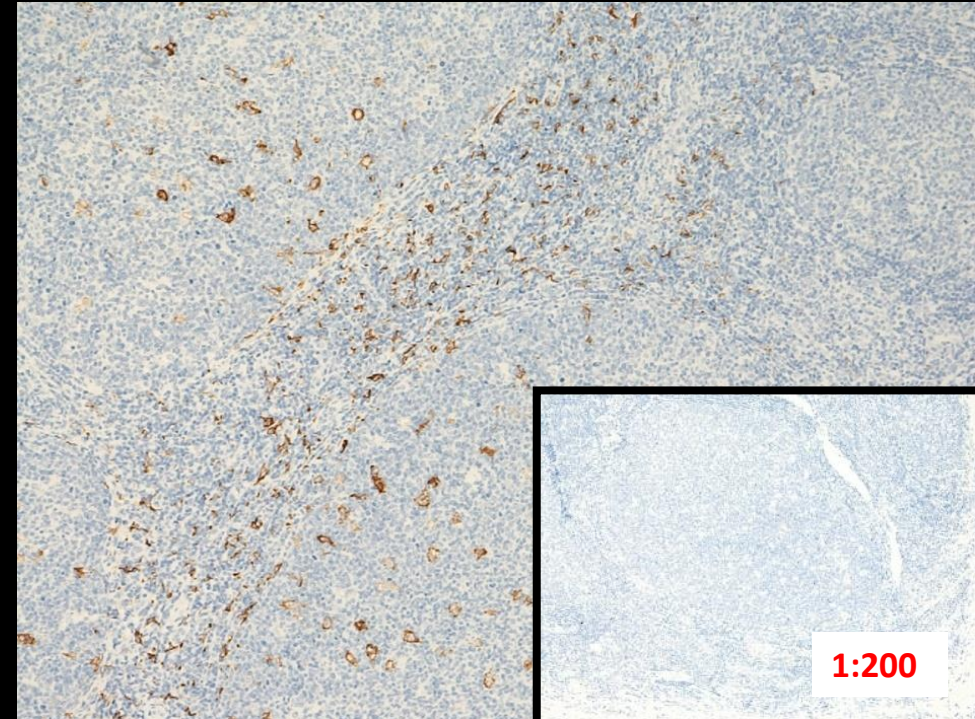
HIER in TRS pH 9

CD163, MRQ-26 (1:200)/Flex+



HIER in TRS pH 6.1

CD163, MRQ-26 (1:25)/Flex+



For app. 85-90% of the epitopes, HIER in buffers at pH 8-9 is preferable to pH6

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Ready To Use products: Recommendations to antigen retrieval provided by the manufactures ?

## CK7 Run 62

Table 3. Proportion of sufficient and optimal results for CK7 for the most commonly used RTU IHC systems

RTU systems	<u>Recommended protocol settings*</u>		<u>Laboratory modified protocol settings**</u>	
	Sufficient	Optimal	Sufficient	Optimal
Dako AS mAb OV-TL 12/30 <b>IR619</b>	100% (12/12)	92% (11/12)	100% (11/11)	91% (10/11)
Dako Omnis mAb OV-TL 12/30 <b>GA619</b>	100% (32/32)	97% (31/32)	100% (26/26)	92% (24/26)
Leica Bond III/MAX mAb RN7 <b>PA0942/PA0138</b>	100% (6/6)	17% (1/6)	100% (11/11)	45% (5/11)
VMS Ultra/XT rmAb SP52 <b>790-4462</b>	100% (16/16)	69% (11/16)	97% (98/101)	85% (86/101)

\* Protocol settings recommended by vendor – Retrieval method and duration, Ab incubation times, detection kit, IHC stainer/equipment.

\*\* Significant modifications: retrieval method, retrieval duration and Ab incubation time altered, detection kit – only protocols performed on the specified vendor IHC stainer integrated.

17% (1/6)

**BERS1 (citrate based HIER buffer)**

83% (5/6)

**BERS2 (alkaline based HIER buffer)**

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Efficient HIER - Influence of time and temperature

Taylor CR et al : *Applied Immunohistochemistry* 1996; 4(3) : 144-166 - *Temperature and time are inversely related* :

Similar strong intensity of staining could be generated by the following heating conditions:

100°C for 20 min = 90°C for 30 min = 80°C for 50 min = 70°C for 10 h

Balaton AJ et al : *Applied Immunohistochemistry* 1996; 4(4) : 259 - 263

Optimal staining intensity could be generated by the following heating conditions:

MWO at 100°C for 20 min = Pressure cooker at 120°C for 3 min



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## HIER buffer - Influence of time and temperature

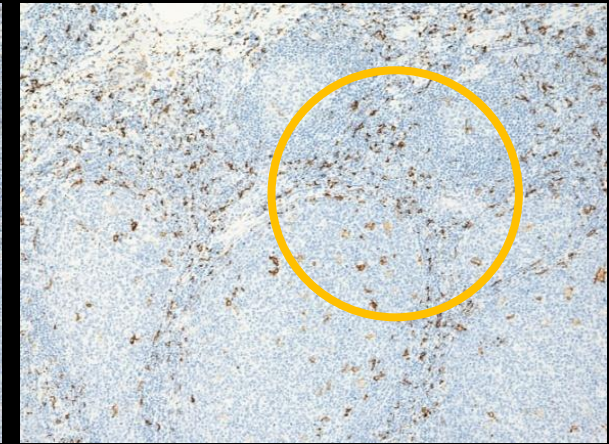
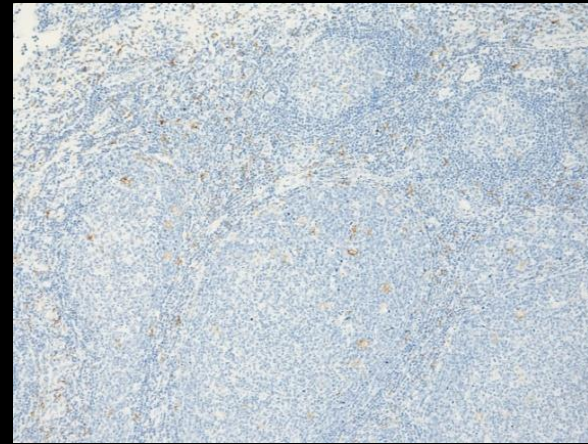
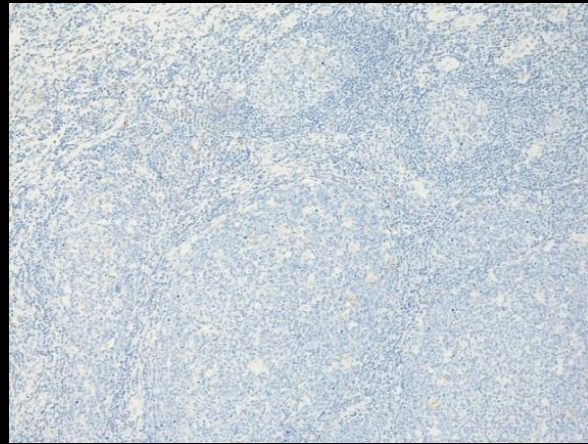
CD163, MRQ-26 (1:200)

10 min

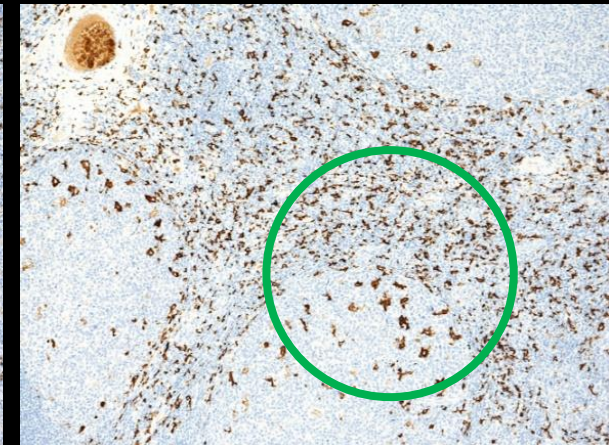
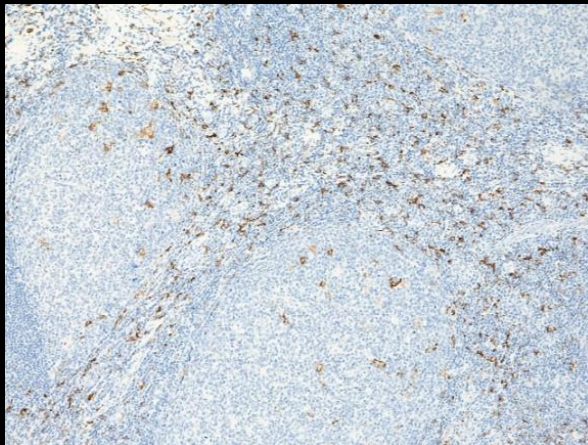
20 min

80 min

HIER at 80°C  
TRS pH 9, Flex+



HIER at 97°C  
TRS pH 9, Flex+



Tonsillar tissue fixed in 10% formalin (48h).



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Efficient HIER - Influence of time and temperature

TRS pH 9 (80°C/60')

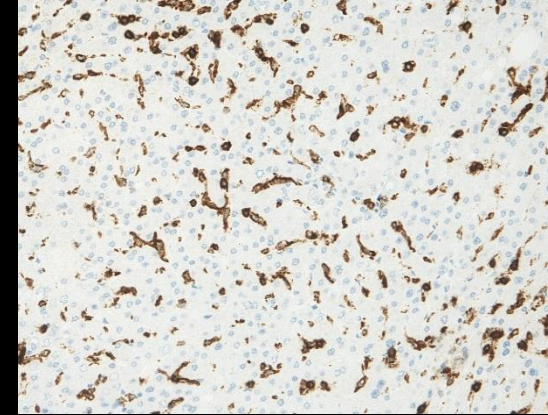
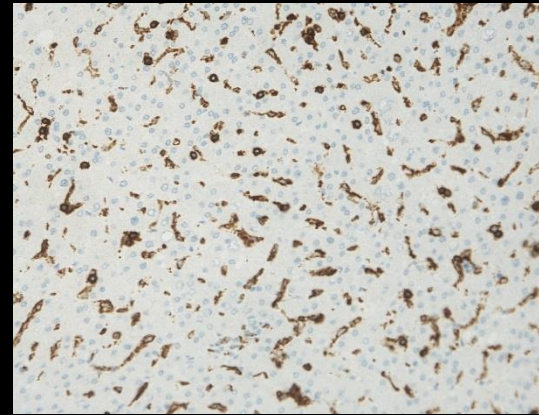
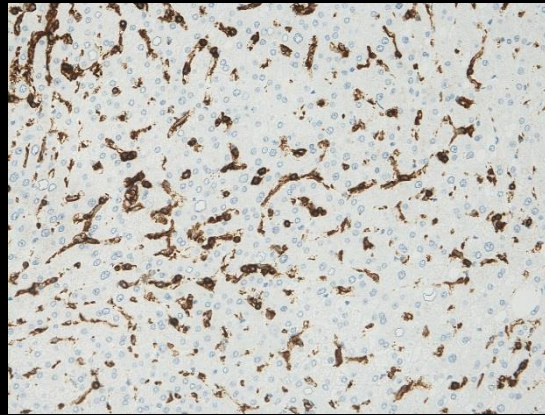
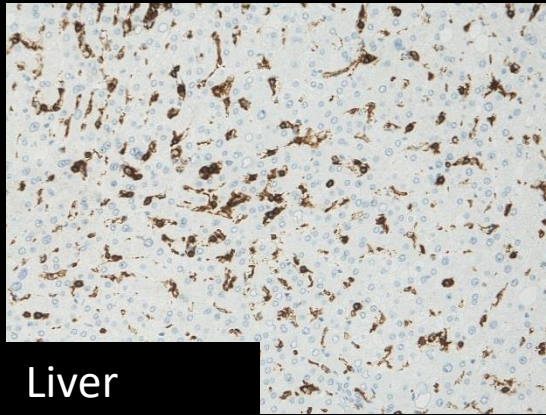
TRS pH 9 (90°C/30')

TRS pH9 (90°C/60')

TRS pH9 (97°C/24')

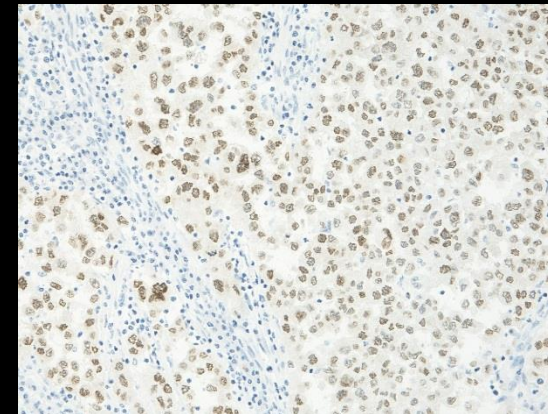
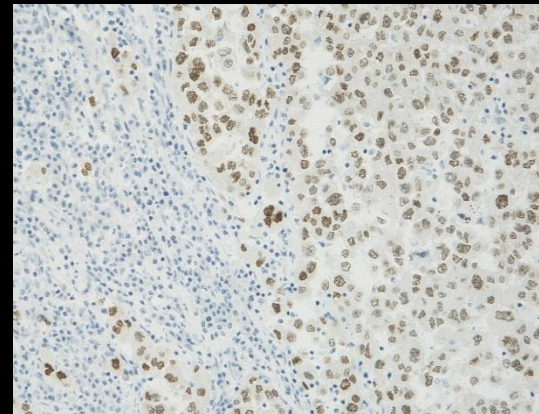
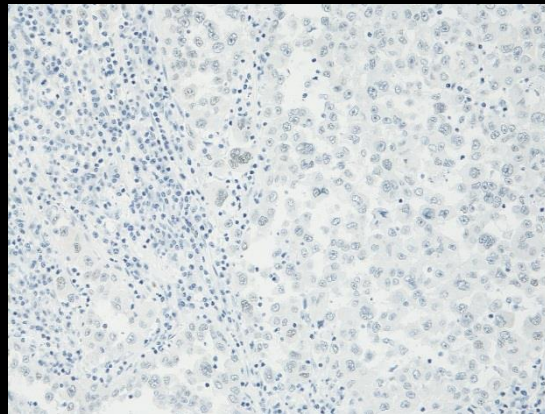
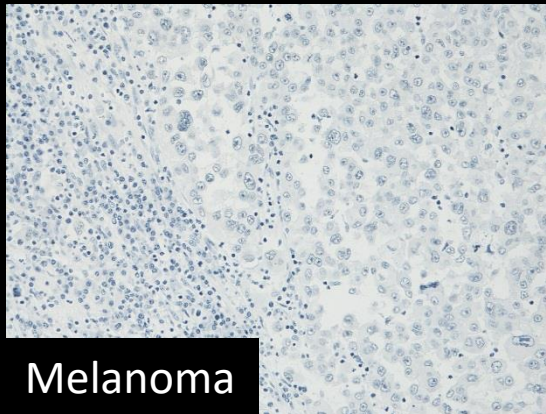
CD45/LCA  
(2B11+PD7/26)

Liver



SOX10  
(BS7)

Melanoma





# ANTIGEN RETRIEVAL TECHNIQUES IN IMMUNOHISTOCHEMISTRY: COMPARISON OF DIFFERENT METHODS

STEFANO A. PILERI<sup>1\*</sup>, GIOVANNA RONCADOR<sup>1</sup>, CLAUDIO CECCARELLI<sup>1</sup>, MILENA PICCIOLI<sup>1</sup>, ASPASIA BRISKOMATIS<sup>1</sup>, ELENA SABATTINI<sup>1</sup>, STEFANO ASCANI<sup>1</sup>, DONATELLA SANTINI<sup>1</sup>, PIER PAOLO PICCALUGA<sup>1</sup>, ORNELLA LEONE<sup>1</sup>, STEFANIA DAMIANI<sup>1</sup>, CESARINA ERCOLESSI<sup>1</sup>, FEDERICA SANDRI<sup>1</sup>, FEDERICA PIERI<sup>1</sup>, LORENZO LEONCINI<sup>2</sup> AND BRUNANGELO FALINI<sup>3</sup>

<sup>1</sup>Second Service of Pathologic Anatomy and Haematopathology Section, Bologna University, Italy

<sup>2</sup>Institute of Pathologic Anatomy, Siena University, Italy

<sup>3</sup>Haematopathology Laboratory, Institute of Haematology, Perugia University, Italy

## Chemical composition of the HIER buffer's

Standard low pH buffer's (e.g., citrate based pH 6.0)

Standard high pH buffer's (e.g., TE based pH 8-10)

Modified low pH buffers pH 6.1-6.2 : S1699/S1700 (Dako) or Diva Decloaker (Biocare)

## Overall best performance:

HIER in EDTA pH 8.0 (compare with Tris-HCL pH 8.0)

Table I—Results obtained under different methodological conditions with the antibodies most commonly used in the study of the haemolymphopoietic system and related disorders

Clone	Specificity	Source	Dilution	No AgR	PT	HBAR+ citrate	HBAR+ Tris-HCl pH8	HBAR+ EDTA pH8
O10	CD1a	Immunotech	1:40	—	—	+— — —	++ — —	++++
Poly	CD3	DAKO	1:300	—	++ — —	+— — —	++ — —	++++
C8/144B	CD8	Dr Mason	1:6	—	—	+— — —	++ — —	++++
			1:400	—	—	+— — —	++ — —	++++
C3D-1	CD15	DAKO	1:6	+ — — —	—	+— — —	++ — —	++++
			1:320	++ — —	—	++++	++ — —	++++
L26	CD20	DAKO	1:200	+ — — —	+ — — —	++++	++ — —	++++
			1:3200	+ — — —	+ — — —	++++	++ — —	++++
IF8	CD21	DAKO	1:10	—	++++	—	—	—
MB16	CD23	DAKO	1:50	—	—	+ — — —	+++ —	++++
Ber-H2	CD30	Professor Stein	1:10	—	—	+ — — —	++++	++++
			1:320	—	—	+ — — —	++++	++++
QBEND-10	CD34	BioGenex	1:20	+ — — —	—	++++	++++	++++
			1:400	+ — — —	—	++++	++++	++++
BerMACDRC	CD35	DAKO	1:5	+ — — —	++++	—	+ — — —	+ — — —
MAC383	CD40	Immunotech	1:100	—	++++	—	—	—
DF-T1	CD43	DAKO	1:200	+ — — —	—	++++	+++ —	++++
			1:1600	+ — — —	—	++++	++++	++++
PD7/26+2B11	CD45	DAKO	1:200	—	+ — — —	++++	++++	++++
			1:4000	—	—	+ — — —	++++	++++
UCHL-1	CD45R0	DAKO	1:120	+ — — —	+ — — —	++++	++++	+++ —
K1-B3	CD45R	Professor Parwaresch	1:80	++ — —	+ — — —	++++	++++	++++
			1:320	++ — —	+ — — —	++++	++++	++++
4KB5	CD45RA	DAKO	1:20	++ — —	—	++++	+++ —	+++ —
	CD57	Becton	1:20	++ — —	++ — —	++++	++++	++++
Y2/51	CD61	DAKO	1:5	—	++++	+ — — —	+ — — —	+++ —
KP1	CD68	DAKO	1:640	+ — — —	++ — —	++++	++ — —	++++
PG-M1	CD68	Professor Falini	1:20	+ — — —	++ — —	++++	++ — —	++++
JCB117	CD79a	Dr Mason	1:10	+ — — —	—	++++	++++	++++
Kim-4p	Follicular dendritic cells	Professor Parwaresch	1:5	—	++++	++ — —	++ — —	+ — — —
DBA.44	Hairy cells	Professor Delsol	1:5	++ — —	—	++++	++++	++++
JC159	GlycophorinA	DAKO	1:320	+ — — —	—	++++	++++	+++ —
NP57	Neutrophilic elastase	DAKO	1:10	++++	—	—	—	—
M616	FVIII RAG	DAKO	1:6	+ — — —	++ — —	++++	++ — —	++++
Poly	Lysozyme	DAKO	1:800	++ — —	++++	++++	++++	++++
Poly	IgA	DAKO	1:2000	+ — — —	++++	++++	++++	++++
Poly	IgG	DAKO	1:5000	++ — —	++++	++++	++++	++++
Poly	IgM	DAKO	1:5000	—	++ — —	++++	++++	++++
Poly	IgD	DAKO	1:1000	—	—	++++	++++	++++
Poly	κ-Ig light chain	DAKO	1:10 000	++ — —	++++	++++	++++	+++ —
Poly	λ-Ig light chain	DAKO	1:12 000	++ — —	++++	++++	++++	+++ —
Poly	Protein S-100	DAKO	1:2000	++ — —	++++	++++	++++	++++
Poly	MPO	DAKO	1:10 000	++ — —	++++	++++	++++	++++

CD—cluster of differentiation; No AgR—no antigen retrieval; PT—proteolytic treatment; HBAR—heat-based antigen retrieval; Poly—polyclonal antibody; FVIII RAG—Factor VIII-related antigen; MPO—myeloperoxidase.

In bold: overnight incubation of the primary antibody+SABC technique.

—=completely negative result; + — — —=weak positivity in a percentage of cells expected to be positive; ++ — —=weak positivity in all cells expected to be positive; +++ —=moderately strong positivity in all cells expected to be positive; ++++=very strong positivity in all cells expected to be positive.



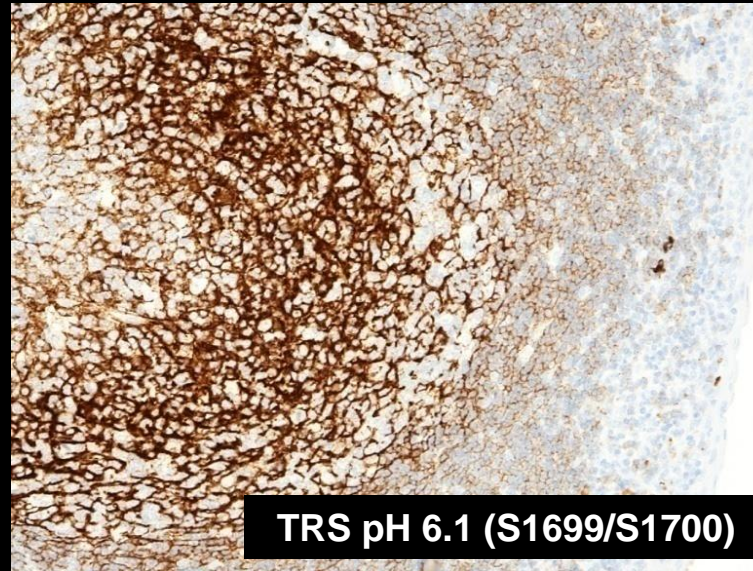
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Modified low pH buffers

CD21, clone 1F8



CD21, clone 1F8



CD21, clone 2G9



HIER time 20' / Flex+

Markers requiring the TRS Low pH 6.1 (Dako, S1699/S1700) or Diva Decloaker pH 6.2 (Biocare, DV2004) :

EPCAM (clone EP-4 or MOC-31 or "VU-1D9"); GP200 (clone SPM 314 or 66.4.C2); CD21 clone 1F8; CD61 clone Y2/51; NGFR clone MRQ-21; Desmoglein-3 clone BC11, PHH3 clone BC37 and .....

Mandatory for : CD7 clone CBC 3.7; CD30 clone ConD6/B5; CD5 clone Leu1



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Modified low pH buffers

CD30, ConD6/D5 (1:50)  
(Hodgkin Lymphoma)

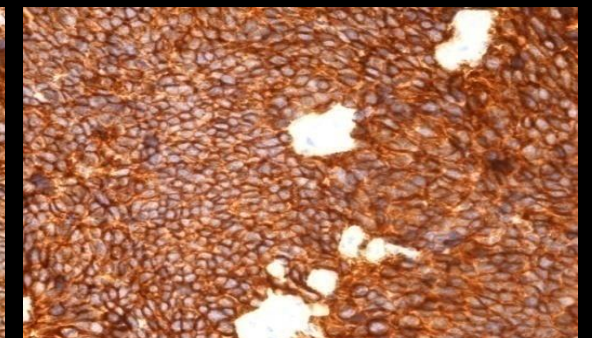
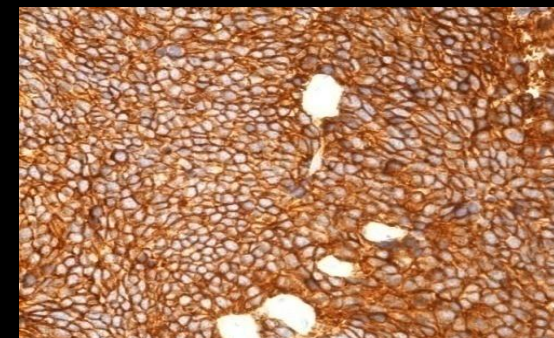
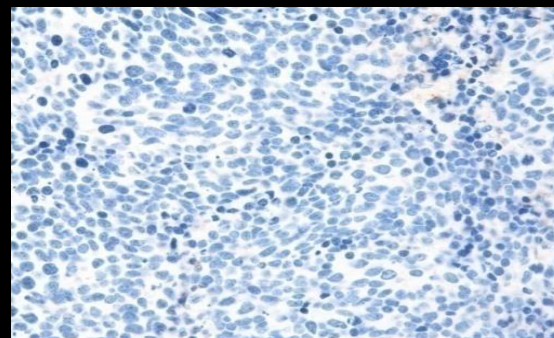
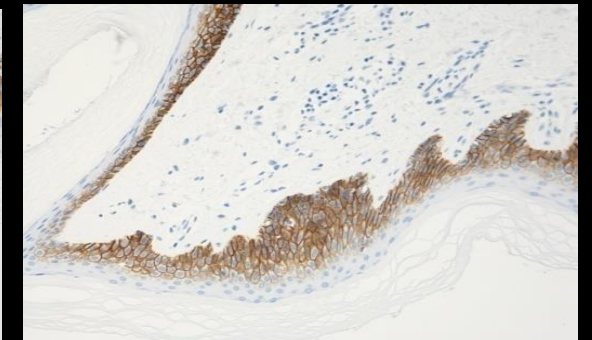
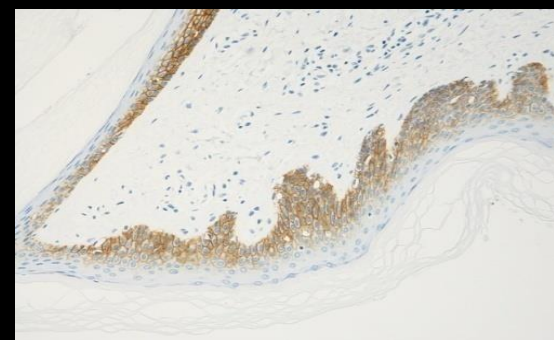
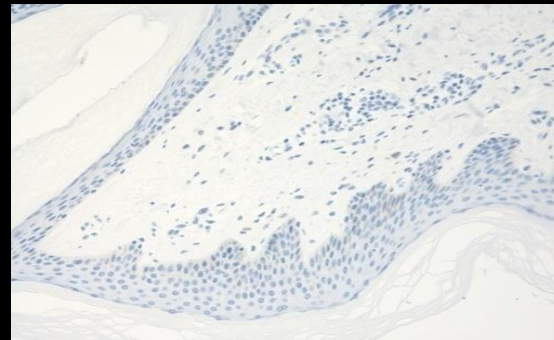
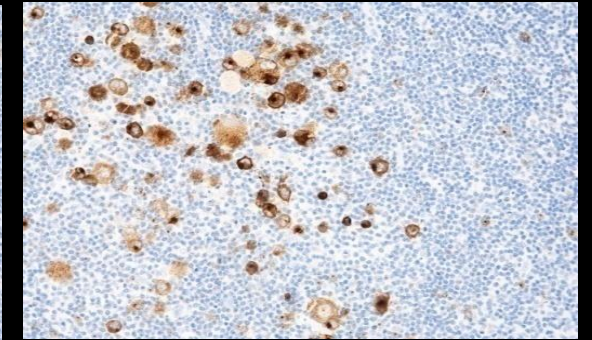
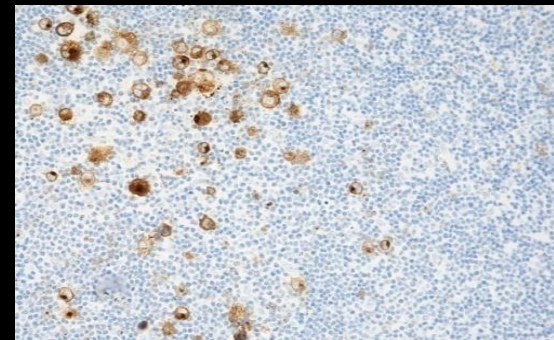
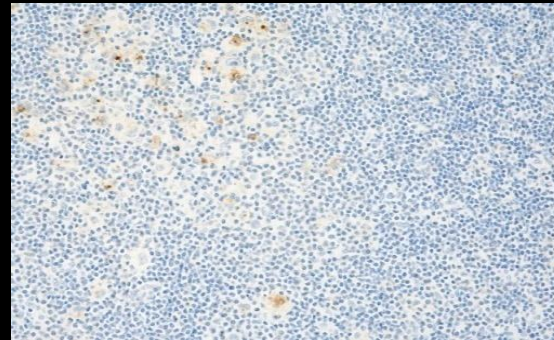
Desmoglein-3, BC11 (1:25)  
(Skin)

EP-CAM, MOC-31 (1:20)/EP4  
(Small cell carcinoma)

TRS pH 9 (Dako)  
PT / 99° / 20 min

TRS pH 6.1 (Dako S1700)  
PT / 99° / 20 min

Diva Decloaker (Biocare)  
PT / 99° / 20 min



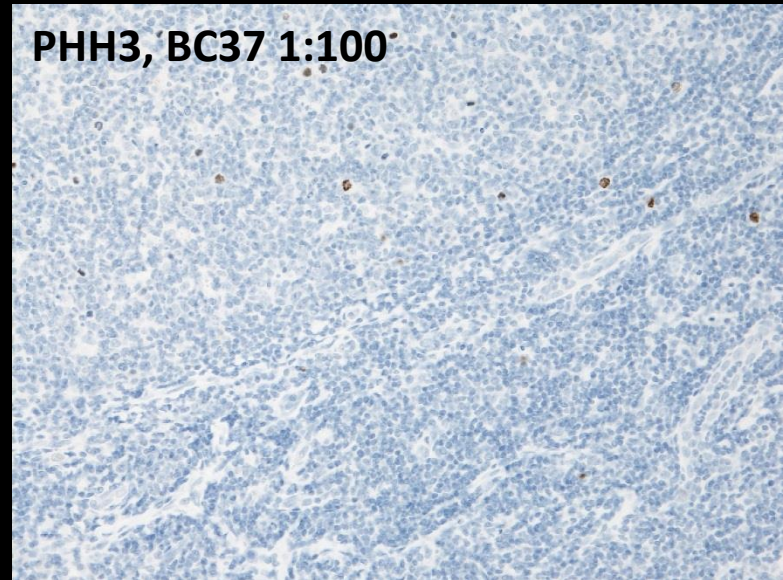
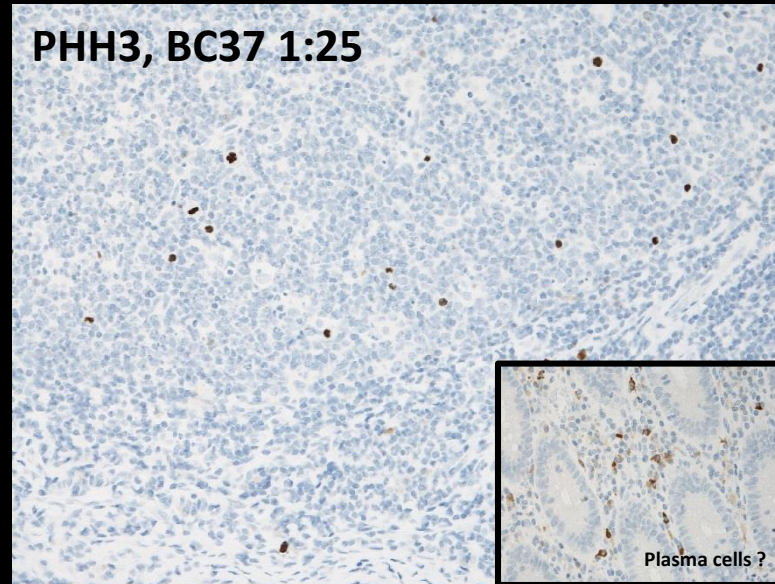


# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Modified low pH buffers

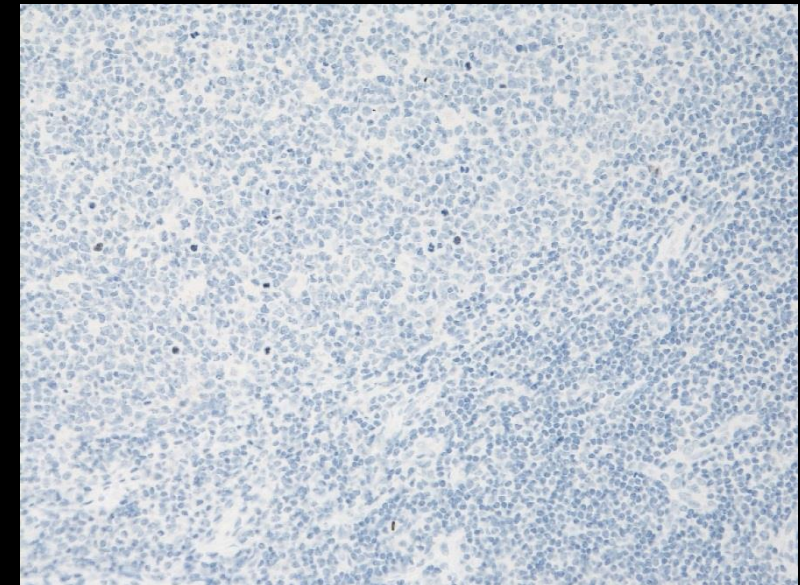
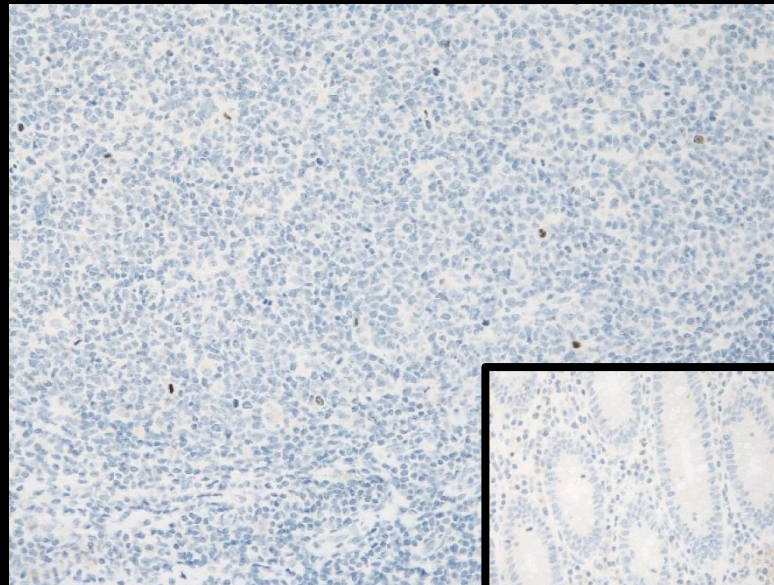
TRS pH 6.1 (Dako S1700)

24`/97 ° C



TRS High pH 9 (3-1)

24`/97 ° C

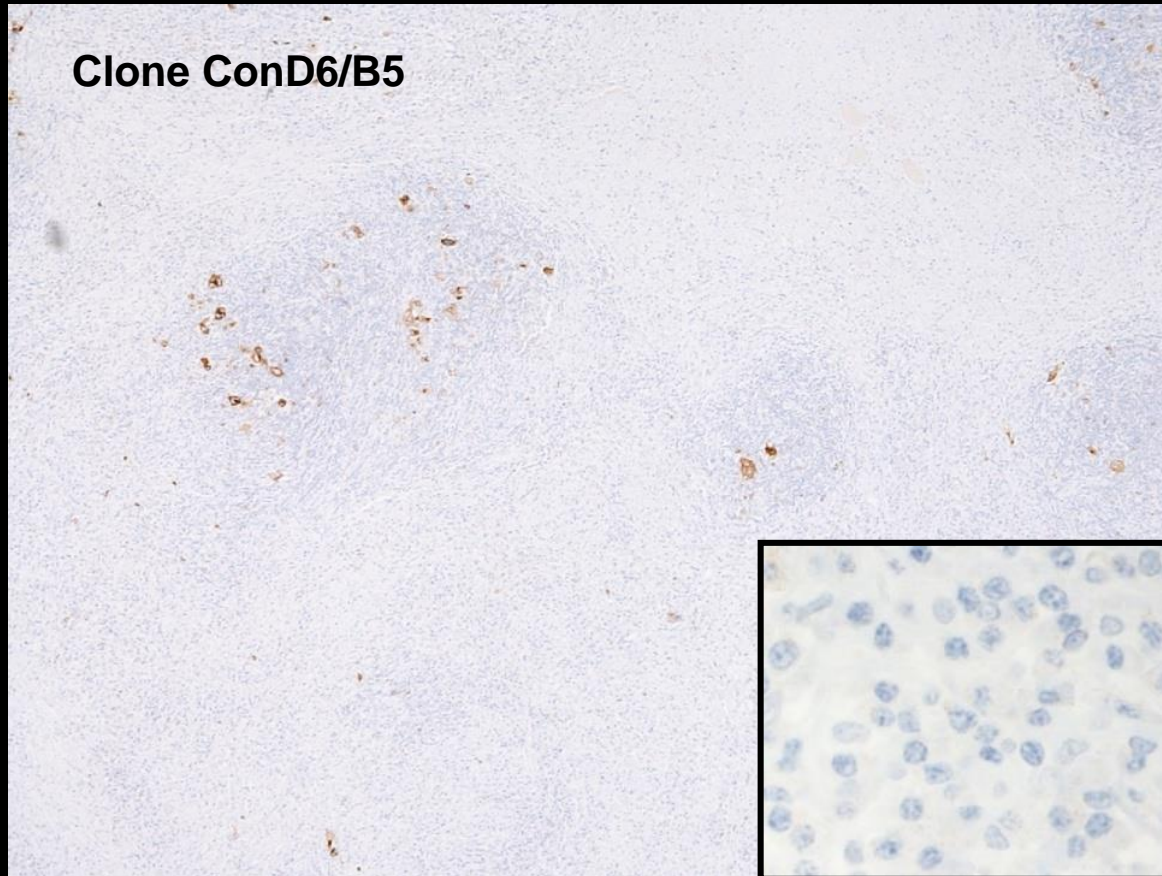




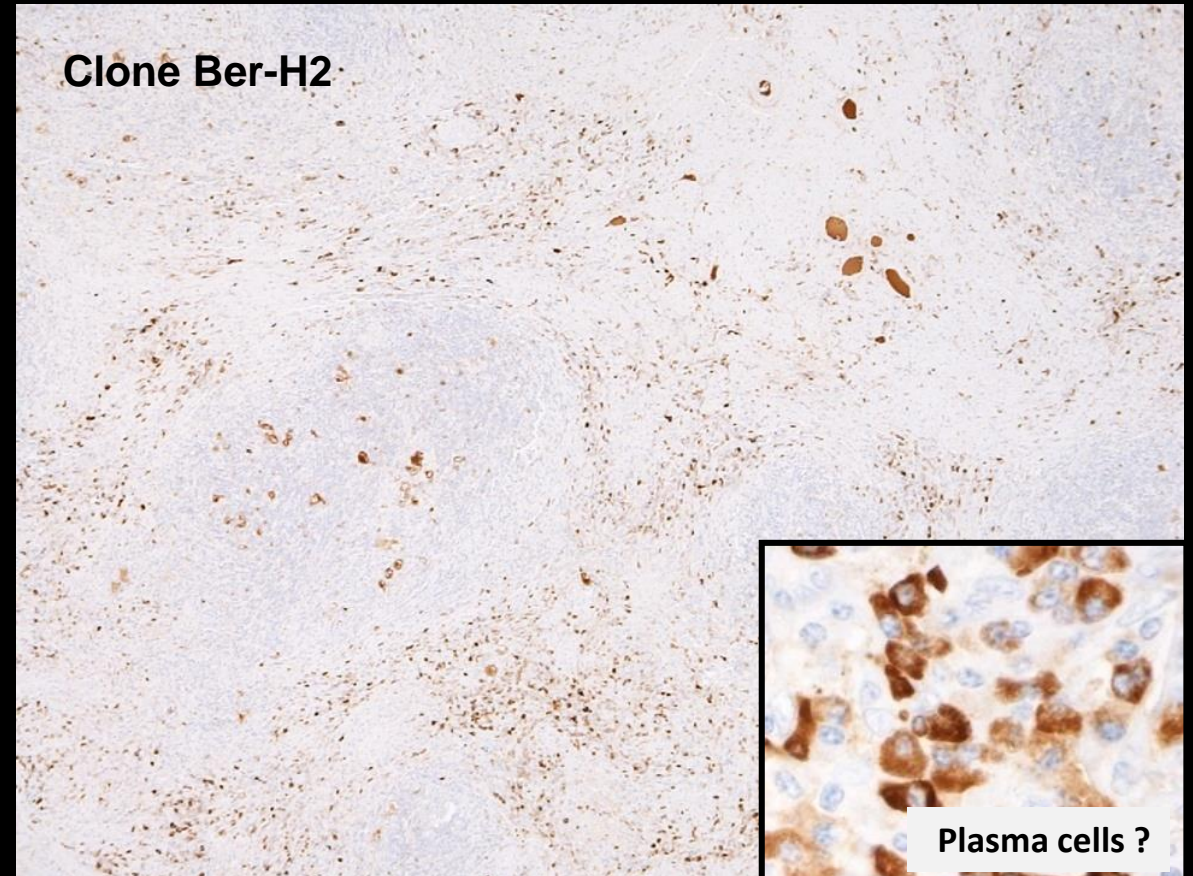
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Modified low pH buffers

### CD30



HIER buffer, TRS pH 6.1 (Dako S 1700)



Hodgkin's Lymphoma



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Modified low pH buffers

### CD30 clone ConD6/B5

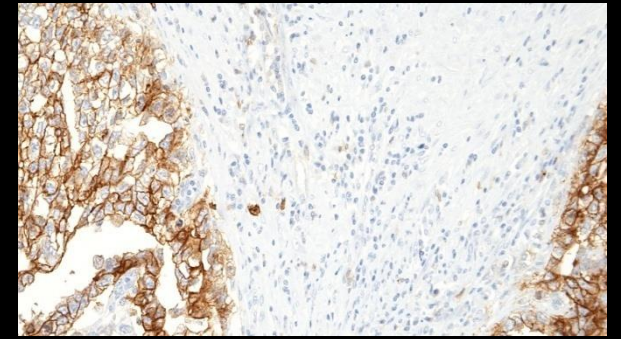
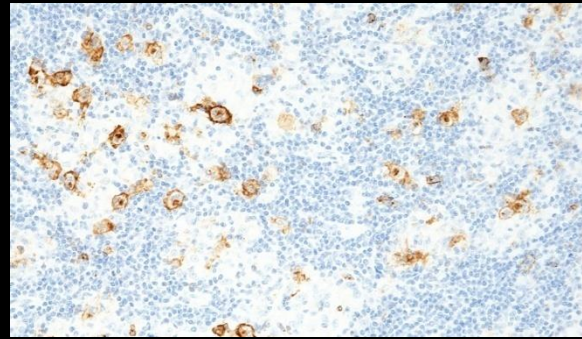
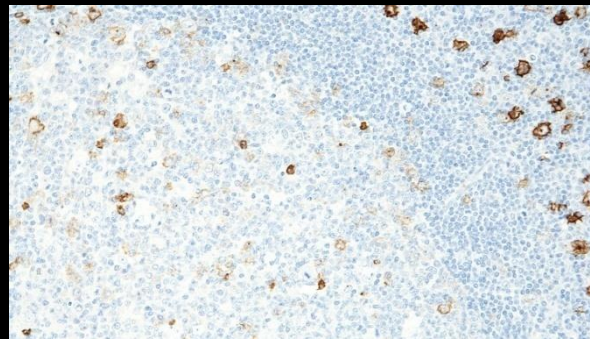
Tonsil

Hodgkin lymphoma

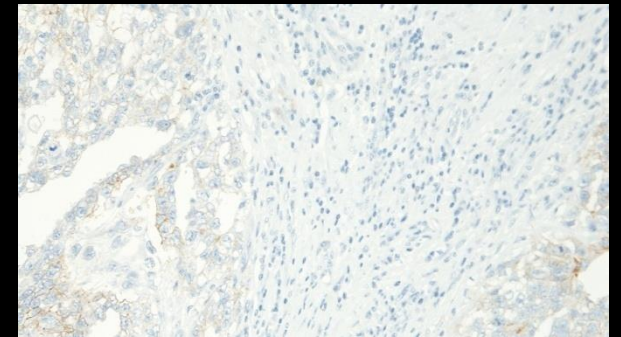
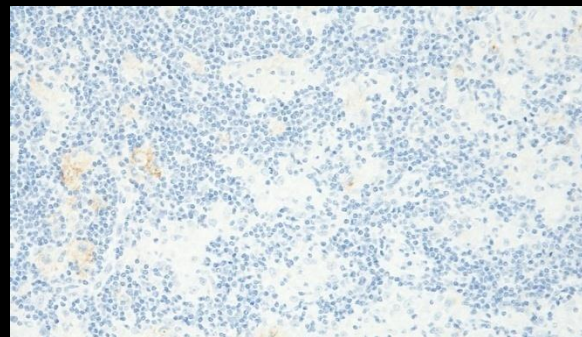
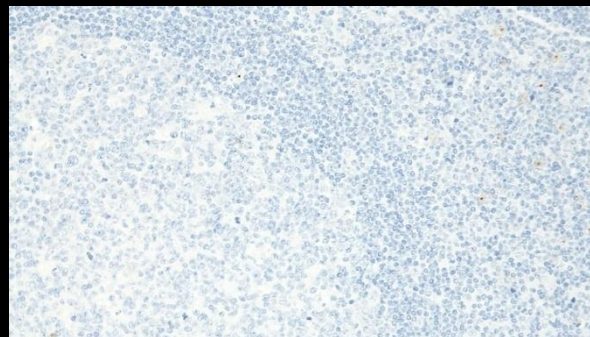
Embryonal carcinoma

HIER buffer, TRS pH 6.1

(Dako S 1700)



HIER buffer, Low pH  
(LabVision TA-999-DHBL)



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

The purpose of antigen retrieval is to unmask antigen epitopes /restore antigenic determinants and recover immuno-reactivity

Antigen retrieval procedures for formalin fixed tissue:

- ❑ Heat Induced Epitope Retrieval (HIER)
- ❑ Tissue digestion using proteolytic enzymes
- ❑ Combined pre-treatment (HIER with proteolytic digestion)



Proteolytic enzymes cleave more or less specific amino acid sequences within peptide chains (and not covalent cross-links formed in tissues during formalin fixation).

→ Improves penetration of immuno-reagents into tissue structures and enhance accessibility to the epitopes of interest.

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Enzymatic digestion - the problem?

A significant proportion of Labs still use enzymatic pre-treatment for e.g, “old” markers as cytokeratins and S100 introduced back in 1980-1990.

Only few markers require enzymatic digestion for “optimal performance”.

### PAN-CK

Table 4. Pass rates for antibody cocktails combined with epitope retrieval methods in nine NordiQC runs

Pass rate for compiled data from run 15, 20, 24, 30, 36, 41, 47, 54 & 58								
	Total		HIER		Proteolysis		HIER + proteolysis	
	Protocols	Sufficient	Protocols	Sufficient	Protocols	Sufficient	Protocols	Sufficient
mAb AE1/AE3	1145	836 (73%)	1075	826 (77%)	49	6 (12%)	9	3 (33%)
mAb AE1/AE3/5D3	48	42 (88%)	47	42 (89%)	1	0	0	0
mAb AE1/AE3/PCK26	361	219 (61%)	48	22 (46%)	48	3 (6%)	258	192 (74%)
mAb MNF116	111	31 (28%)	53	9 (17%)	48	22 (46%)	9	2 (22%)

For mAb clone AE1/AE3 (PAN-CK) and S100: App. 4-5 % of all protocols were based on enzymatic pre-treatment

### S100

Table 5. Pass rates for S100 antibody combined with epitope retrieval methods in the last three NordiQC runs

Pass rate for compiled data from run 45, 50 & 59									
	Total		HIER		Proteolysis		HIER + proteolysis		No pretreatment
	Protocols	Sufficient	Protocols	Sufficient	Protocols	Sufficient	Protocols	Sufficient	Protocols
mAb 4C4.9	137	80 (58%)	110	71 (65%)	4	0	2	1	21
pAb NCL-L-S100p	30	18 (60%)	21	14 (67%)	6	2 (33%)	0	0	3
pAb Z0311	494	417 (84%)	444	386 (87%)	26	15 (58%)	3	2	21
pAb 760-2523	97	68 (70%)	82	62 (76%)	2	1	0	0	13
Total	758	583 (77%)	657	533 (81%)	38	18 (47%)	5	3	58



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## "Optimal" enzymatic digestion depends on:

Enzyme type

Concentration

Time

Temperature

Fixation type & time

Tissue type

### Most common Enzymes

Proteinase K

Pronase XIV

Pronase XXIV

Pepsin

Trypsin

Difficult to control and to standardizes within routine LAB

### Markers requiring enzymatic pretreatment :

FVIII (poly), LMV CK (CAM 5.2), PAN CK (MNF116), EGFR (various), TCR- $\beta$  (8A3).....

Extracellulare matrix proteins (COLL-III (poly), Laminin (poly) and COLL-IV (CIV-22) .....

Short time formalin fixation = gentle proteolysis

Long time formalin fixation = prolonged proteolysis

≤ 2% of all commonly used antibodies require enzymatic pre-treatment

## Choice of proteolytic enzyme

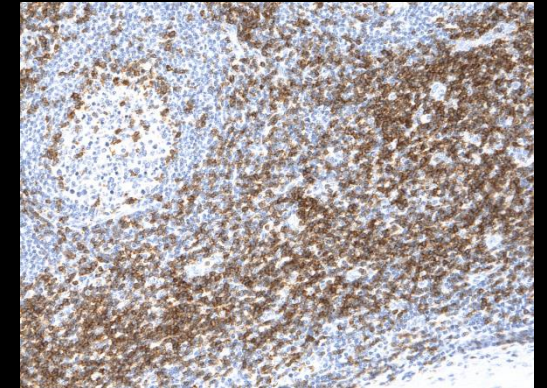
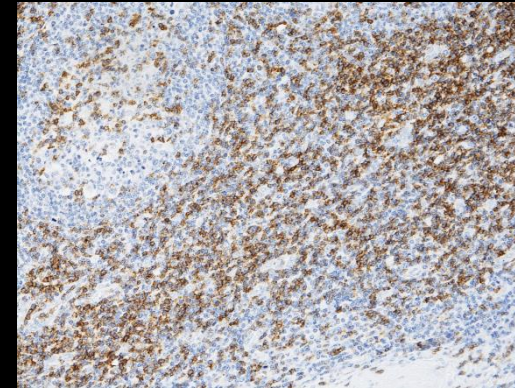
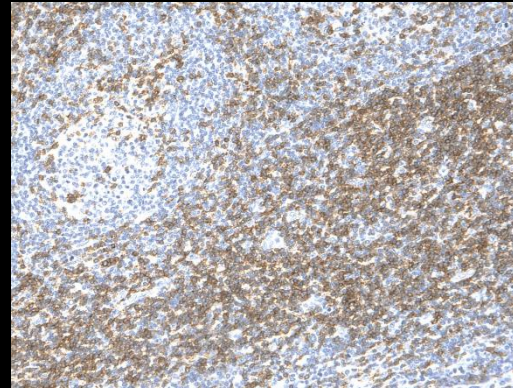
Trypsin (Biocare, RTU)  
40`

Pepsin (ZytoVision, RTU-H)  
15`

Proteinase K (Dako, RTU)  
dil. 1:4 / 5`

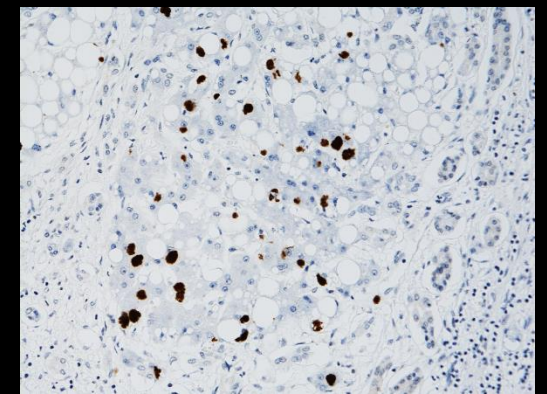
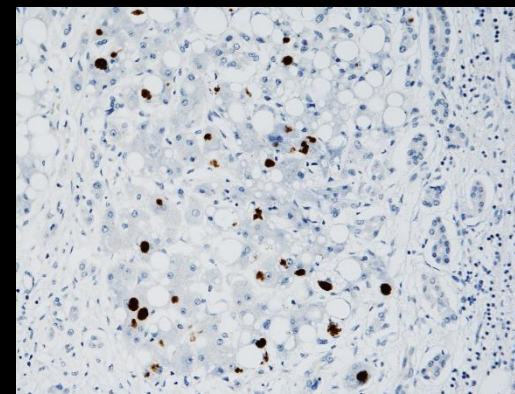
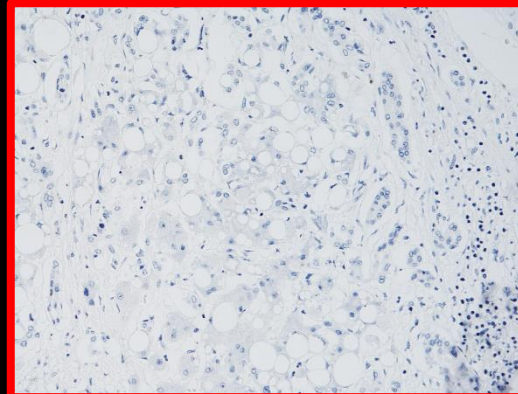
TCR- $\beta$ , 8A3, 1:200 RR

Tonsil



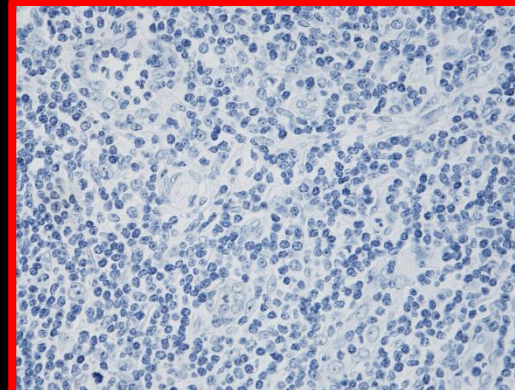
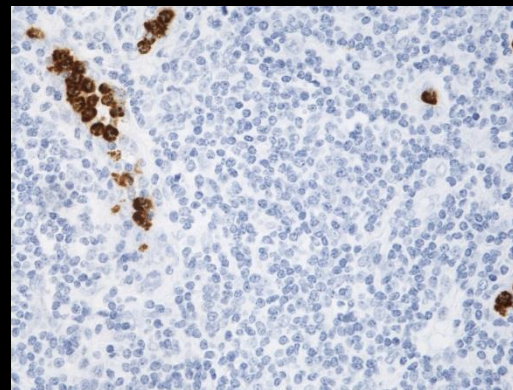
Ubiquitin, Ubi-1 1:750

Liver/ Mallory bodies



Neutrophil Elastase, NP57 1:1000

Tonsil





Proteolytic enzyme & digestion time ?

Tonsil NBF 48h

Digestion temp. 32°C

Proteinase K  
(RTU S3020, Dako)

Proteinase K dil. 1:4  
(RTU S3020, Dako)

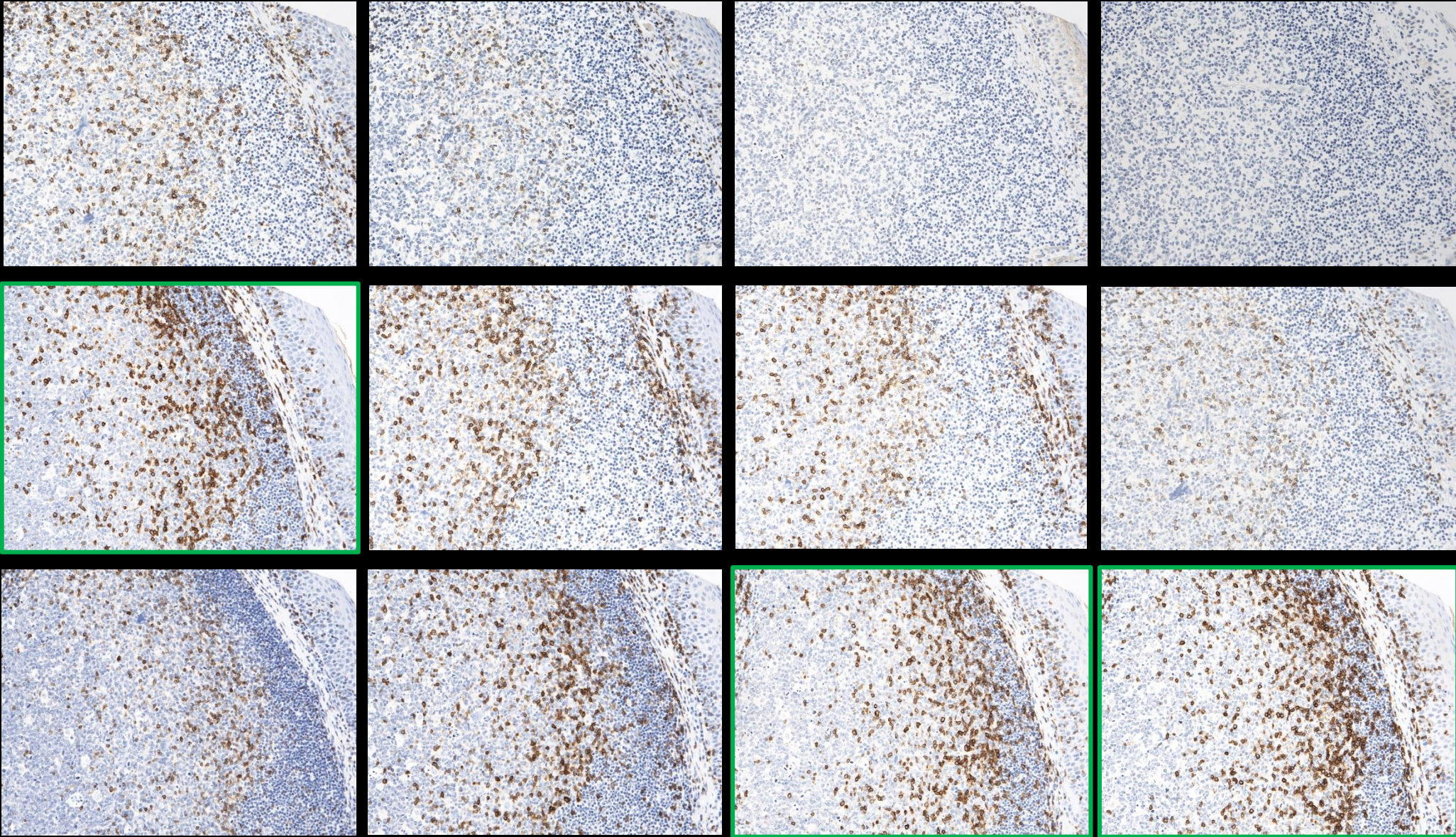
Trypsin  
(RTU, Biocare)

5 min

10 min

20 min

40 min



TCR-β clone 8A3 (1:200 RR) / Flex+ (Omnis)

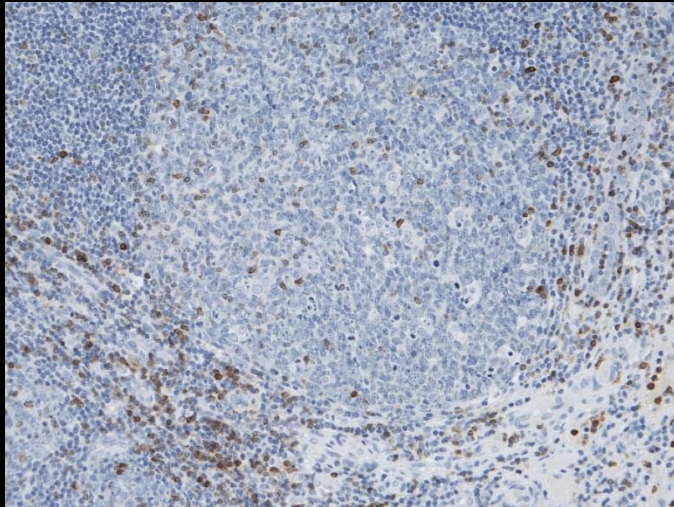


# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

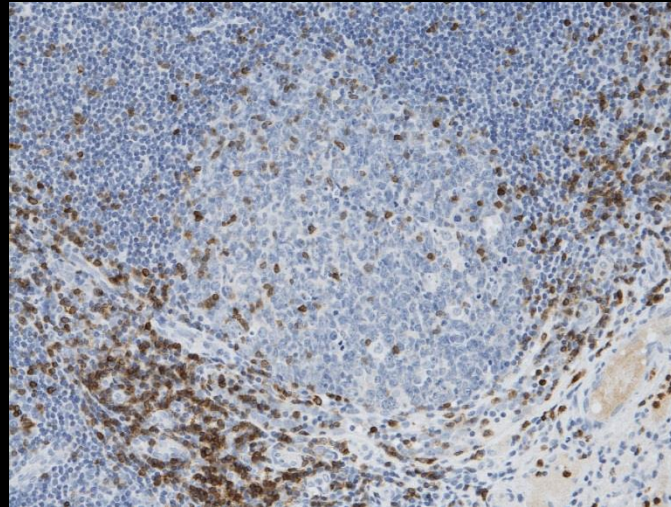
## Proteolytic enzyme & digestion temperature ?

TCR  $\beta$  clone 8A3 (1:200 RR) / Flex+ (Tonsil NBF 48h)

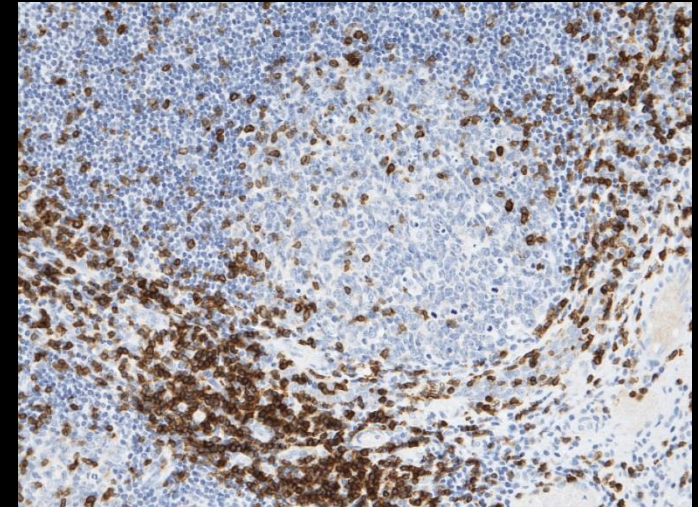
Trypsin Digestion temp. 4°C (10 `)



Trypsin Digestion temp. 24°C (10 `)



Trypsin Digestion temp. 37°C (10 `)



Increased intensity of TCR  $\beta$  positive T-cells



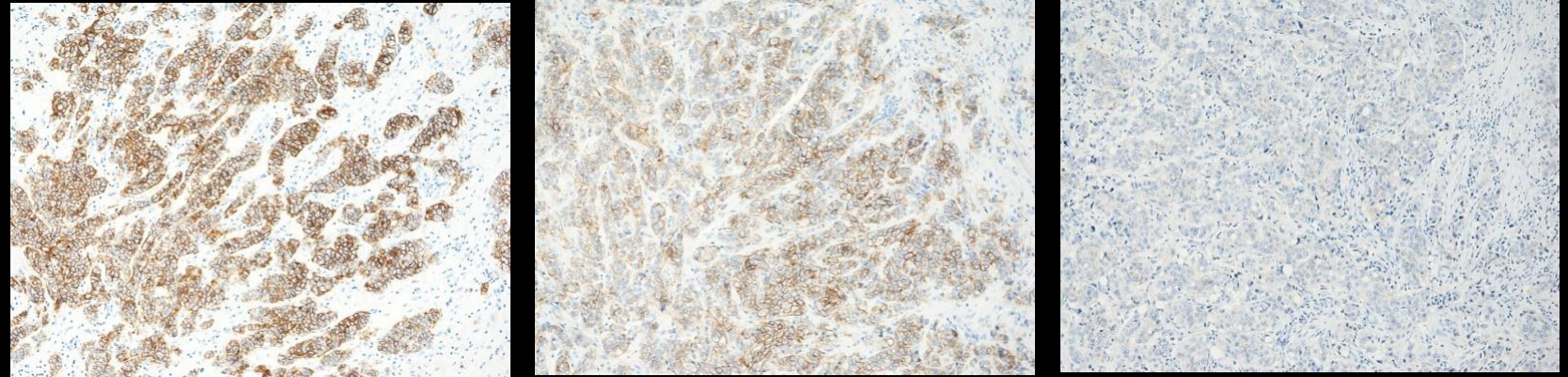
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Enzymatic digestion (Influence of fixation time)

EPCAM, clone MOC-31, dilution 1:20

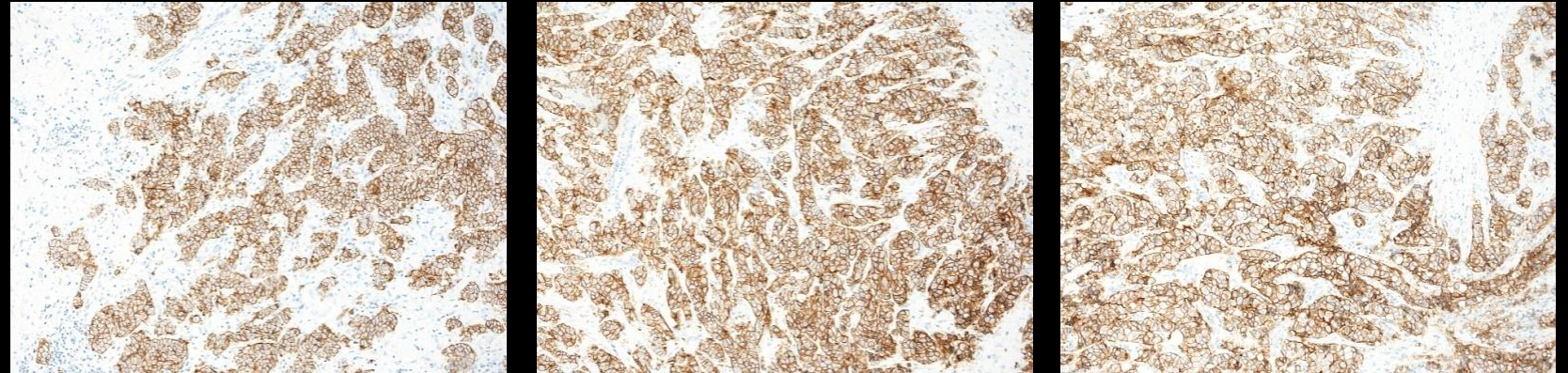
Pepsin / (Dako, S3002)

10 min/37°C



HIER , Low pH (S1700)

20 min / 97°C



NBF 24 h

NBF 48 h

NBF 120h

Adenocarcinoma (Breast) fixed in 10% Formalin

## Proficiency testing in immunohistochemistry—experiences from Nordic Immunohistochemical Quality Control (NordiQC)

Mogens Vyberg<sup>1,2</sup> · Søren Nielsen<sup>1</sup>

Virchows Arch (2016) 468:19–29

Table 3 Major causes of insufficient staining reactions

1. Less successful antibodies (17 %)
  - a. Poor antibodies<sup>a</sup>
  - b. Less robust antibodies<sup>b</sup>
  - c. Poorly calibrated RTUs
  - d. Stainer platform dependent antibodies
2. Insufficiently calibrated antibody dilutions (20 %)
3. Insufficient or erroneous epitope retrieval (27 %)
4. Error-prone or less sensitive visualization systems<sup>c</sup> (19 %)
- 5 Other (17 %)
  - a. Heat-induced impaired morphology
  - b. Proteolysis induced impaired morphology
  - c. Drying out phenomena
  - d. Stainer platform-dependant protocol issues
  - e. Excessive counterstaining impairing interpretation

<sup>a</sup> Consistently gives false negative or false positive staining or a poor signal-to-noise ratio in one or more assessment runs

<sup>b</sup> Frequently giving inferior staining results, e.g., due to mouse-anti-Golgi reactions or sensitive to standard operations as blocking of endogenous peroxidase

<sup>c</sup> Biotin-based detection kit for cytoplasmic epitopes, use of detection kits providing a too low sensitivity, or use of detection kits and chromogens giving imprecise localization of the staining signals complicating the interpretation

### Problems related to the choice of antigen retrieval method :

- Use of non-alkaline HIER buffer (low pH buffer)
- Use of inefficient / too short HIER period
- Use of no / enzymatic pre-treatment instead of HIER
- Use of excessive retrieval procedure → impaired morphology

**False positive or false negative results**



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Proteolytic pretreatment (excessive retrieval procedures)

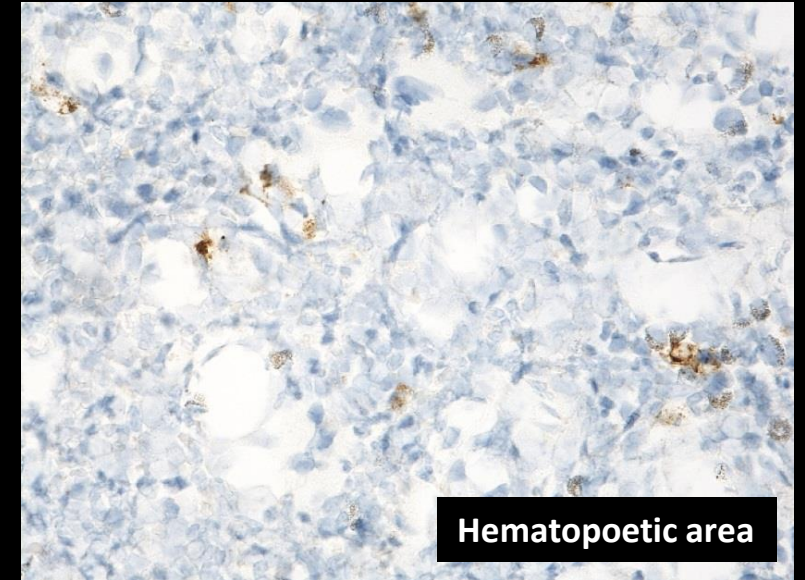
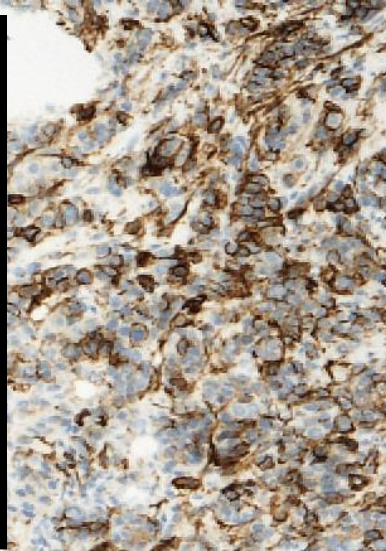
- over digestion (e.g. not calibrated to the fixation time in NBF)

## HIER (excessive retrieval procedures)

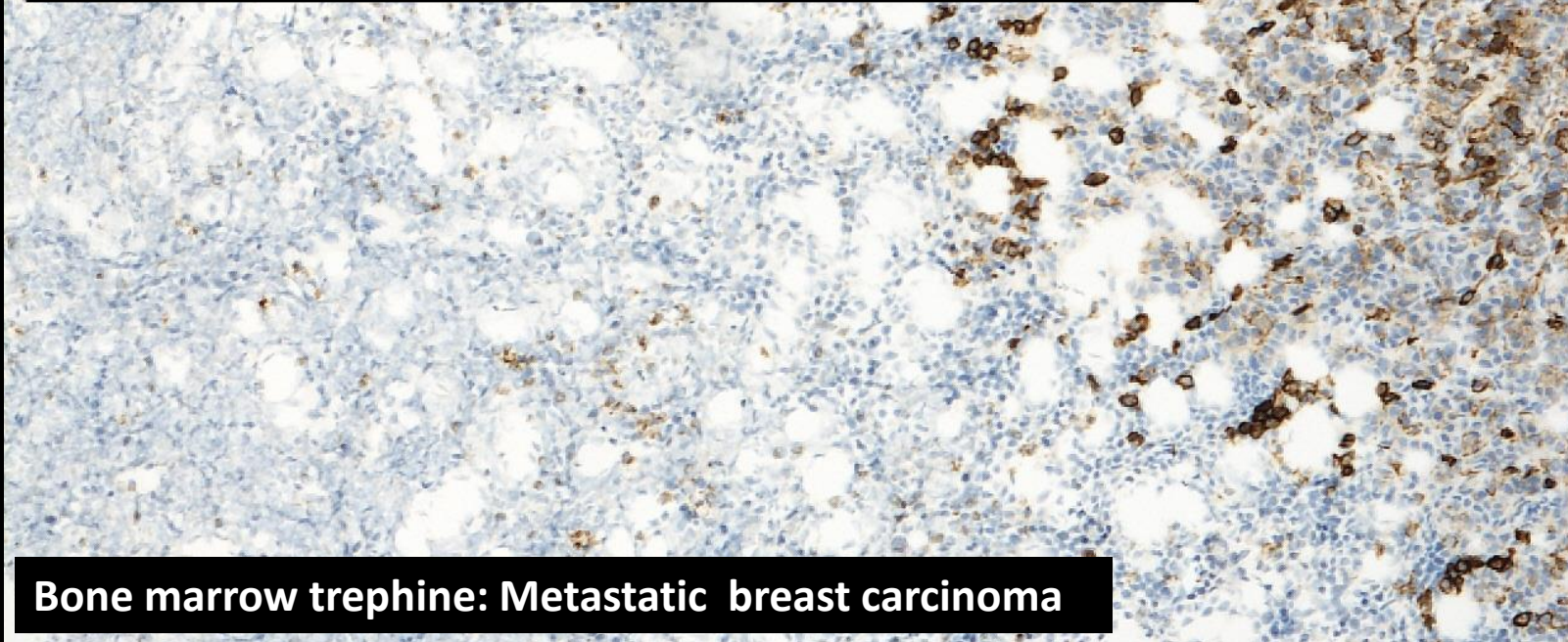
- too high temperature for too long time especially in alkaline retrieval buffers

## HIER applying standard procedures (excessive retrieval procedures)

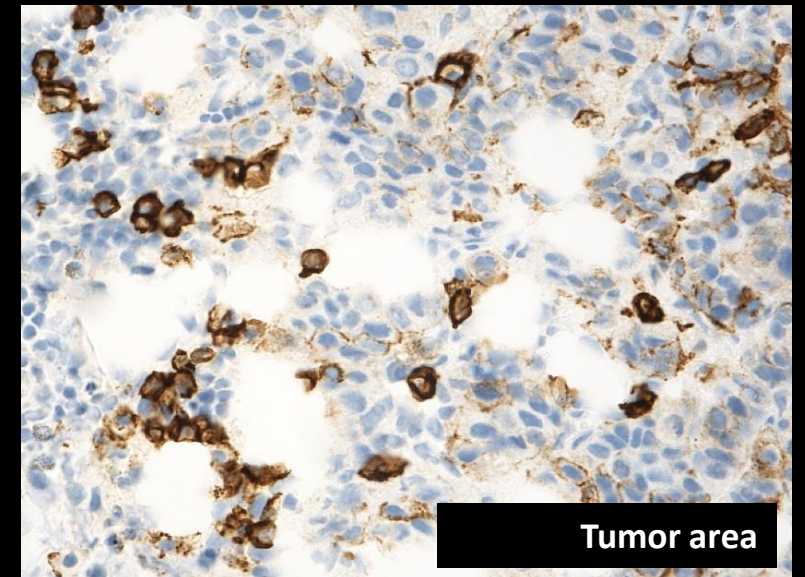
- fragile tissue/cell material (cell clot's )



Hematopoietic area



Bone marrow trephine: Metastatic breast carcinoma



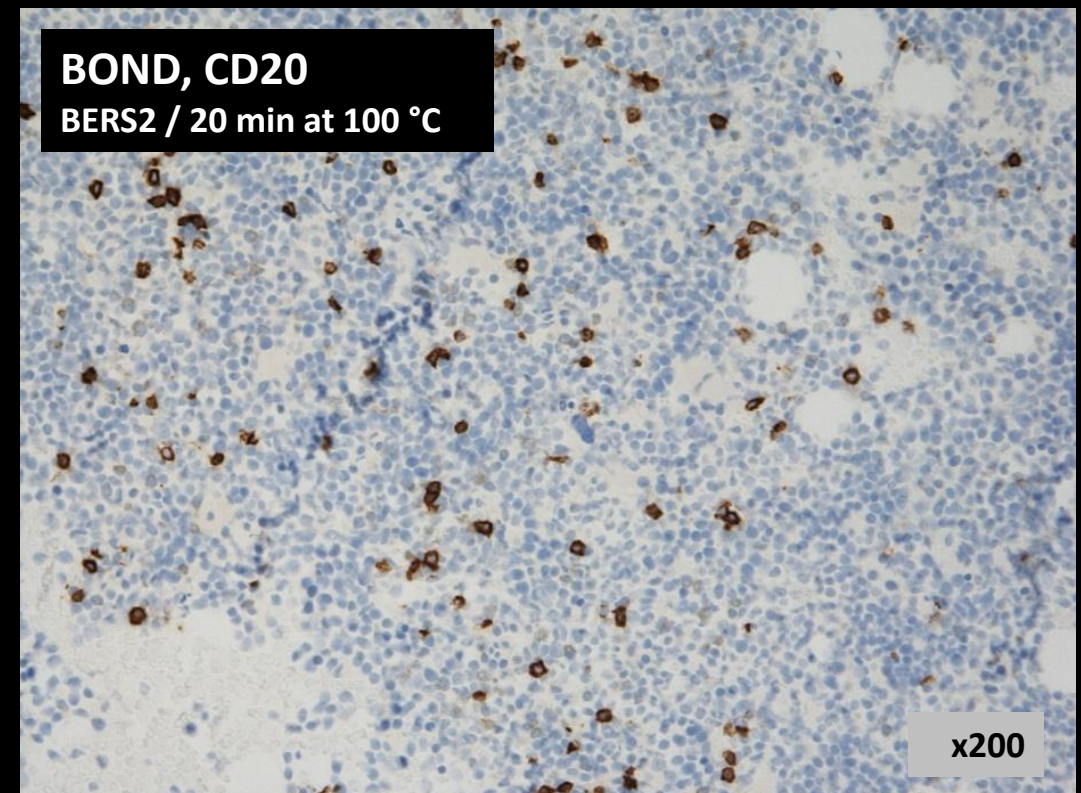
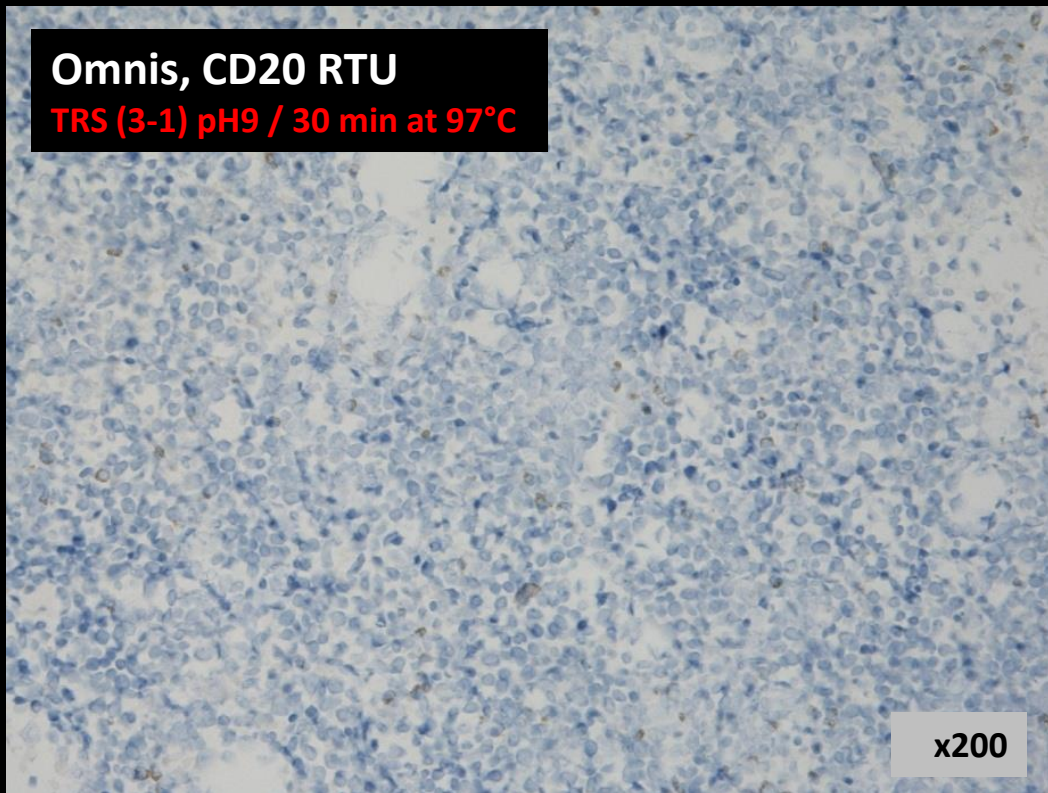
Tumor area



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## CD20 clone L26

Bone Marrow Coagulum/Clot (fixed for 24h in NBF)



HIER settings: Recommendations of the manufacturer`s



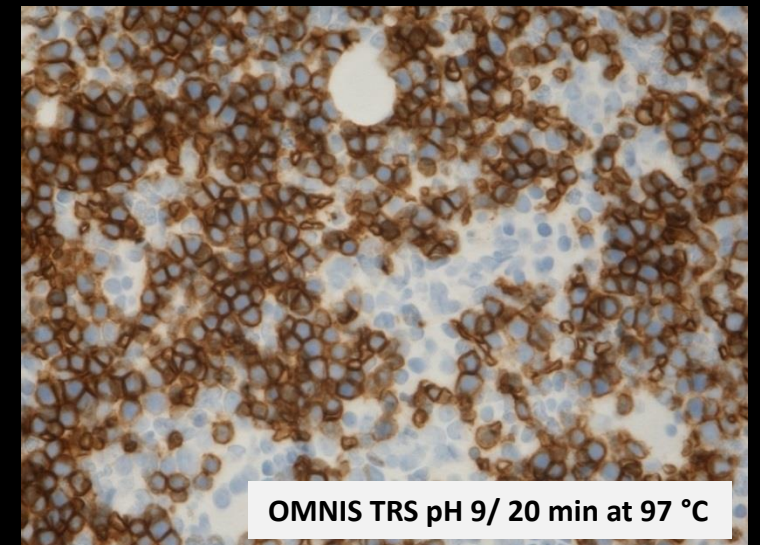
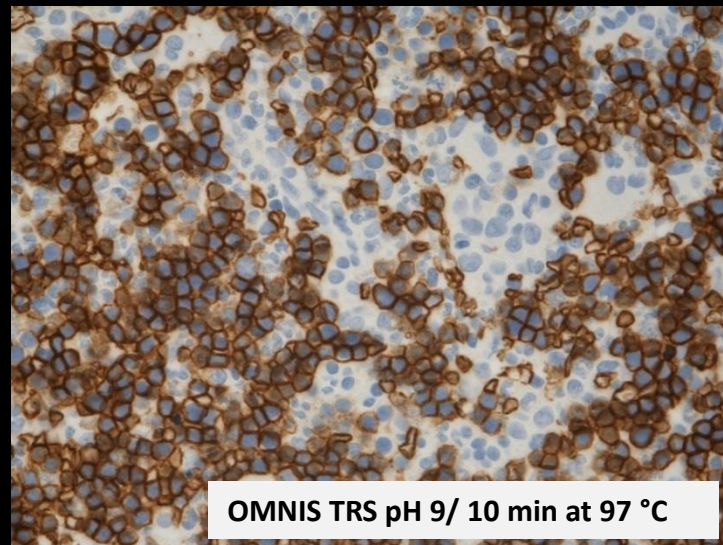
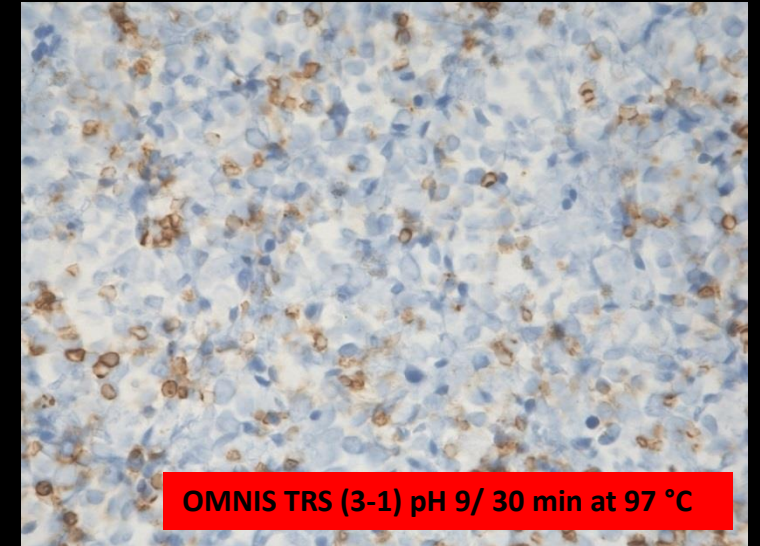
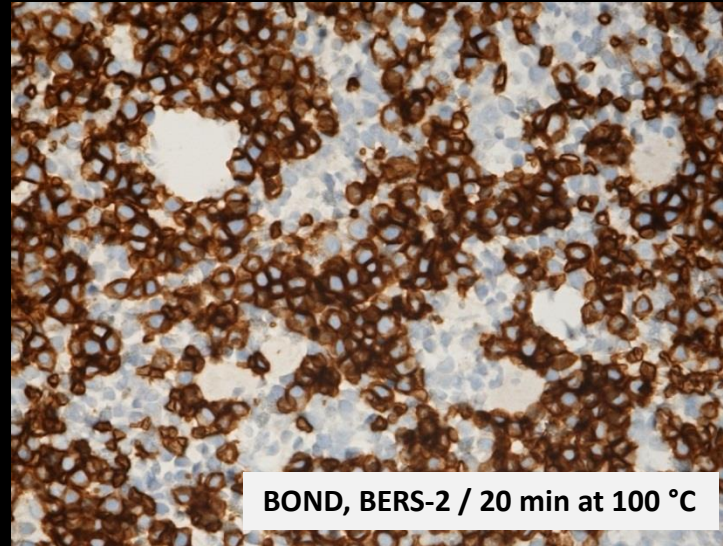
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Bone marrow clot (NBF 24h)

Morphology ?

Glycophorin A clone JC159 (1:500)

Flex+





# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Bone Marrow cloth (NBF 24 h)

Morphology ?

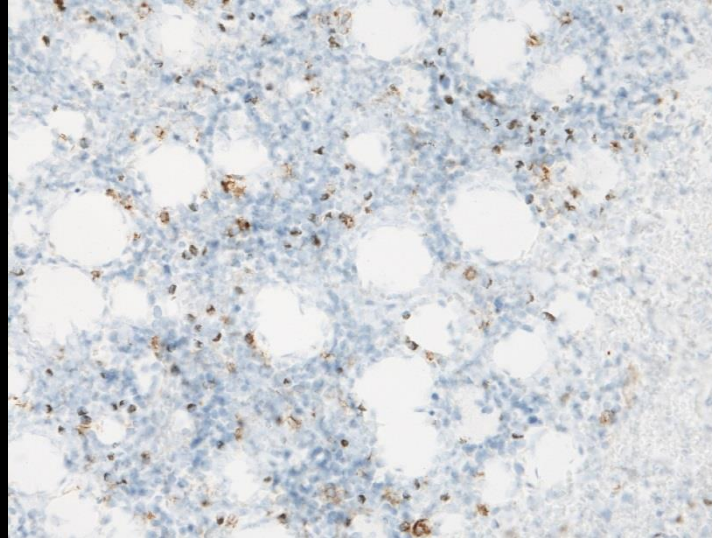
TRS (3-1) High pH 9, 24` at 97C,

Agilent/Dako (Omnis, Flex+)

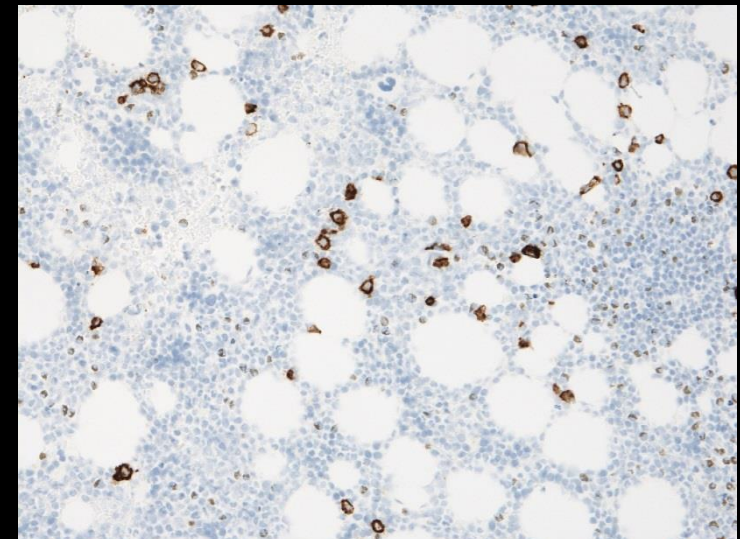
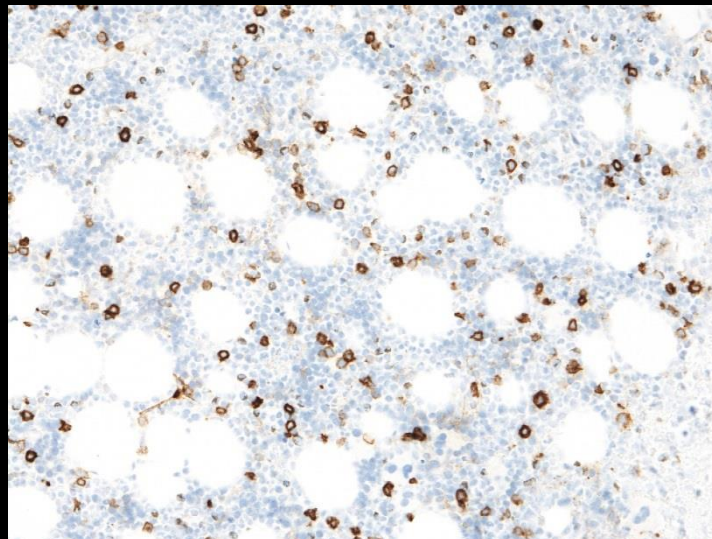
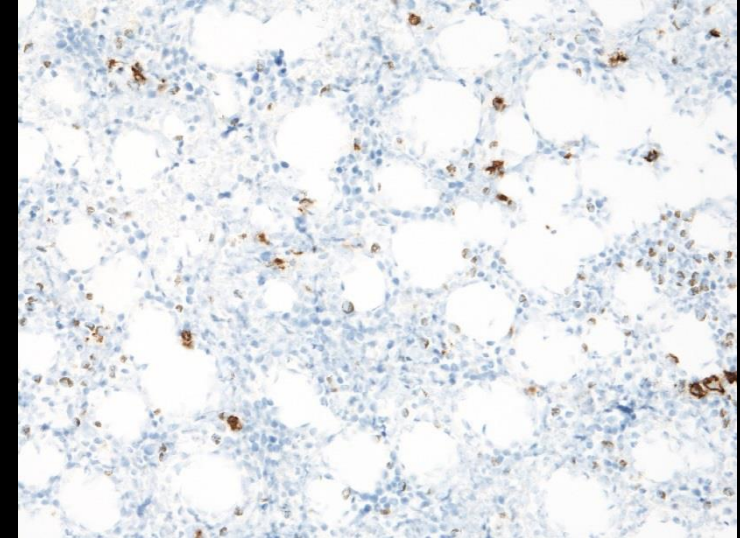
HIER buffer H, 24` at 97C

Thermo S./ LabVision (Omnis, Flex+)

CD117, EP10 (1:25 RR)



CD138, B-A38 (1:1000)



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Bone marrow clot

AS: PT-Link, High pH buffer's at 97°C / 20`

Morphology ?

High pH (3-1) (Dako)

Recommended settings:

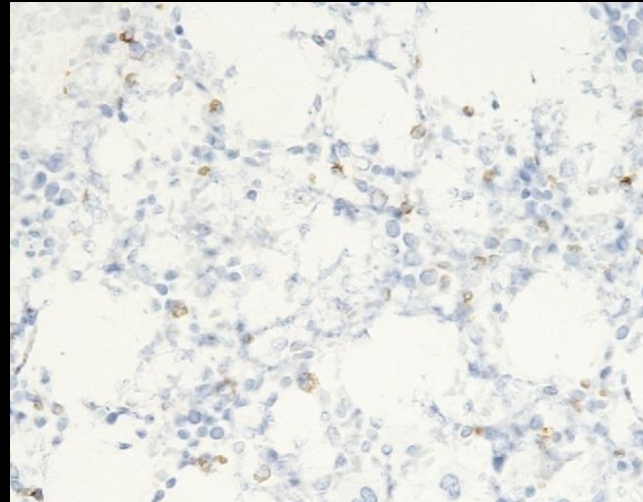
65°C

HIER buffer H (LabVision)

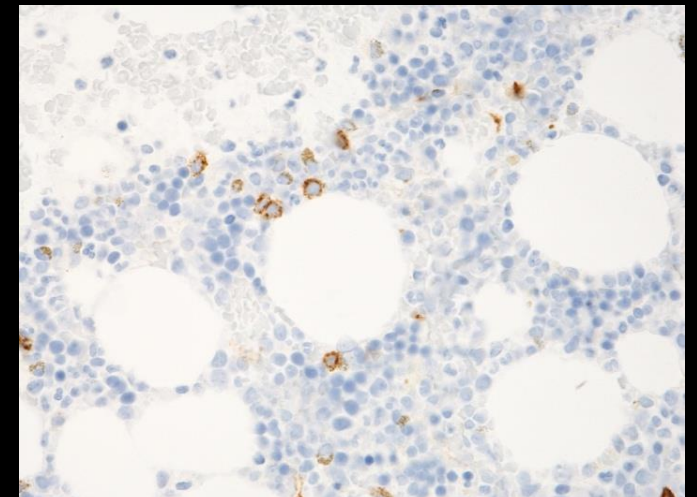
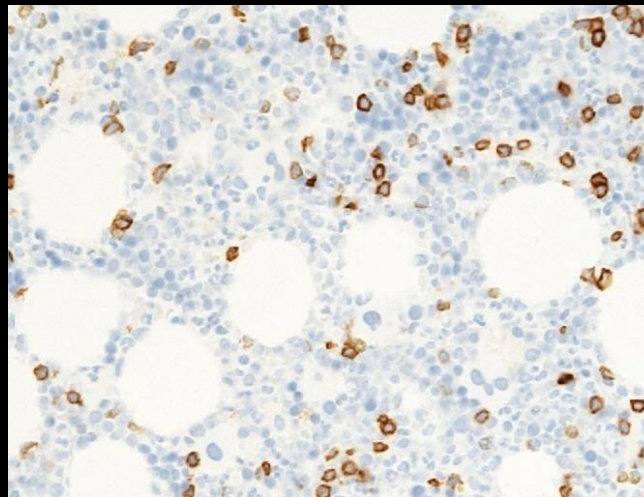
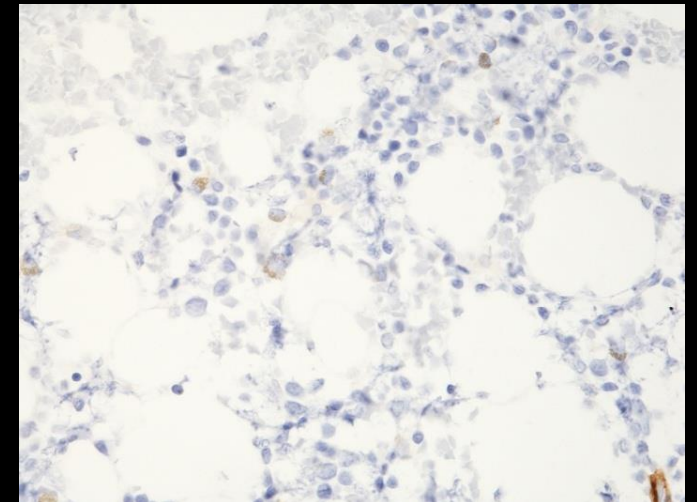
Recommended settings:

85°C

CD5 clone SP19



CD34 clone QBEND-10





# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Excessive retrieval on bone marrow material– what to do ?

- Optimize pre-analytic conditions

At arrival in our department , fresh fixative is added to the bone marrow specimens and a post fixation step for 24-48h at 42°C (stirring) has improved morphology.

Never dry cut slides horizontally – always vertically at room temperature for 30-60 min followed by 30-60 min at 60°C.

Decalcifying agent (shorten time and/or substitute ?)

- Shorten HIER time

Labor intensive: Require validated protocols for the each hematological marker and might require two different protocols depending on the material e.g., solid tissue (lymph nodes) versus bone marrow material.

- Use a more gentle HIER buffer (if possible)

- Consider change platform providing appropriate buffers not causing morphological problems as e.g., the TRS High pH buffer (Dako).





Pause

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Antibody company (producer) - Quote:

In the world of next generation immuno-oncology research, having confidence in your immunoassay results is vital.

Unfortunately, 75% of antibodies in today's market are non-specific or simply do not work at all.

## Primary Antibodies

Volume 57(1): 7-8, 2009  
Journal of Histochemistry & Cytochemistry  
<http://www.jhc.org>

### PERSPECTIVE

## Commercial Antibodies: The Good, Bad, and Really Ugly

John R. Couchman

Biomedicine Institute, University of Copenhagen, Biocenter, Copenhagen, Denmark

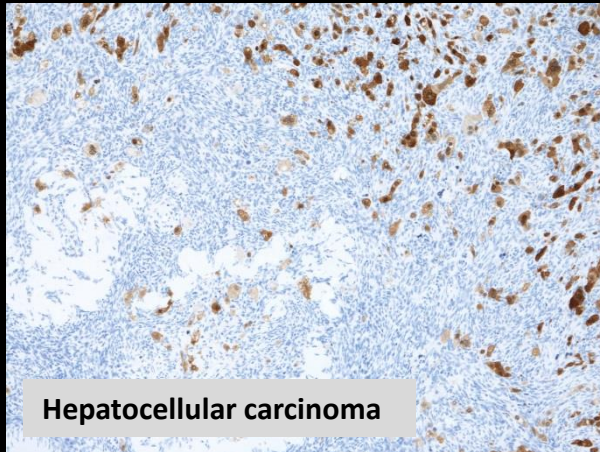
**SUMMARY** The range of antibodies available commercially grows ever larger. Perhaps as a consequence, quality control is not always what it could and should be. Investigators must be aware of potential pitfalls and take steps to assure themselves that the specificity of each antibody is as advertised. Additionally, companies should provide the necessary information about the antigen and antibody to investigators, including references, so that the appropriate controls can be included. (J Histochem Cytochem 57:7-8, 2009)



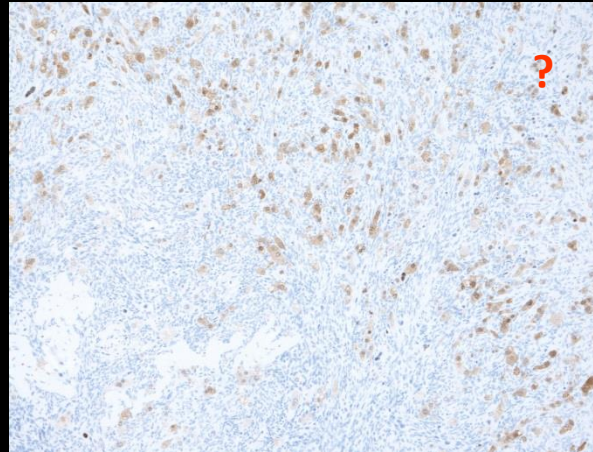
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Antibody choice: Specificity (different vendors)

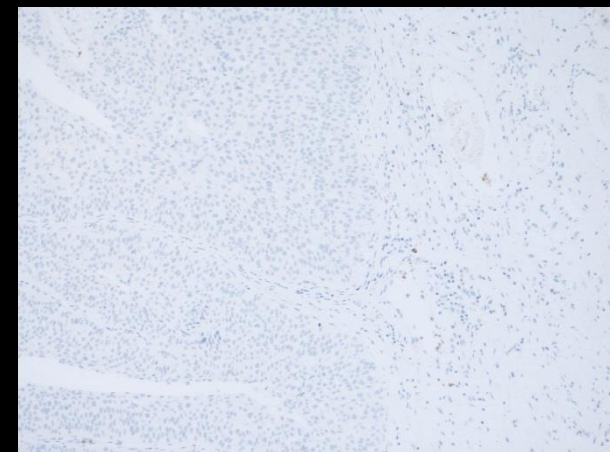
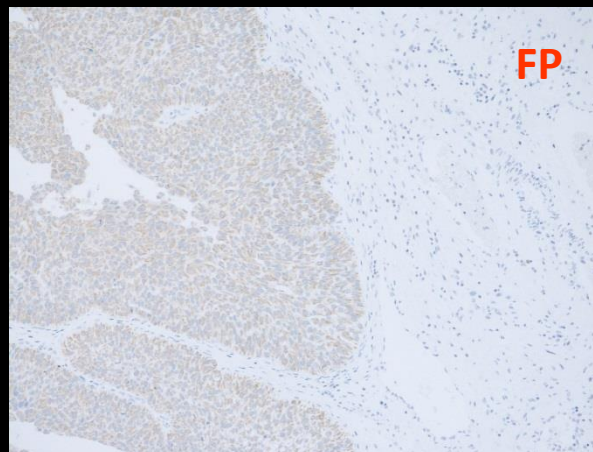
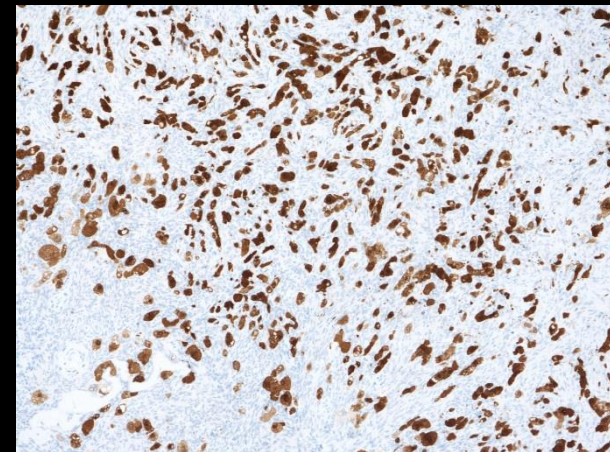
ARG1, SP156 Cell M 1:50



ARG1, SP156 Cell M 1:100



ARG1, SP156 Spring B. 1:25



## Proficiency testing in immunohistochemistry—experiences from Nordic Immunohistochemical Quality Control (NordiQC)

Mogens Vyberg<sup>1,2</sup> · Søren Nielsen<sup>1</sup>

Virchows Arch (2016) 468:19–29

Table 3 Major causes of insufficient staining reactions

1. Less successful antibodies (17 %)
  - a. Poor antibodies<sup>a</sup>
  - b. Less robust antibodies<sup>b</sup>
  - c. Poorly calibrated RTUs
  - d. Stainer platform dependent antibodies
2. Insufficiently calibrated antibody dilutions (20 %)
3. Insufficient or erroneous epitope retrieval (27 %)
4. Error-prone or less sensitive visualization systems<sup>c</sup> (19 %)
- 5 Other (17 %)
  - a. Heat-induced impaired morphology
  - b. Proteolysis induced impaired morphology
  - c. Drying out phenomena
  - d. Stainer platform-dependant protocol issues
  - e. Excessive counterstaining impairing interpretation

**37% insuff.**

<sup>a</sup> Consistently gives false negative or false positive staining or a poor signal-to-noise ratio in one or more assessment runs

<sup>b</sup> Frequently giving inferior staining results, e.g., due to mouse-anti-Golgi reactions or sensitive to standard operations as blocking of endogenous peroxidase

<sup>c</sup> Biotin-based detection kit for cytoplasmic epitopes, use of detection kits providing a too low sensitivity, or use of detection kits and chromogens giving imprecise localization of the staining signals complicating the interpretation

### Problems related to the choice and/or use of the primary antibody:

- **Inappropriate primary antibody**
  - Provide low sensitivity/specificity
- **Appropriate primary antibody**
  - Inapp. titre (too low or too high concentration)
- **Platform (stainer) dependent antibodies**
  - Provide low sensitivity / specificity

**False positive or false negative results**



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

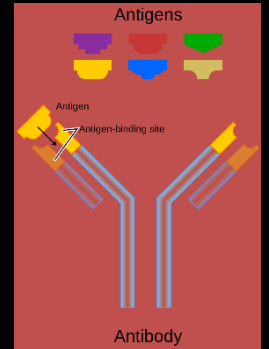
## Parameters affecting antibody-antigen reactions in tissue

**Antibody choice – Sensitivity/Specificity**

**Antibody Titer**

**Antibody performance related to the chosen automated platform**

**Antibody diluents**



**Incubation time**

**Incubation temperature**

**Sensitive to endogenous peroxidase blocking e.g., BCL6 (PG-B6p) and CD4 (1F6)**

**Storage of concentrated primary antibodies**

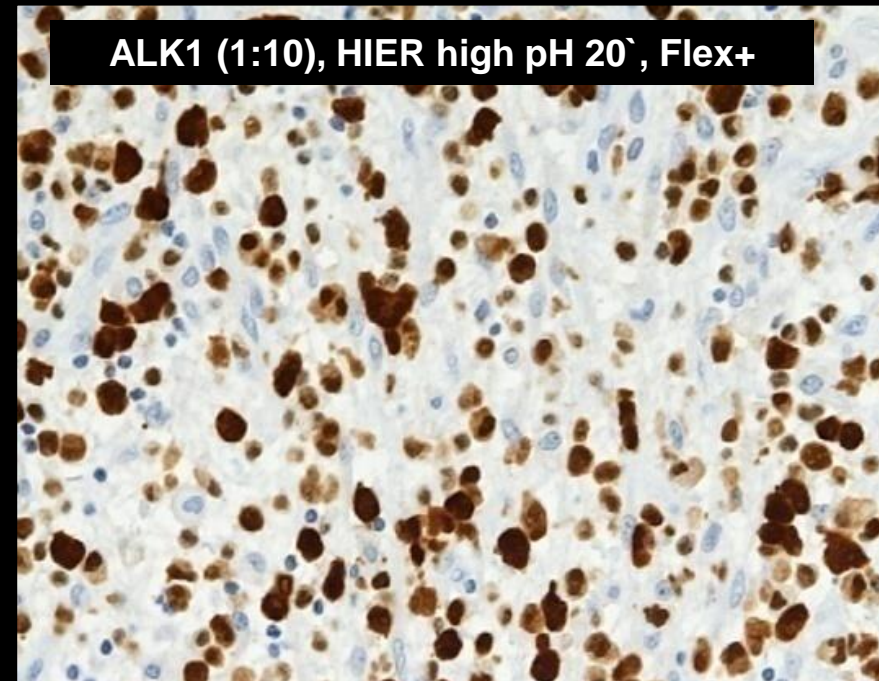
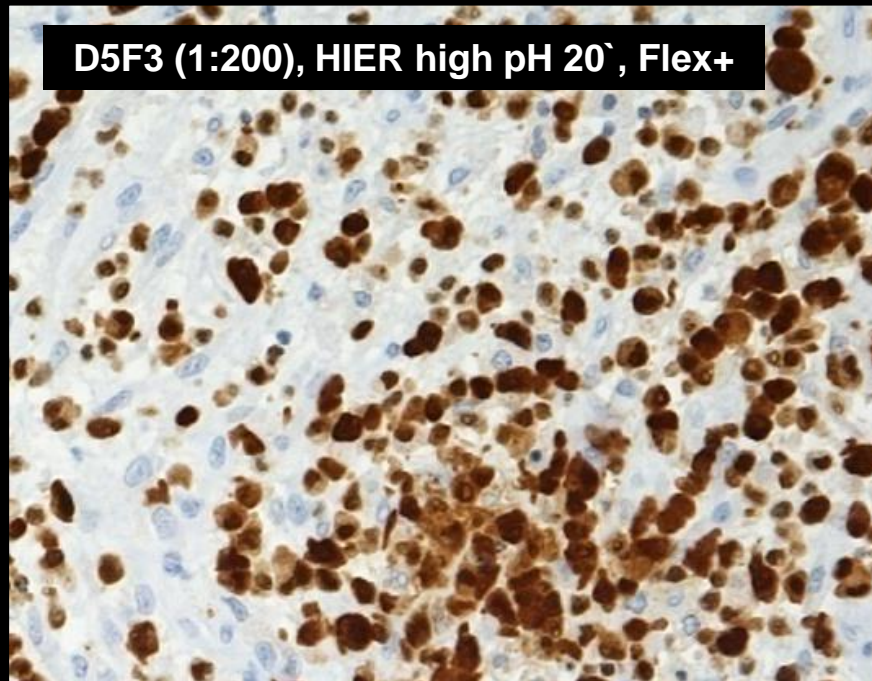
**Storage of diluted primary antibodies**

**Provided that efficient antigen retrieval has been performed and a sensitive detection system has been used**

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Anaplastic lymphoma kinase (ALK)

Anaplastic large cell lymphoma (ALCL) (ALK-NPM rearrangement)



Anything wrong ?



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## IHC-Type 2 marker

### Clinical Cancer Research



#### A Novel, Highly Sensitive Antibody Allows for the Routine Detection of *ALK*-Rearranged Lung Adenocarcinomas by Standard Immunohistochemistry

Mari Mino-Kenudson, Lucian R. Chirieac, Kenny Law, et al.

*Clin Cancer Res* 2010;16:1561-1571. Published OnlineFirst February 23, 2010.

## Lung tumors

Low concentration of ALK Fusion protein (e.g., EML-4/ALK) = require a sensitive antibody for detection

Intended use & “fit-for-purpose”

Human Pathology (2013) 44, 1656–1664



ELSEVIER

Human  
PATHOLOGY

[www.elsevier.com/locate/humpath](http://www.elsevier.com/locate/humpath)

Original contribution

#### Expression of anaplastic lymphoma kinase in Merkel cell carcinomas<sup>☆</sup>

Bettina Ekvall Filtenborg-Barnkob MD\*, Michael Bzorek HT\*

Department of Pathology, Naestved and Slagelse Hospital, Hospital South, Denmark

Received 30 August 2012; revised 13 November 2012; accepted 13 November 2012

## MCC

ALK,D5F3 = 94% pos

ALK,5A4 = 88% pos

ALK, ALK1 = 13% pos

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Table 1. Antibodies and assessment marks for ALK (lung), run 65

Concentrated antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff. <sup>1</sup>	OR <sup>2</sup>
mAb clone <b>5A4</b>	26	Leica Biosystems	8	9	14	4	49%	23%
	2	Monosan						
	1	Abcam						
	1	DBS						
	2	Biocare Medical						
mAb clone <b>OTI1A4*</b>	2	Zytomed Systems	16	6	0	0	100%	73%
	1	Invitrogen						
	19	Origene						
	1	Nordic Biosite						
	1	Cell Signaling						
mAb clone <b>IHC509</b>	1	GenomeMe	0	0	1	0	-	-
rmAb clone <b>D5F3</b>	19	Cell Signaling	7	9	3	0	84%	36%
rmAb clone <b>ALK1</b>	3	Dako/Agilent	0	0	0	4	-	-
	1	Cell Marque						
rmAb clone <b>QR017</b>	1	Quartett	0	1	0	0	-	-
rmAb clone <b>SP8</b>	1	BioGenex	0	0	0	1	-	-
rmAb clone <b>ZR305</b>	1	Zeta Corporation	0	0	1	0	-	-
Ready-To-Use antibodies								
mAb clone <b>5A4</b> <b>PA0306**/PA0831 (VRPS)<sup>3</sup></b>	2	Leica Biosystems	1	1	0	0	-	-
mAb clone <b>5A4</b> <b>PA0306*/PA0831 (LMPS)<sup>4</sup></b>	10	Leica Biosystems	4	3	2	1	70%	40%
mAb clone <b>5A4</b> <b>API3041</b>	1	BioCare	0	0	1	0	-	-
mAb clone <b>5A4</b> <b>CAM-0170</b>	1	Celnovte	0	1	0	0	-	-
mAb clone <b>5A4</b> <b>MAD-001720QD</b>	1	Master Diagnostica	0	0	1	0	-	-
mAb clone <b>ALK1</b> <b>GA641</b>	3	Dako/Agilent	0	0	0	3	-	-
mAb clone <b>ALK1</b> <b>IR641</b>	4	Dako/Agilent	0	0	0	4	-	-
mAb clone <b>ALK1</b> <b>790/800-2918 (LMPS)<sup>4</sup></b>	10	Ventana/Roche	1	0	1	8	10%	10%
mAb clone <b>137E9E8</b> <b>PA132</b>	1	Abcarta	0	0	0	1	-	-
mAb clone <b>OTI1A4 / 1A4</b> <b>8344-C010</b>	1	Sakura Finetek	1	0	0	0	-	-
mAb clone <b>OTI1A4 / 1A4</b> <b>GA785 (VRPS)<sup>3</sup></b>	12	Dako/Agilent	12	0	0	0	100%	100%
mAb clone <b>OTI1A4 / 1A4</b> <b>GA785 (LMPS)<sup>4</sup></b>	4	Dako/Agilent	4	0	0	0	-	-
rmAb clone <b>D5F3</b> <b>790-4794 (VRPS)<sup>3</sup></b>	73	Ventana/Roche	62	7	1	3	95%	85%
rmAb clone <b>D5F3</b> <b>790-4794 (LMPS)<sup>4</sup></b>	48	Ventana/Roche	36	9	3	0	94%	75%
rmAb clone <b>SP8</b> <b>RMPD007</b>	1	Diagnostic BioSystems	0	0	0	1	-	-
Total	256		152	46	28	30		
Proportion			59%	18%	11%	12%	77%	

1) Proportion of sufficient stains (optimal or good) (≥5 assessed protocols).

2) Proportion of Optimal Results (≥5 assessed protocols).

3) Vendor Recommended Protocol Settings (VRPS) to a specific RTU product applied on the vendor recommended platform(s) (≥5 assessed protocols).

4) Laboratory Modified Protocol Settings (LMPS) to a specific RTU product (≥5 assessed protocols).

\*) OTI1A4 is called 1A4 by some vendors

\*\*) Product no. PA0306 has been terminated and replaced by PA0831.

NQC Run 45, 51, 57 and 65 (ALK Lung)

87 protocols were based on clone ALK1:

Only two protocols (2%) were assessed as sufficient

NQC Run 65 (assessment spring/summer 2022)

1/21 protocols were assessed as sufficient

Don't use clone ALK1 to detect ALK rearranged lung adenocarcinomas

It does not "fit-for-purpose"

**D5F3, OTI1A4, 5A4**

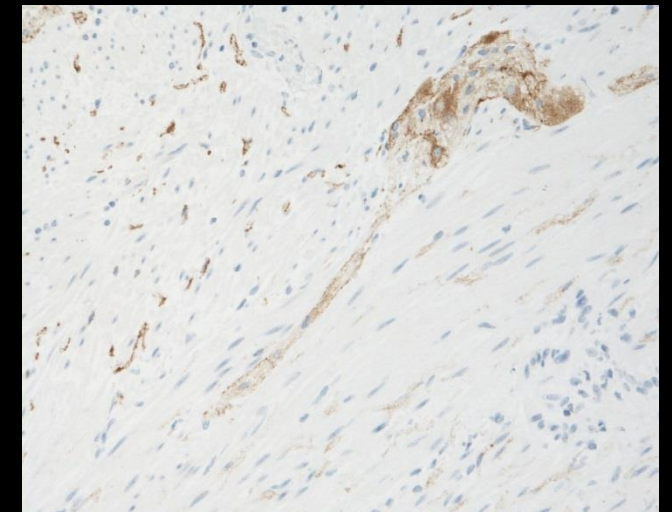
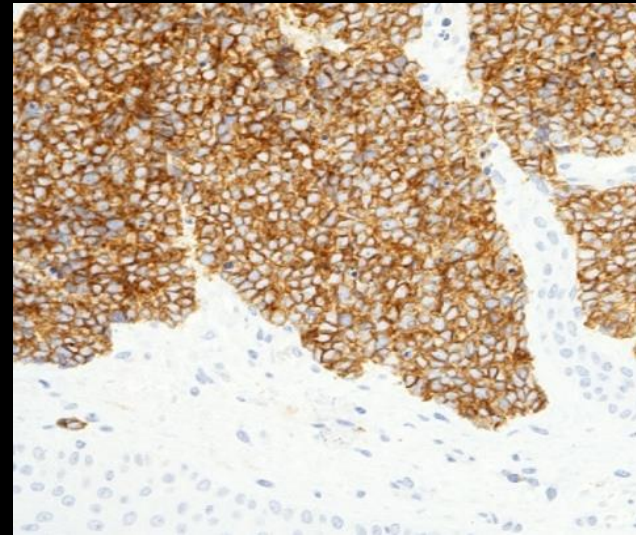
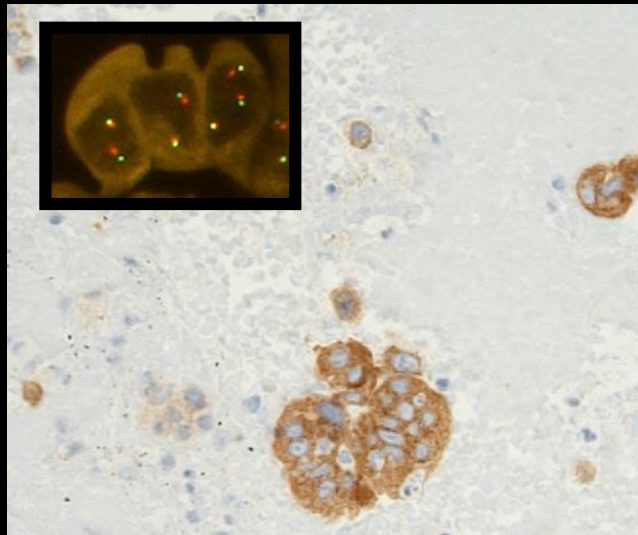


Lung AC (ALK-EML4)

MCC (Skin)

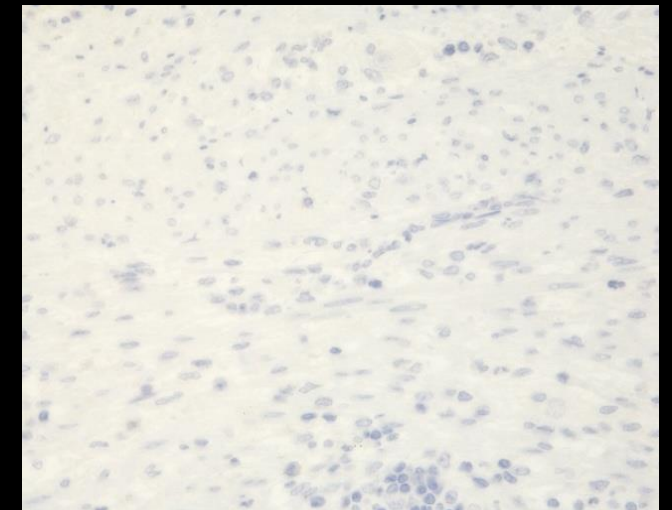
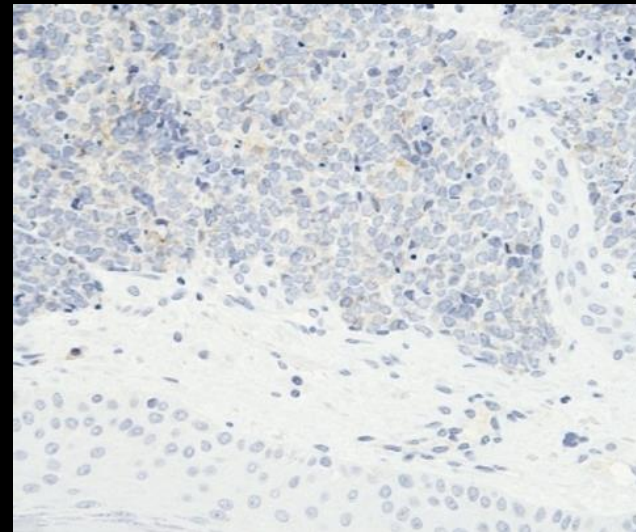
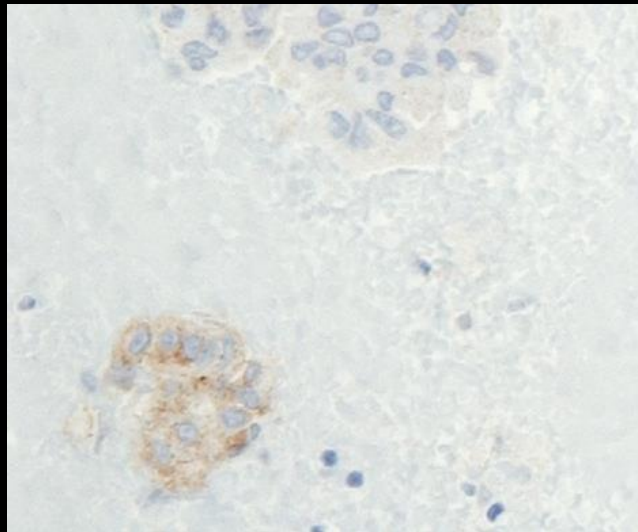
Appendix

ALK, D5F3 (1:200)



Ganglion and peripheral nerve cells

ALK, ALK1 (1:10)



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Problem: Primary antibody provides low sensitivity

## URO II/III

Table 1. Antibodies and assessment marks for URO II/III, run 59

Concentrated antibodies	Reactivity	n	Vendor	Optimal	Good	Borderline	Poor	Suff. <sup>1</sup>	OR <sup>2</sup>
mAb <b>BC21</b>	URO II	24	BioCare Medical	13	10	3	1	85%	37%
		3	Zytomed Systems						
mAb <b>AU-1</b>	URO III	2	Cell Marque	-	-	-	2	-	-
rmAb <b>SP73</b>	URO III	7	Cell Marque	-	-	1	7	0%	0%
		1	Immunologic						
rmAb <b>ERP18799</b>	URO II	1	ABCAM	-	1	-	-	-	-
rmAb <b>EP321</b>	URO III	1	Bio SB	-	-	1	-	-	-
rpAb <b>AB82173</b>	URO III	2	ABCAM	-	-	-	2	-	-
Ready-To-Use antibodies									Suff. <sup>1</sup> OR <sup>2</sup>
mAb <b>BC21</b>	URO II	1	Biocare Medical	-	1	-	-	-	-
<b>AVI 3051 KG</b>									
mAb <b>BC21</b>		1	Biocare Medical	-	1	-	-	-	-
<b>API 3051 AA</b>									
mAb <b>BC21</b>		1	Vitro SA	1	-	-	-	-	-
<b>MAD-000773QD</b>									
mAb <b>BC21</b>	URO II/III	1	Zytomed Systems	-	1	-	-	-	-
<b>MSG102</b>									
mAb <b>BC21+BC17</b>		3	Biocare Medical	-	2	1	-	-	-
<b>API 3094 AA</b>									
rmAb <b>SP73</b>		1	Roche/Ventana	-	-	1	-	-	-
<b>760-4533 (VRPS)<sup>3</sup></b>	URO III								
rmAb <b>SP73</b>		16	Roche/Ventana	-	-	2	14	0%	0%
<b>760-4533 (LMPS)<sup>4</sup></b>									
rmAb <b>SP73</b>	URO III	1	Cell Marque	-	-	-	1	-	-
<b>345R-17/18</b>									
Total		66		14	16	9	27		
Proportion				21%	24%	14%	41%	45%	

1) Proportion of sufficient stains (optimal or good). (≥5 assessed protocols).

2) Proportion of Optimal Results (≥5 assessed protocols).

3) Vendor Recommended Protocol Settings (VRPS) to a specific RTU product applied on the vendor recommended platform(s) (≥5 assessed protocols).

4) Laboratory Modified Protocol Settings (LMPS) to a specific RTU product (≥5 assessed protocols).

The problem:

Abs raised exclusively against Uroplakin III show too low analytical sensitivity - detecting urothelial cancers among carcinomas of unknown origin.



# Comparison of Antibodies to Detect Uroplakin in Urothelial Carcinomas

Heidi L. Kristoffersen, BMS, Rasmus Røge, MD, PhD, and Søren Nielsen, BMS

(*Appl Immunohistochem Mol Morphol* 2022;30:326–332)

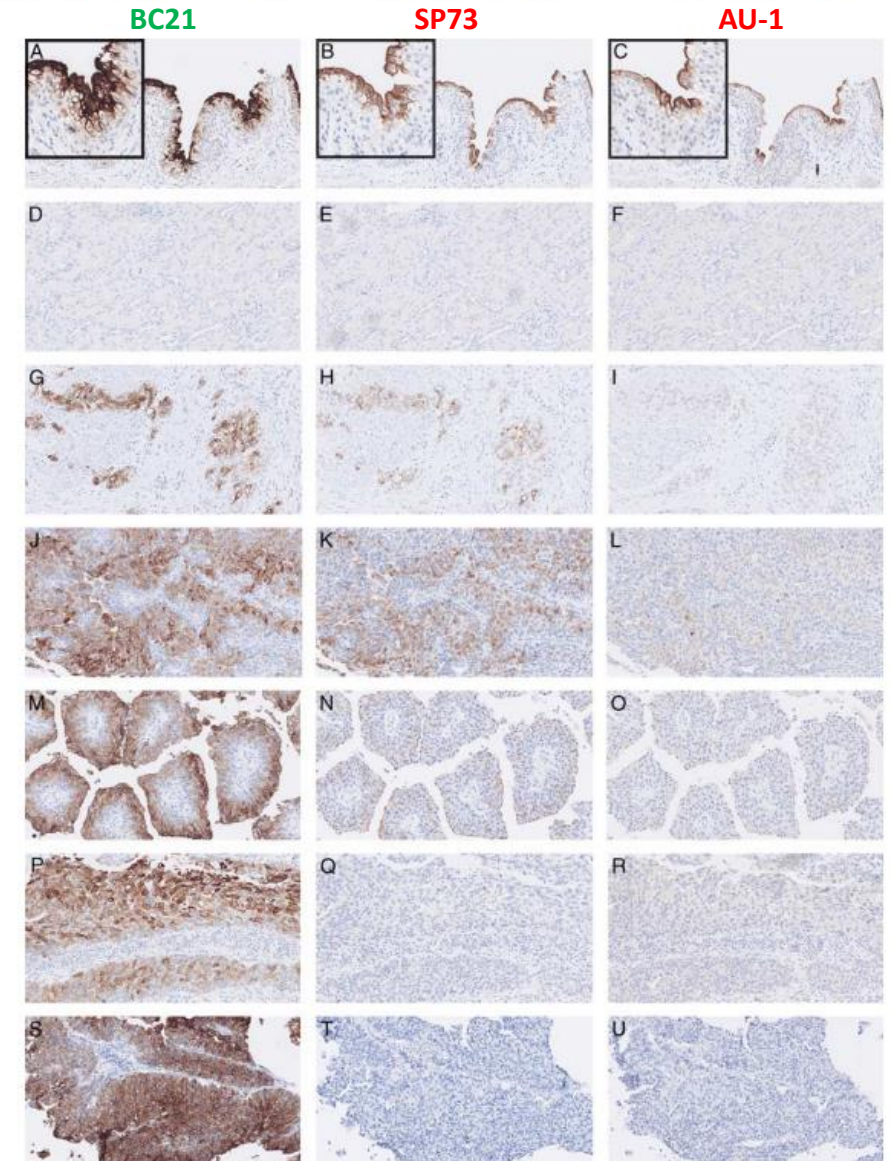
TABLE 3. Results, H-scores

	mmAb BC21 Uroplakin II			rmAb SP73 Uroplakin III			mmAb AU-1 Uroplakin III		
	n	n%	Mean H-score	n	n%	Mean H-score	n	n%	Mean H-score
Urothelial carcinomas (n = 58)									
Positive total	40	69	72	17	29	19	11	19	4
High-expressor (H-score 150-300)	8	13	182	0	0	0	0	0	0
Medium-expressor (H-score 10-149)	23	38	60	8	13	36	2	3	10
Low-expressor (H-score 1-9)	9	15	6	9	15	3	9	15	3
Negative (H-score <1)	18	31	0	41	71	0	47	81	0
Nonurothelial carcinomas (n = 111)									
Positive total	3	3	18	0	0	0	0	0	0
High-expressor (H-score 150-300)	0	0	0	0	0	0	0	0	0
Medium-expressor (H-score 10-149)	2	2	27	0	0	0	0	0	0
Low-expressor (H-score 1-9)	1	1	2	0	0	0	0	0	0
Negative (H-score <1)	108	97	0	111	100	0	111	100	0

mmAb indicates mouse monoclonal antibody; rmAb, rabbit monoclonal antibody.

**Demonstrated that UPII klon BC21 outperforms UPIII Abs and at present should be the preferred choice of UP marker.**

**However, should be used in a panel with other Urothelial markers as e.g., GATA3 due to moderate analytical sensitivity (and due to positivity in a minority of nonurothelial carcinomas - app. 3%)**



**FIGURE 1.** Examples of UP staining in normal tissue and UCs with various intensity using UPII mmAb clone BC21 (A, D, G, J, M, P, S), UPIII rmAb clone SP73 (B, E, H, K, N, Q, T), and UPIII mmAb clone AU-1 (C, F, I, L, O, R, U). A–C: normal urethra, insert high powerfield. D–F: normal kidney. G–I, J–L, M–O, P–R, and S–U: different UCs. mmAb indicates mouse monoclonal antibody; rmAb, rabbit monoclonal antibody; UCs, urothelial carcinomas; UP, uroplakin.

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Problem: Primary antibody provides low sensitivity

## CK8/18

Table 1. Antibodies and assessment marks for CK8/18, run 57

Concentrated antibodies	reactivity	n	Vendor	Optimal	Good	Borderline	Poor	Suff. <sup>1</sup>	Suff. OPS <sup>2</sup>
mAb clone <b>34betaH11</b>	CK8	1	Diagnostic BioSystems	0	0	1	0	-	-
mAb clone <b>5D3</b>	CK8/18	16	Leica/Novocastra	5	6	4	10	44%	-
		3	Diagnostic BioSystems						
		2	Biocare						
		2	Thermo Scientific						
		1	DCS						
		1	Monosan						
mAb clones <b>B22.1/B23.1</b>	CK8/18	11	Cell Marque	8	3	2	1	79%	92%
		1	Bio SB						
		1	Immunologic						
		1	Menarini						
mAb clone <b>C51</b>	CK18	1	Zymed	0	0	0	1	-	-
mAb clone <b>CAM5.2</b>	CK8(7)	3	Zytomed	0	1	2	0	-	-
mAb clone <b>CY90</b>	CK18	1	Nordic Mubio	0	0	1	1	-	-
		1	Sigma						
mAb clone <b>DC10</b>	CK18	11	Agilent/Dako	0	0	9	7	0%	-
		3	Thermo Scientific						
		1	Biocare						
		1	Immunologic						
mAb clone <b>TS1</b>	CK8	2	Thermo Scientific	0	2	1	0	-	-
		1	Leica/Novocastra						
mAb clone <b>TS1+DC10</b> ("homebrew")	CK8/18	1	Thermo + Dako	1	0	0	0	-	-
rmAb clone <b>BSR15</b>	CK8	1	Nordic Biosite	1	0	0	0	-	-
rmAb clone <b>EP17</b>	CK8	4	Epitomics	4	0	1	0	-	-
		1	Cell Marque						
rmAb clones <b>EP17/EP30</b>	CK8/18	9	Dako/Agilent	9	1	0	0	100%	-
		1	Cell Marque						
Ready-To-Use antibodies									
mAb clone <b>DC10</b> <b>IR618/IS618</b>	CK18	5	Agilent/Dako	0	1	3	1	-	
mAb clone <b>DC10</b> <b>GA618</b>	CK18	9	Agilent/Dako	0	2	6	1	-	

The problem:

Abs raised exclusively against CK18 may show partial or complete loss of CK18 in 25% of breast carcinomas depending on their histological type (included in this run 57).



# NordiQC assessment CK 8/18

EP17/EP30 = CK8/18

DC10 = CK18 only

Abs raised exclusively against CK18 may show partial or complete loss of CK18 in 25% of breast carcinomas depending on their histological subtype

EP17/EP30

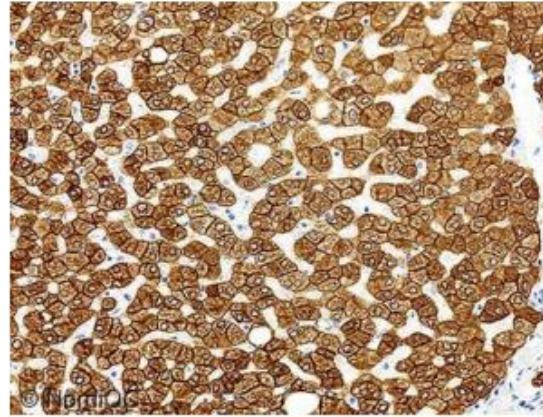


Fig. 3a  
Optimal CK8/18 staining of the liver using the rmAb clone cocktail EP17/EP30 as a concentrate in an optimally calibrated LD assay, performed on the Dako Omnis. The vast majority of hepatocytes show a distinct, moderate staining reaction with a membrane enhancement. Compare with Figs. 4a – 6a, same protocol.

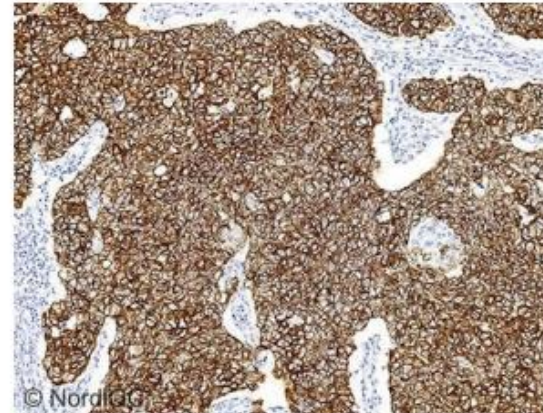


Fig. 5a  
Optimal CK8/18 staining of the breast ductal carcinoma using same protocol as in Figs. 3a - 4a. Virtually all neoplastic cells show a strong and distinct cytoplasmic staining reaction. The use of a cocktail of CK8 and CK18 antibodies secures optimal staining reaction despite the apparent loss of CK18 in the tumour. Compare with Fig. 5b – same field.

DC10

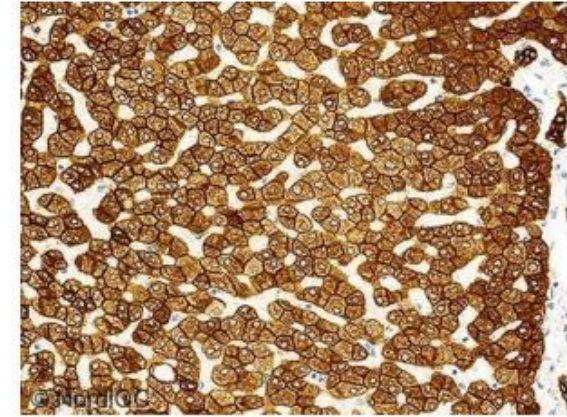


Fig. 3b  
Optimal CK8/18 staining of the liver using an insufficient protocol based on the mAb clone DC10. Clone DC10 reacts with CK18 and in this (DC10) CK18-protocol the vast majority of hepatocytes show a distinct, moderate staining reaction with a membrane enhancement. Compare with Figs. 4b – 6b, same protocol. Compare also with Fig. 3a – same field.

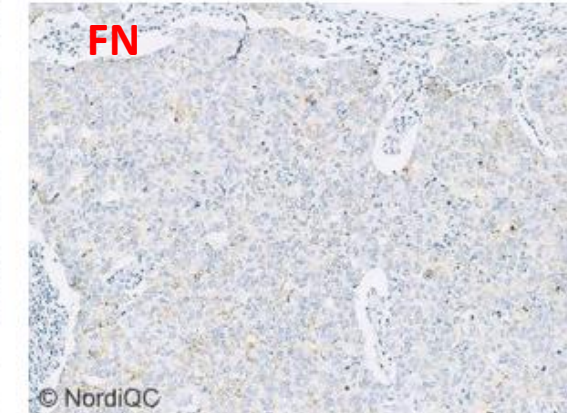


Fig. 5b  
Insufficient CK8/18 staining of the breast ductal carcinoma using same protocol as in Figs. 3b - 4b. Despite producing optimal staining in liver (and the SCLC), virtually all neoplastic cells are unstained. This tumour belongs to the approx. 25% of breast carcinomas that shows partial or complete loss of CK18 expression, making CK18 antibodies like clone DC10 less suitable identifying non-squamous carcinomas including adenocarcinoma of unknown origin. Compare with Fig. 5a – same field.



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

**Problem: Primary antibody provides low specificity and/or poor signal-to-noise ratio**

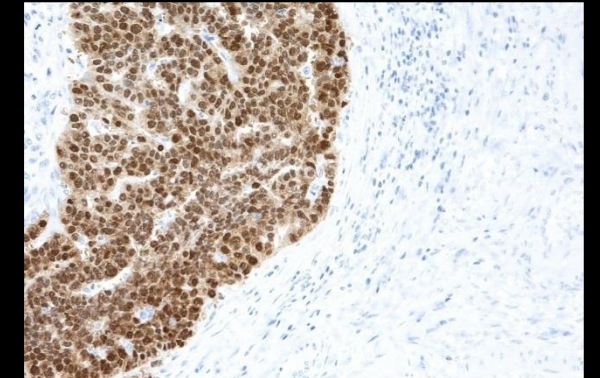
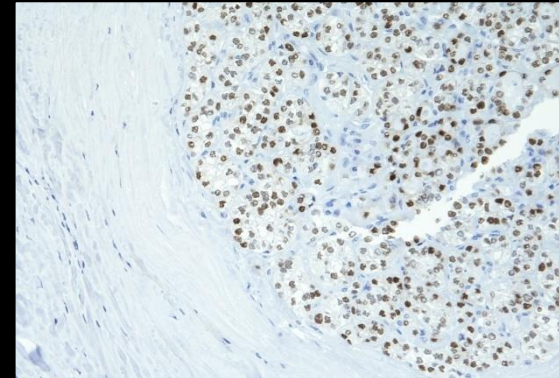
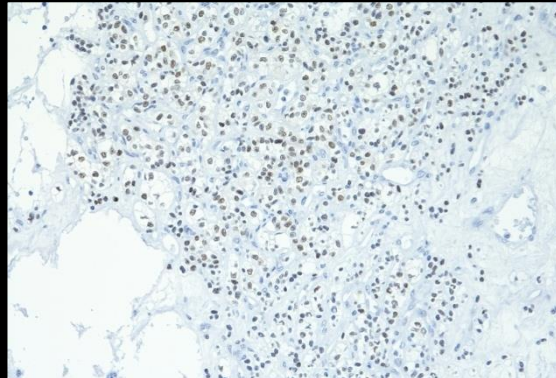
**Which antibody ?**

Renal Cell Carcinoma (CC)

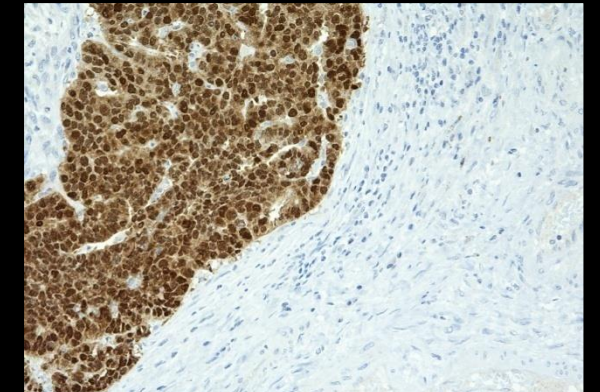
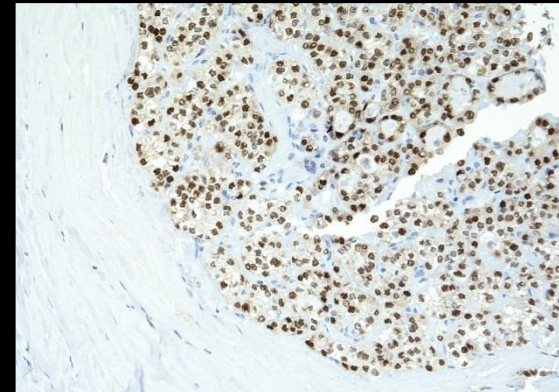
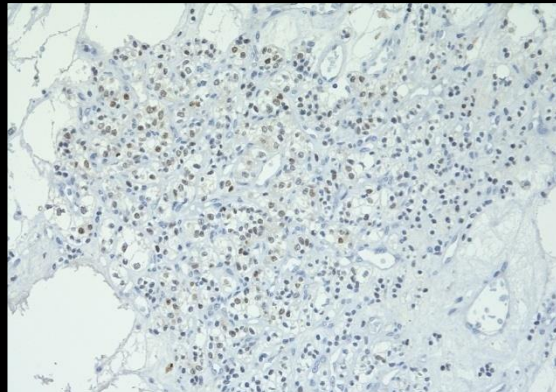
Thyroid Carcinoma (Pa)

Ovary Carcinoma (Se)

Pax-8, MRQ-50 (1:2000)  
HIER High pH/Flex+ (AS)



Pax-8, BC12 (1:150) HIER  
High pH/Flex+ (AS)





# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

**Problem: Primary antibody provides low specificity and/or poor signal-to-noise ratio**

***Liau J-Y et al.: Appl Immunohistochem Mol Morphol . 2016 Jan;24(1):57-63***

**Demonstrated that neuroendocrine tumors (NET's) from a large variety of organs were immuno-reactive with the two less specific antibodies (pAb Proteintech & mAb MRQ-50) - cross-reacting with other PAX proteins**

**Also, all NET's were immuno-negative with the two monoclonal antibodies raised against the C-terminal part of PAX8 protein (PAX8R1 & BC12)**

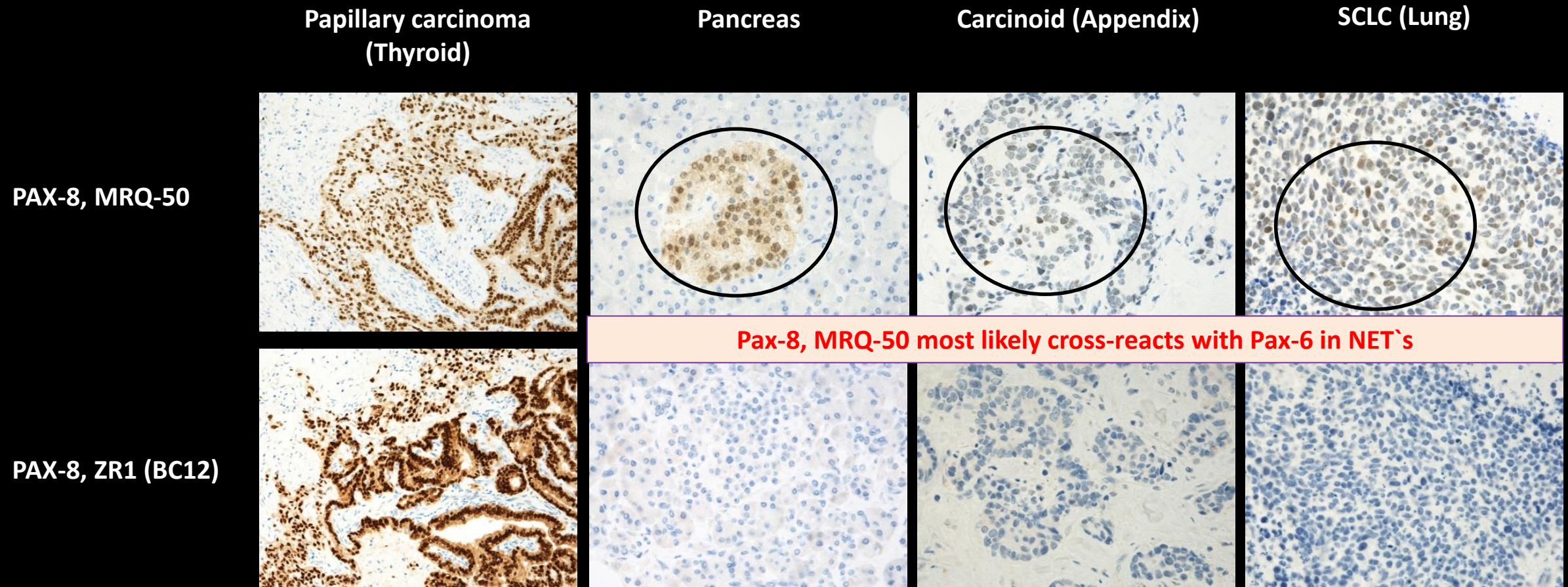
**Moretti L et al. : Mod Pathol. 2012; 25 : 231-236**

**Demonstrated that an N-terminal PAX-8 polyclonal antibody cross-react with N-terminal region of PAX-5 and is responsible for reports of PAX-8 positivity in malignant lymphomas.**

**Also, PAX8 mRNA levels were not detected in any of the B-cell lymphoma cell lines studied. These results indicate that benign and malignant B-cells do not express PAX8.**

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

**Problem: Primary antibody provides low specificity and/or poor signal-to-noise ratio**



Pax8, MRQ-50 most likely raised against the N-terminal part of the PAX8 protein (cross-reacts with other PAX proteins)

Pax8, ZR1 or BC12 raised against the C-terminal part of the PAX8 protein (no cross-reacting with other PAX proteins)



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

**Problem: Primary antibody provides low specificity and/or poor signal-to-noise ratio**

**Table 1. Antibodies and assessment marks for PAX8, run 62**

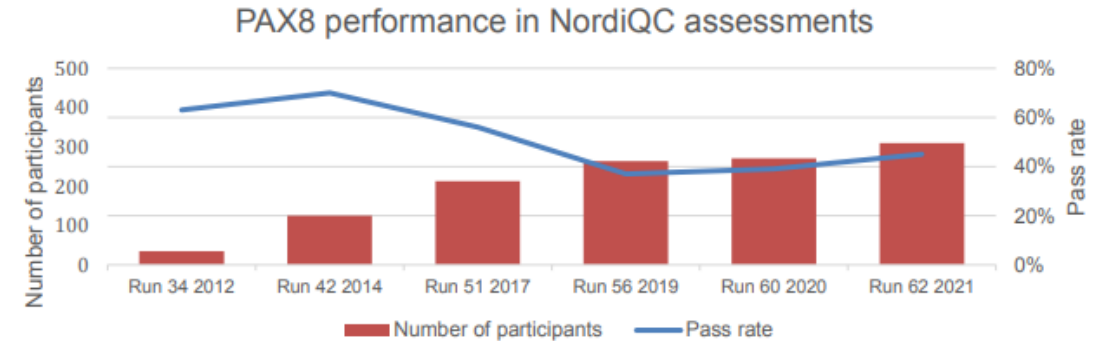
Concentrated antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff. <sup>1</sup>	OR <sup>2</sup>
mAb clone <b>BC12*</b>	4 1 1	Biocare Zytomed Systems Diagnostic Biosystems	-	3	1	2	50%	-
mAb clone <b>MRQ-50</b>	34	Cell Marque	-	18	8	8	55%	-
mAb clone <b>PAX8R1</b>	2	Abcam	-	2	-	-	-	-
rmAb clone <b>EP298*</b>	7 4 3 1	Cell Marque Epitomics <sup>5</sup> BIO SB Nordic Biosite	4	4	6	1	53%	27%
rmAb clone <b>EP331*</b>	8 4 1	Cell Marque Epitomics Abcam	-	5	7	1	39%	-
rmAb clone <b>SP348*</b>	55 5	Abcam Gennova	47	10	1	2	95%	78%
rmAb clone <b>ZR-1*</b>	5 1 1	Zeta Corporation Abcam Bio SB	3	1	3	-	57%	43%
pAb, <b>10336-1-AP</b>	21	Proteintech	-	8	9	4	38%	-
pAb, <b>363A-15</b>	1	Cell Marque	-	1	-	-	-	-
pAb, <b>CP379</b>	6	Biocare	-	1	4	1	-	-
Ab <b>QR016*</b>	2	Quartett	-	1	-	1	-	-
Unknown	2		-	-	2	-	-	-

mAb clone MRQ-50/pAbs cross react with other Pax proteins in the family (e.g. PAX5):

Should we use primary antibodies that cross react with other proteins in the same family ?

Would we accept cross-reactivity in the family of CD's and CK's - e.g., CD20 to CD3 or CK5 to CK8 ?

**Graph 1. Proportion of sufficient results for PAX8 in the six NordiQC runs performed**



## Conclusion

Optimal staining results could be obtained with the rmAb clones **EP298**, **SP348**, **ZR-1**, **BC12**, **IHC048** and **RM436**. Irrespective of the clone applied, efficient HIER, use of a sensitive 3-step polymer/multimer based detection system and careful calibration of the primary antibody were the most important prerequisites for an optimal staining result.

The rmAbs clones **EP298**, **SP348** and **ZR-1** gave encouraging results and a high proportion of sufficient results on the main fully automated platforms and no cross-reaction with e.g. PAX5 was observed. In contrast, the mAb clone **MRQ-50** provided a poor performance especially on the Ventana BenchMark and Dako Omnis platforms and at the same time also labelled PAX5 in B-cells. The **EP331** also provided a low pass rate due to aberrant nuclear staining reaction in non-PAX8 expressing cells and poor signal-to-noise ratio.

**BC12 (platform dependent)**

**“ZR1” (lot variations/antibody diluent dependent)**

**EP298**

**SP348**

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Problem: Primary antibody provides low specificity and/or poor signal-to-noise ratio

Table 1. Antibodies and assessment marks for SATB2, run 58

Concentrated antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff. <sup>1</sup>	OR <sup>2</sup>
mAb clone <b>CL0276</b>	5 2 1	Atlas Antibodies Sigma Aldrich Novus Biologicals	0	0	0	8	0%	0%
mAb clone <b>CL0320</b>	1	Atlas Antibodies	0	0	1	0	-	-
mAb clone <b>SATBA4B10</b>	3 2 2	Abcam Santa Cruz Zytomed Systems	0	0	2	5	0%	0%
mAb clone <b>OTI5H7</b>	1	ZSBio	1	0	0	0	-	-
rmAb clone <b>EP281</b>	30 12 1 1 1	Epitomics Cell Marque Immunologic BioSB Biocare Medical Unknown	22	14	4	6	78%	82%
rmAb clone <b>SP281</b>	4 1	Abcam Spring Bioscience	2	1	1	1	60%	40%
rmAb clone <b>ZR167</b>	1	Nordic Biosite	1	0	0	0	-	-
rmAb clone <b>EPNCIR130A</b>	5	Abcam	0	0	0	5	0%	0%
pAb <b>HPA001042</b>	5	Sigma Aldrich	0	0	2	3	0%	0%
pAb <b>Ab69995</b>	1	Abcam	0	0	0	1	-	-
Ready-To-Use antibodies							Suff. <sup>1</sup>	OR <sup>2</sup>
rmAb clone <b>EP281 384R-17/18</b>	19	Cell Marque	7	10	1	1	89%	37%
rmAb clone <b>EP281 PR/HAR239</b>	2	PathnSitu	2	0	0	0	-	-
rmAb clone <b>EP281 API3225</b>	1	Biocare Medical	0	1	0	0	-	-
rmAb clone <b>EP281 MAD-000747QD</b>	1	Máster Diagnostica	0	0	1	0	-	-
rmAb clone <b>EP281 BSB3199</b>	2	BioSB	0	0	0	2	-	-
Total	105		35	26	12	32	-	
Proportion			33%	25%	11%	31%	58%	

FP/FN

FP/FN

FN

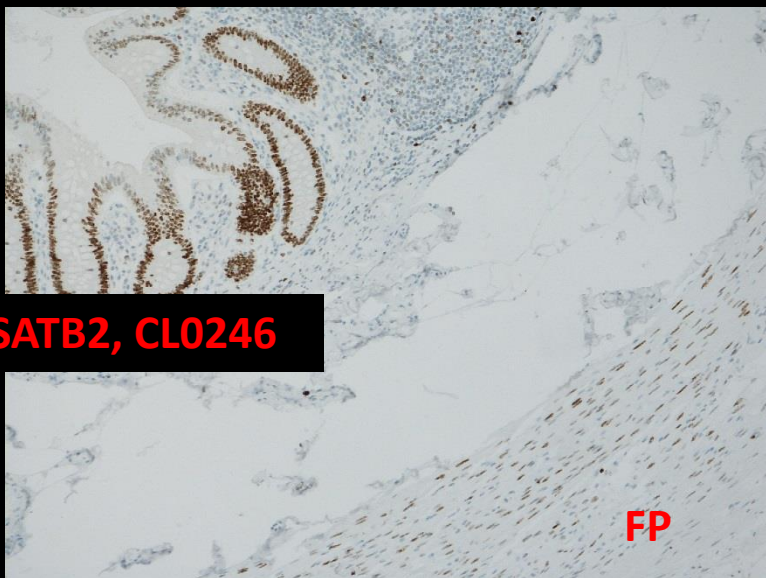
FP/FN + aberrant cytoplasmic staining

FN + aberrant cytoplasmic staining

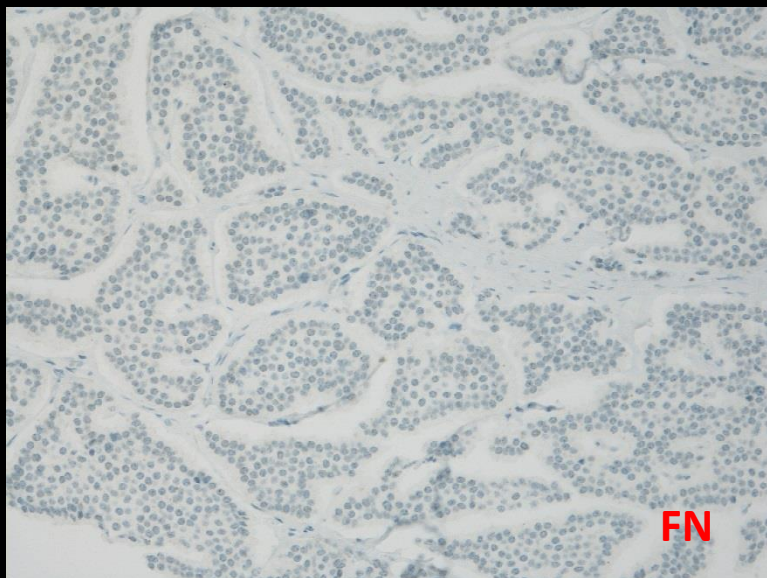
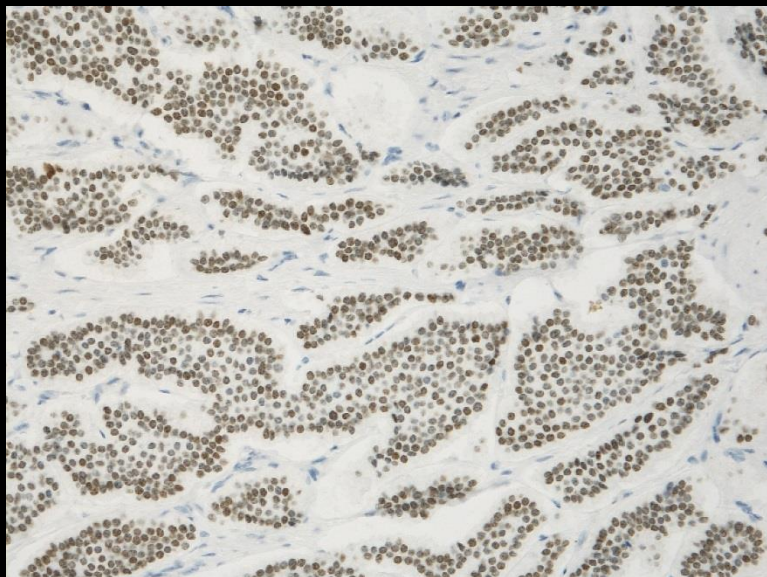
Impossible to calibrate correctly



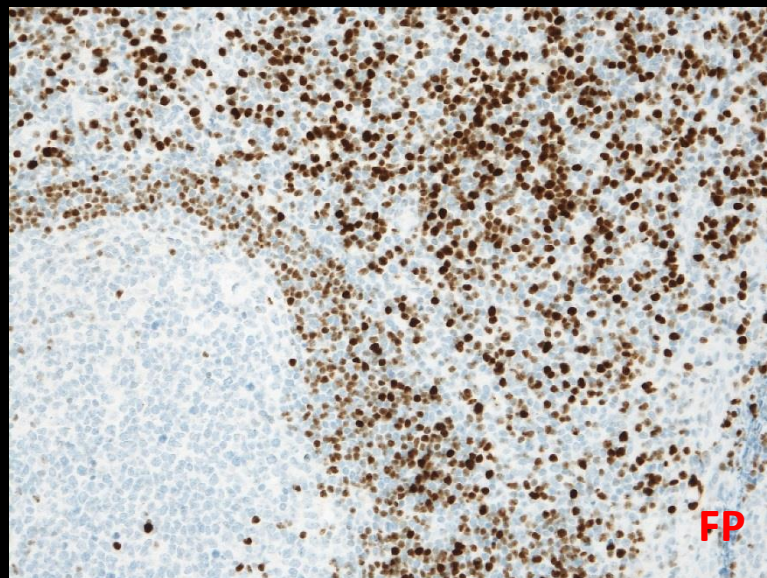
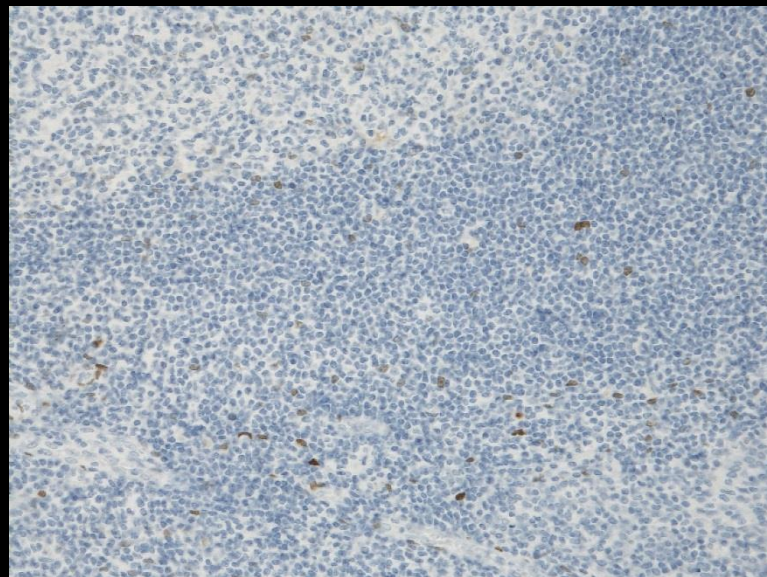
Appendix



Neuroendocrine tumour (colon)



Tonsil

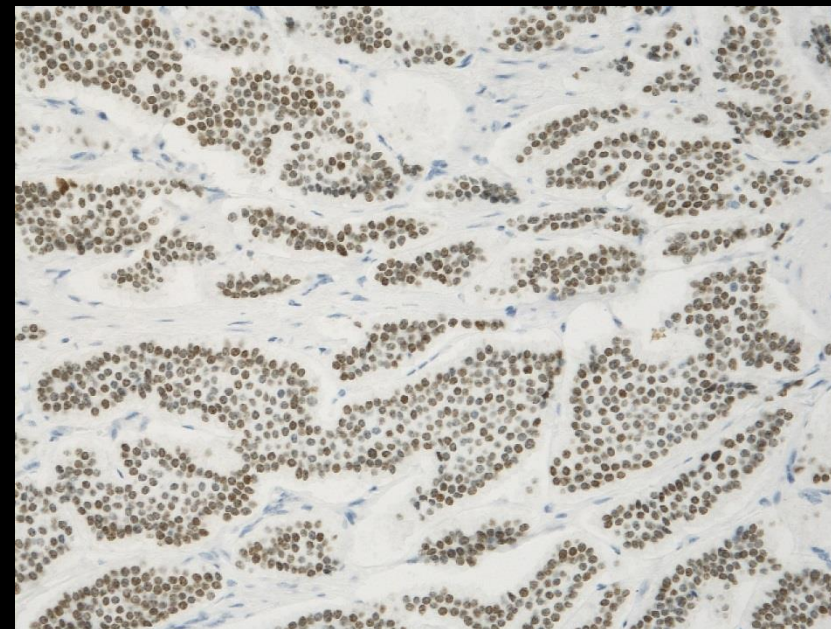




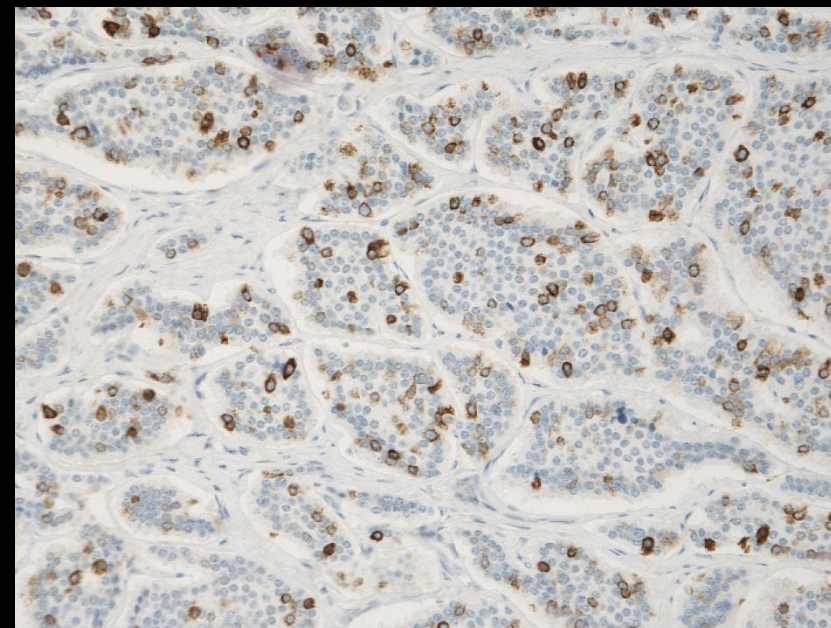
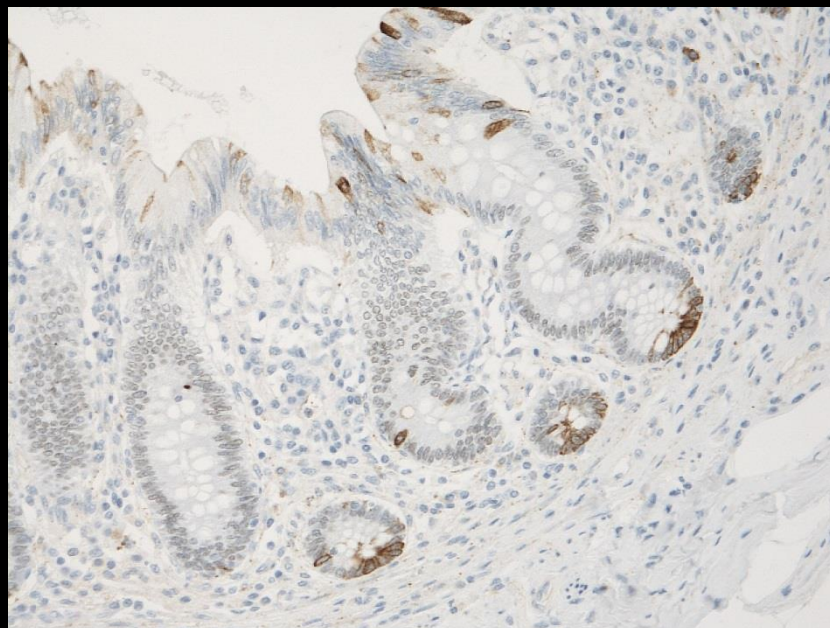
Appendix

Neuroendocrine tumour (colon)

SATB2, EP281



SATB2, Pab 69995





# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Antibody choice: Sensitivity & Specificity

Abs providing low sensitivity	Abs providing low specificity and/or aberrant staining
SATB2 clone CL0276 & SATBA4310	SATB2 clone CL0276 & SATBA4310
SATB2 clone EPNCIR130A	PAX8 clone EP331 & MRQ-50
Uroplakin II+III clone AU-1 & SP73 & EP321 (all = Uro III)	MUM1 clone MRQ-43 & BC5
P16 clone G175-405	CK-HMW clone 34βE12
TTF1 clone 8G7G3/1	PR clone 1E12
ERG (Ets-Related-Gene) clone 9FY	ECAD clone EP700Y
ALK clone ALK1	PAX5 clone SP34
CK8/18 clone DC10 & C51 & CY90 (all = CK18)	SMAD4 clone RBT-SMAD4 & SP306
CEA clone II-7	MLH1 clone M1 & G168-728
CGA clone DAK-A3	CD79a clone HM57
P63 clone 7JUL	MSH6 clone 44
CD79a clone HM57 & 11E1	Many pAbs (e.g., P40 and SOX10)
ALK clone ALK1	.....
.....	

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

**Problem: Poorly calibrated primary antibody (false negative or false positive)**

**The right primary antibody**

**The right protocol (AR procedure and detection system)**

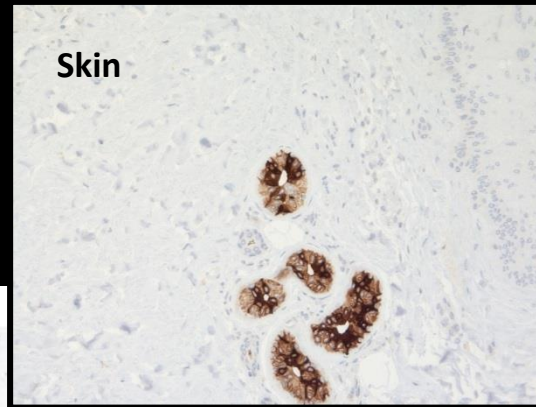
**Poorly calibrated primary Ab ?**

**Tissue controls are the key element**

**Normal skin is the preferred positive control for GCDFP-15. The epithelial cells of the eccrine sweat glands must show an as strong as possible positive cytoplasmic staining reaction, while all other cells should be negative.**

**Normal breast tissue can also be used as control in which epithelial cells of the ductal glands must show an as strong as possible staining reaction.**

**Background staining may be seen in vicinity of highly positive tissue structures (e.g. eccrine sweat glands)**



## Gross cystic disease fluid protein-15 (GCDFP-15)

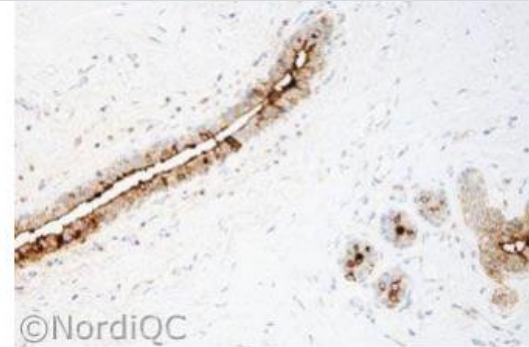


Fig. 1a  
Optimal staining for GCDFP-15 of the breast hyperplasia using the mAb clone 23A3 optimally calibrated as a concentrate, HIER in an alkaline buffer and a polymer based detection system. The majority of the ductal epithelia cells show a distinct moderate to strong cytoplasmic staining reaction. Also compare with Fig. 2a – same protocol.

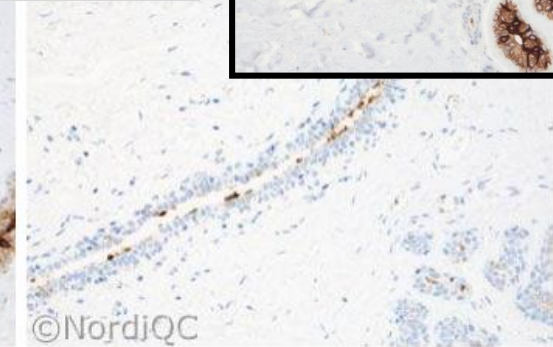


Fig. 1b  
Insufficient staining for GCDFP-15 of the breast hyperplasia applying the mAb clone 23A3 as a concentrate using exactly the same protocol settings as used in Fig 1a, except for a 20 fold dilution of the primary antibody. The proportion and the intensity of the cells demonstrated are significantly reduced compared to the result in Fig. 1a. Also compare with Fig. 2b – same protocol.

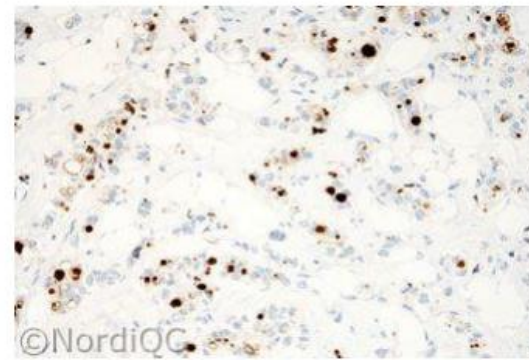


Fig. 2a  
Optimal staining for GCDFP-15 of the breast carcinoma no. 5 using same protocol as in Fig. 1a.  
The majority of the neoplastic cells show a moderate to strong dot-like cytoplasmic staining reaction.

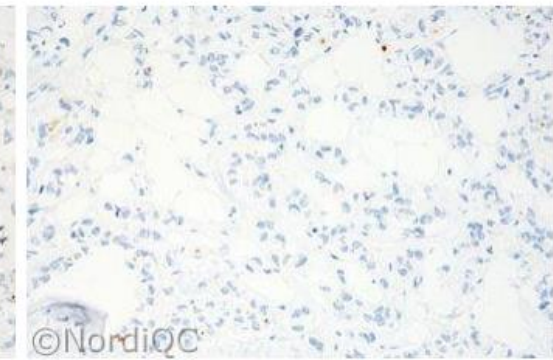
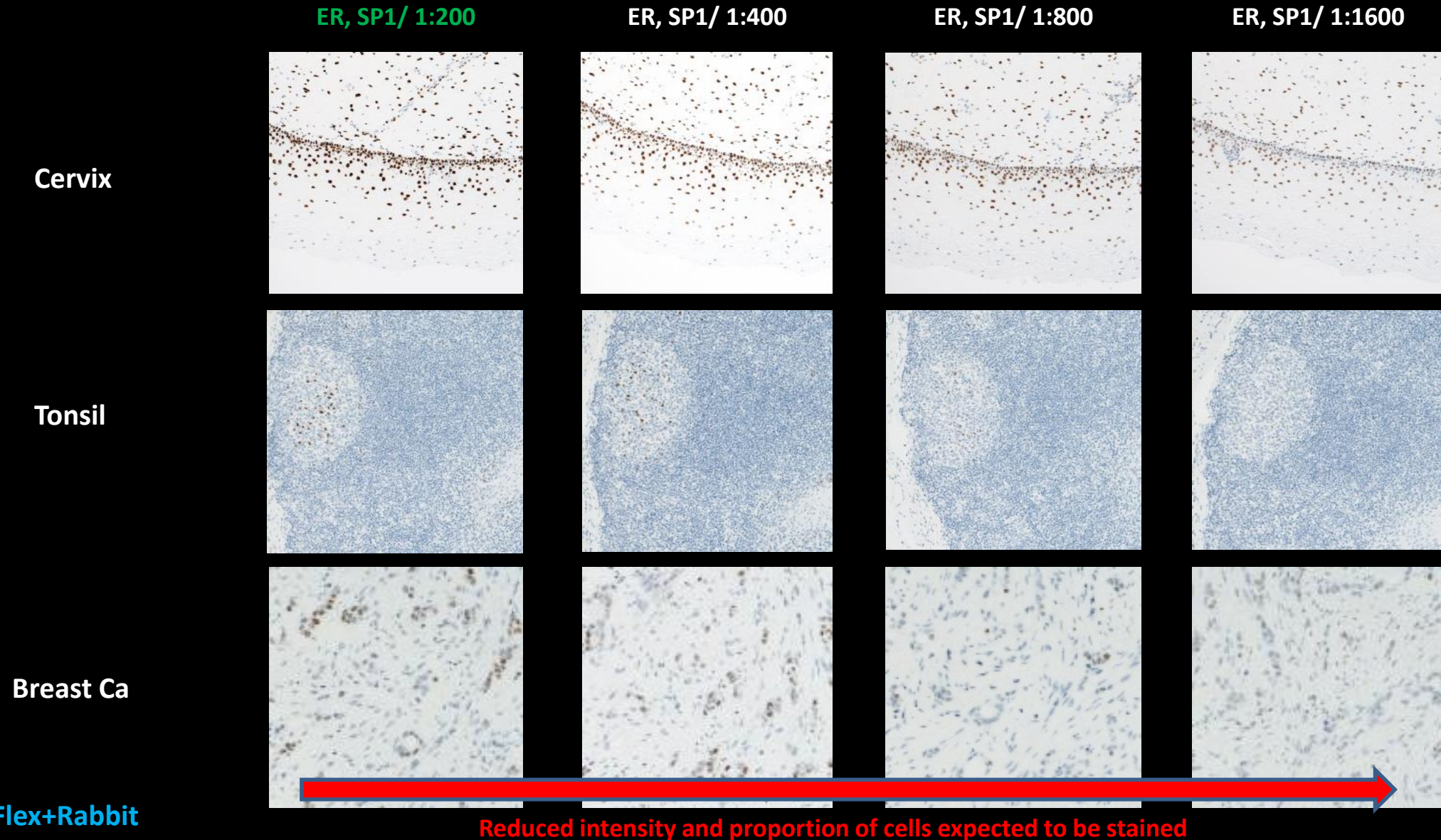


Fig. 2b  
Insufficient staining GCDFP-15 of the breast carcinoma no. 5 using same protocol as in Fig. 1b. - same field as in Fig. 2a.  
Only scattered neoplastic cells show a faint dot-like reaction.



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

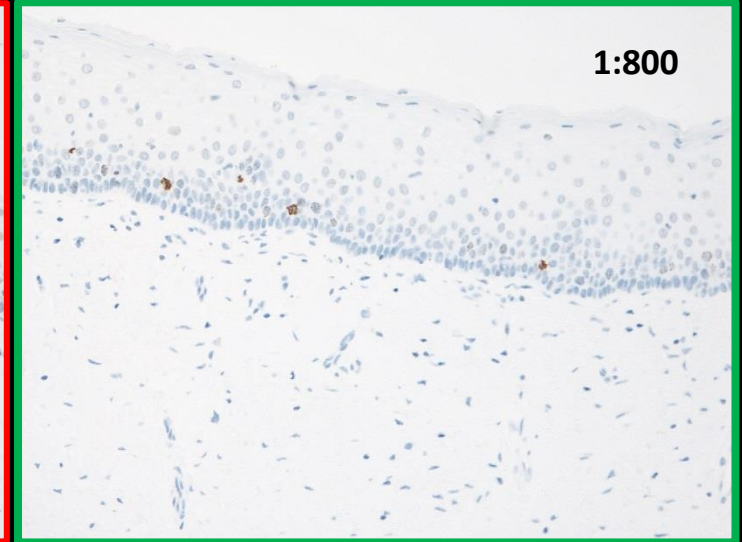
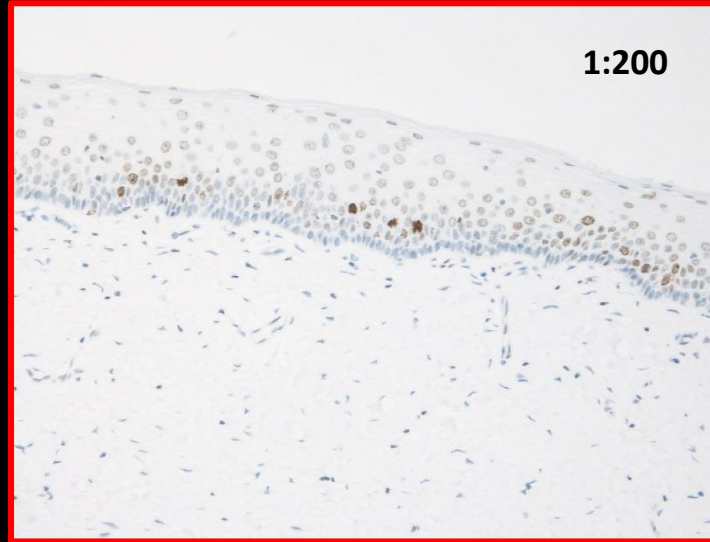
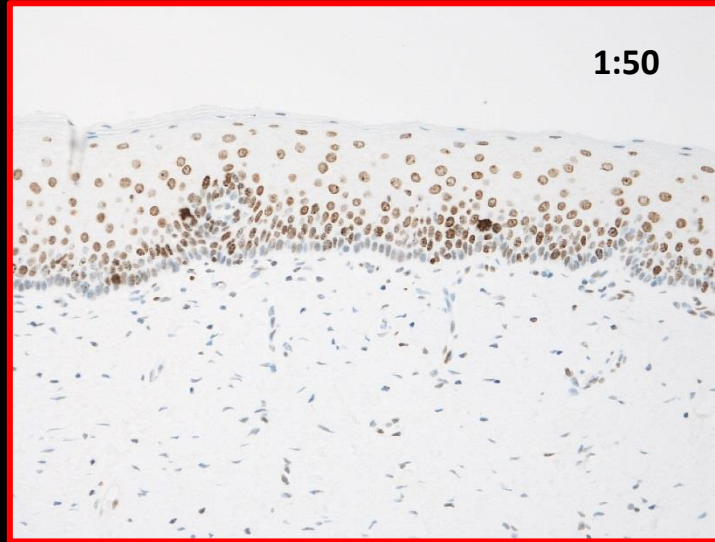
Problem: Primary antibody poorly calibrated (false negative)



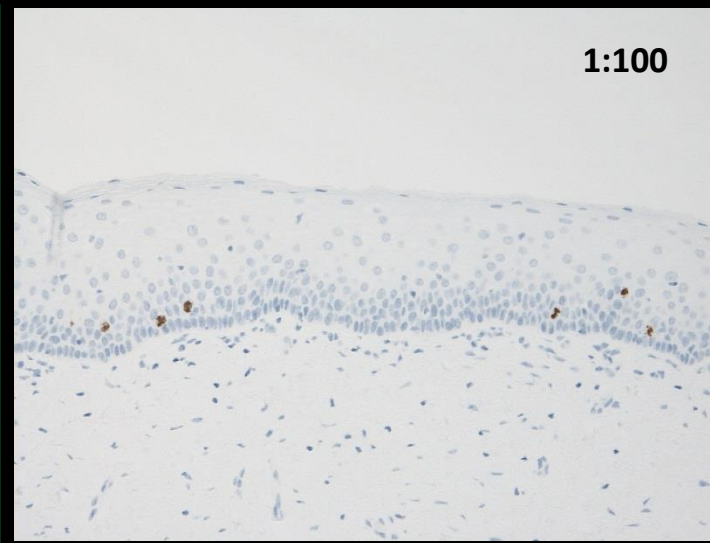
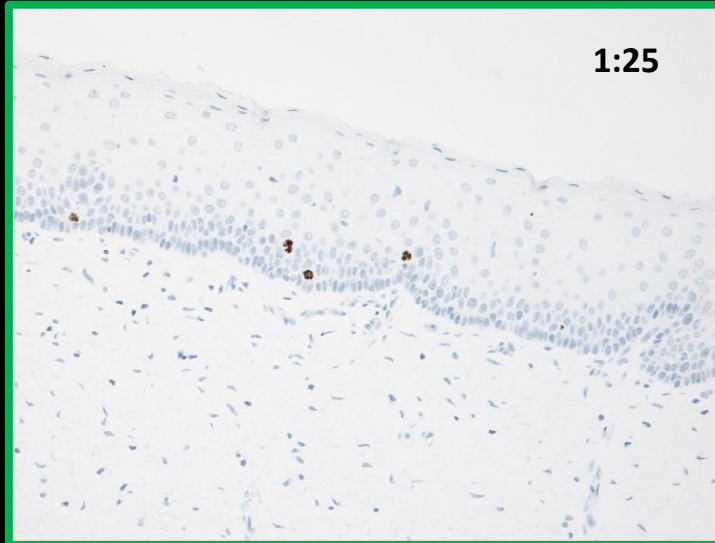
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Problem: Primary antibody poorly calibrated (false positive)

PHH3, poly  
(Cell Marque)



PHH3, BC37  
(Biocare)



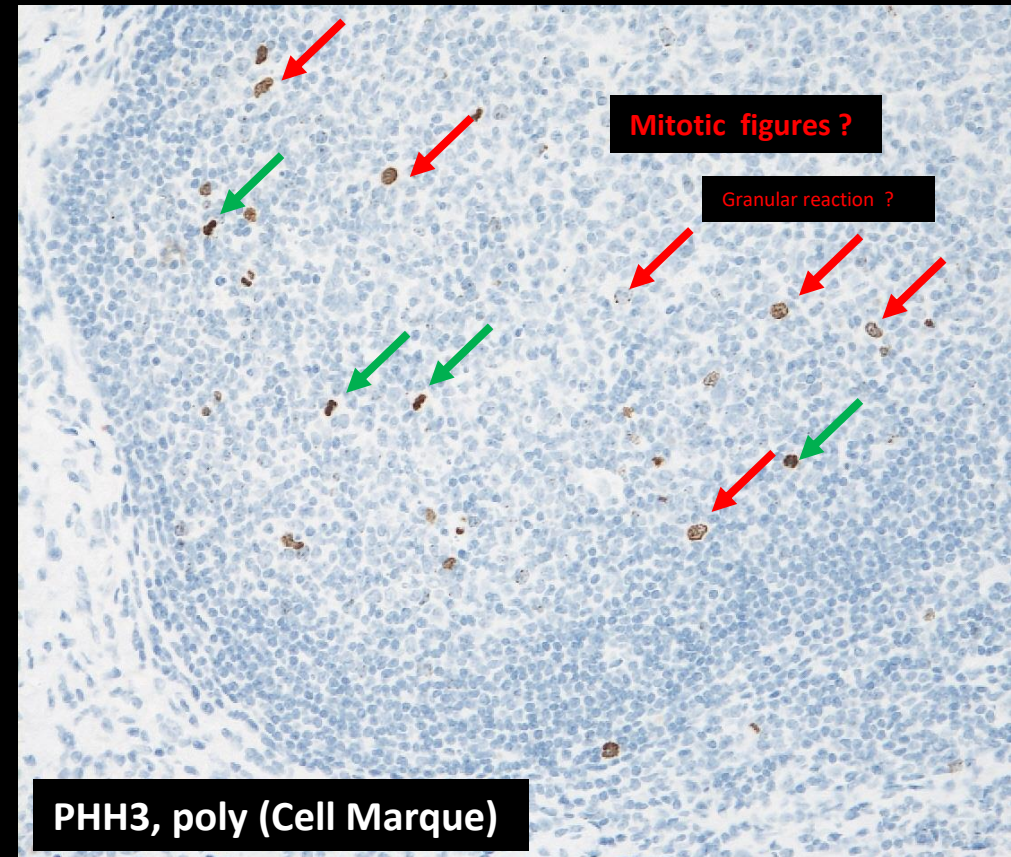
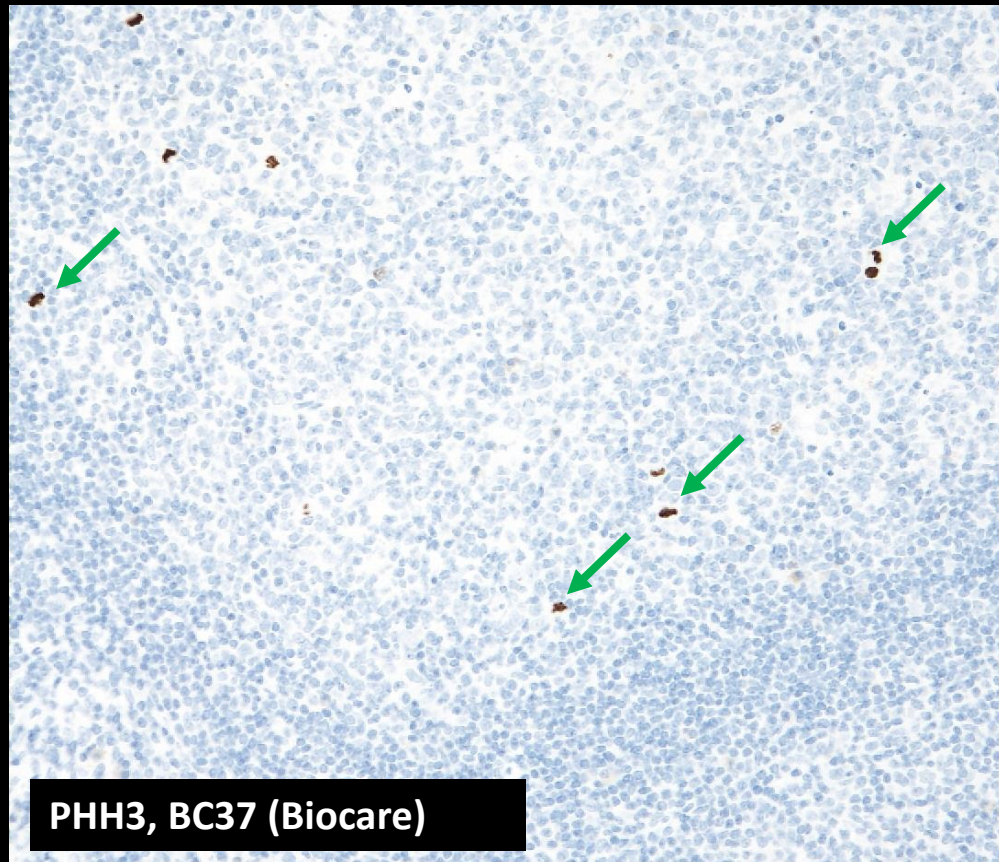
HIER Low pH/Flex+

PHH3 should only be positive in cells in the late G2 and M phase (mitotic cells )



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Problem: Primary antibody poorly calibrated (false positive)



PHH3, poly might not be completely phospho-specific and might cross-reacts with cells in e.g. interphase

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## RESEARCH ARTICLE

---

### Performance of 4 Immunohistochemical Phosphohistone H3 Antibodies for Marking Mitotic Figures in Breast Cancer

*Cornelia M. Focke, MD,\*† Kai Finsterbusch, MSc,\* Thomas Decker, MD,\*  
and Paul J. van Diest, MD, PhD†*

**Conclusions:** Performance and reliability varied significantly between the 4 tested antibodies. For faster identification of mitotic hot spots and as potential marker in digital image analysis, the Merck antibodies seem to be most suitable.

**Key Words:** PHH3, breast cancer, proliferation, mitotic activity, phosphohistone H3

*(Appl Immunohistochem Mol Morphol 2016;00:000–000)*

OBS – Præsenteres ikke ved workshoppen ?



## Proficiency testing in immunohistochemistry—experiences from Nordic Immunohistochemical Quality Control (NordiQC)

Mogens Vyberg<sup>1,2</sup> · Søren Nielsen<sup>1</sup>

Virchows Arch (2016) 468:19–29

Table 3 Major causes of insufficient staining reactions

1. Less successful antibodies (17 %)
  - a. Poor antibodies<sup>a</sup>
  - b. Less robust antibodies<sup>b</sup>
  - c. Poorly calibrated RTUs
  - d. Stainer platform dependent antibodies
2. Insufficiently calibrated antibody dilutions (20 %)
3. Insufficient or erroneous epitope retrieval (27 %)
4. Error-prone or less sensitive visualization systems<sup>c</sup> (19 %)
- 5 Other (17 %)
  - a. Heat-induced impaired morphology
  - b. Proteolysis induced impaired morphology
  - c. Drying out phenomena
  - d. Stainer platform-dependant protocol issues
  - e. Excessive counterstaining impairing interpretation

**37% insuff.**

<sup>a</sup> Consistently gives false negative or false positive staining or a poor signal-to-noise ratio in one or more assessment runs

<sup>b</sup> Frequently giving inferior staining results, e.g., due to mouse-anti-Golgi reactions or sensitive to standard operations as blocking of endogenous peroxidase

<sup>c</sup> Biotin-based detection kit for cytoplasmic epitopes, use of detection kits providing a too low sensitivity, or use of detection kits and chromogens giving imprecise localization of the staining signals complicating the interpretation

### Problems related to:

#### ❑ The choice and use of the primary antibody (Concentrate or RTU)

- Inappropriate primary antibody
  - Provide low sensitivity/specificity
- Appropriate primary antibody
  - Inapp. titre (too low or too high concentration)
- Platform (stainer) dependent antibodies
  - Provide low sensitivity / specificity

**False positive or false negative results**

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Primary antibodies sensitive to the chosen platform



Implementing a new platform has been a challenge

ALK clone D5F3 or 5A4  
HCL, clone DBA44  
GATA3, clone L50-823  
MART-1/Melan A, clone 103  
PAX 8, clone BC12  
SMAD4, clone B8  
WT1, clone WT49  
MMR  
ASMA, 1A4  
.....

Changing the primary Ab

Changing Ab-Ag reaction microenvironment (Diluent)

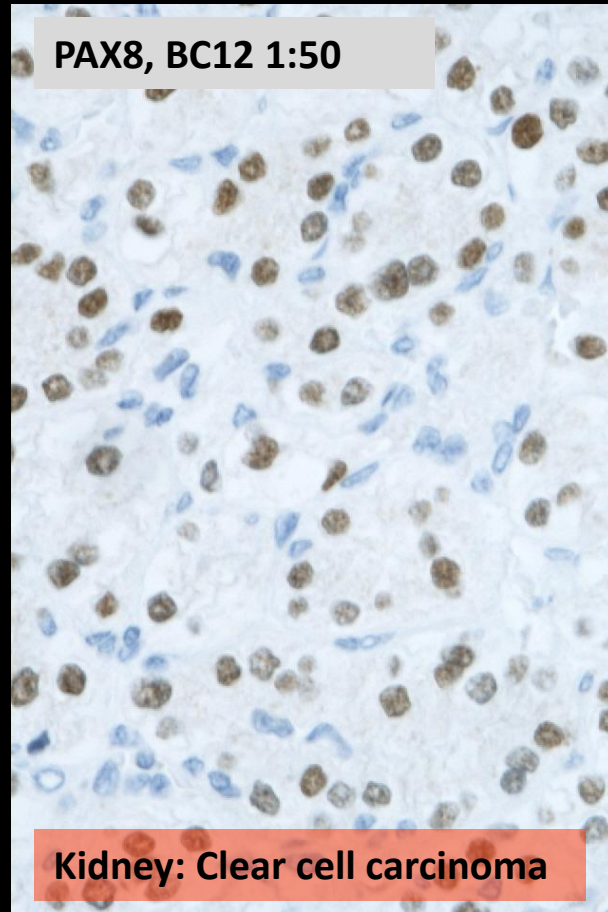
Low affinity primary antibodies



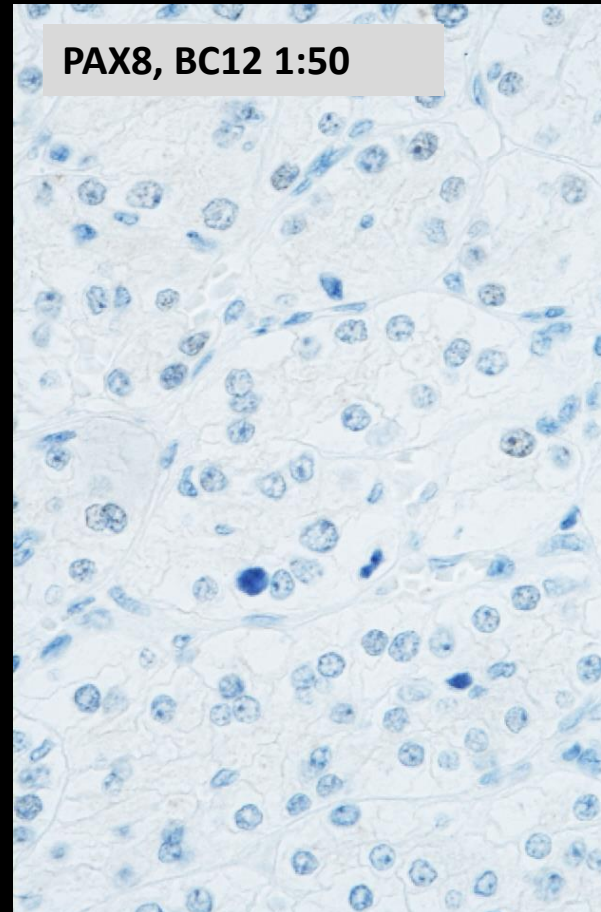
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Primary antibodies  
sensitive to the  
chosen platform

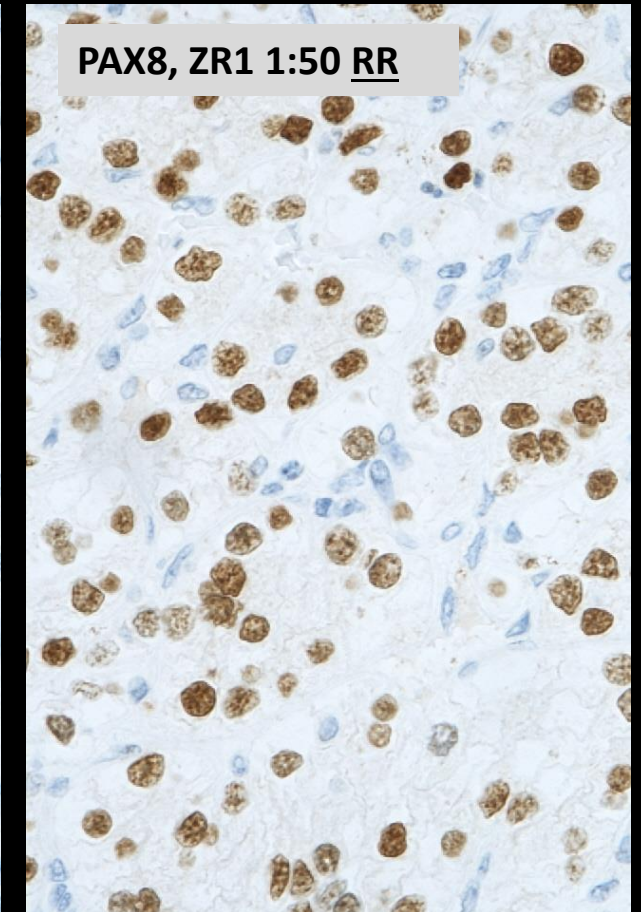
Autostainer



Omnis



Omnis



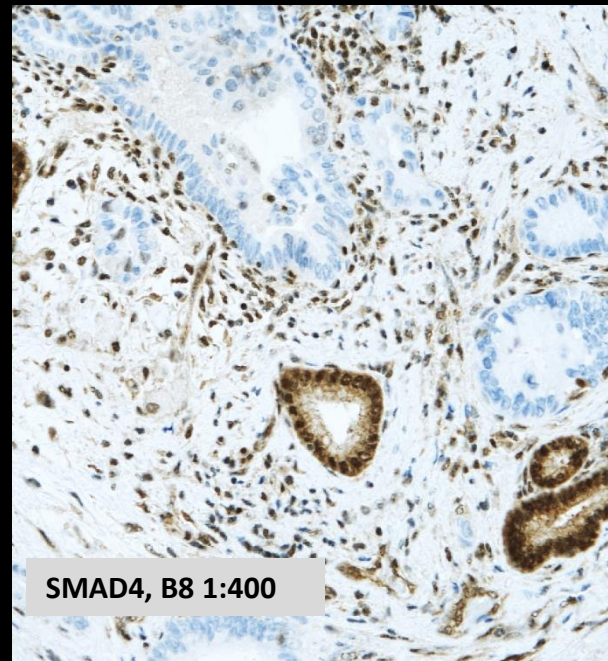
HIER High pH 20', Flex+ (10+20)

HIER High pH 48', Flex+ (10+20)

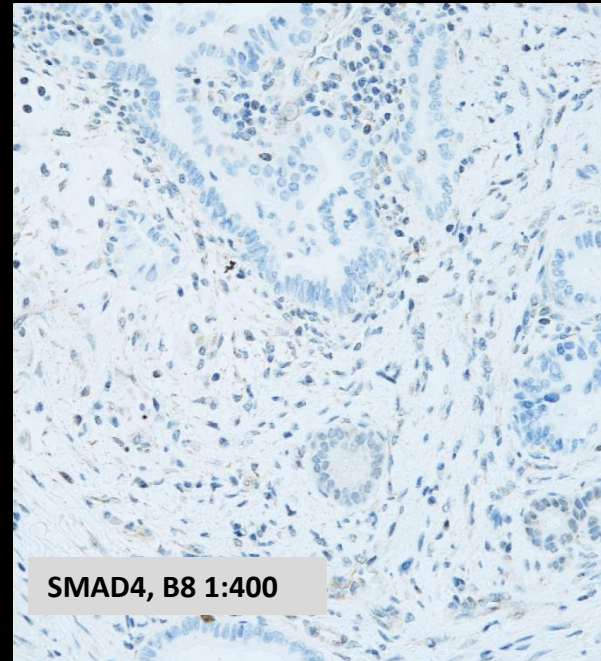
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Primary antibodies  
sensitive to the  
chosen platform

**Autostainer**



**Omnis**



**Omnis**



HIER High pH 20', Flex+ (10+20)

HIER High pH 24', Flex+ (10+20)

**Pancreatic Adenocarcinoma**



## Challenging and platform dependent antibodies

Marker	Problematic clone	Alternative
ASMA	1A4	BS66
BCL2	124	"E17" or SP66
CD3	F7.2.38	LN10
CD4	4B12	EP204
CD23	1B12	DAK-CD23
CD56	123C3 & 123C3.D5	MRQ-42
CDX2	DAK-CDX2	EPR2764Y or EP25
CEA	II-7	CEA31
CK (LMW)	5D3	EP17/EP30
PMS2	A16-4	EP52 or EPR3947

Marker	Problematic clone	Alternative
CR	DAK-Calret1	CAL6
Desmin	D33	BS21
EPCAM	BER-EP4	BS14
Melan A	A103	EP43 (Melanomas)
OCT 3/4	C-10	MRQ-10 or N1NK
PAX8	MRQ-50	SP348
P16	E6H4	JC2 or MX007
WT1	6F-H2	D817F or EP122
SMAD4	B-8	EP618Y
.....		

Experience from Dept. of surgical Pathology, Region Zealand, Denmark on the Omnis - These alternative clones might also work on other platforms (e.g., Benchmark Ultra or Bond MAX/III).

Go to the NordiQC website for information of the individual markers in relation to the chosen platform

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Problem: Platform dependent antibodies

NordiQC results (2018-2022)

Antibody	Clone	Platform(s)
CD56	123C3	BenchMark (Ventana) and Omnis (Dako)
PMS2	A16-4	BenchMark
Melan A	A103	Benchmark and "Omnis"
P16 (RTU format's Ventana)	E6H4	Omnis
Alpha Smooth Muscle Actin	1A4	Benchmark
SMAD4	B8	Benchmark and Omnis
CK8/18	5D3	Benchmark
EPCAM	BER-EP4	Benchmark and BOND (Leica)
BRAFmut	VE1	In general challenging on most platforms except for the Benchmark Ultra (Ventana)
CR	DAK-Calret1	Benchmark and Omnis
Desmin	D33	Omnis
.....		

Problem: Does the automatic platform come with appropriate reagents fulfilling purpose and intended use of the assay e.g., Antigen Retrieval Solutions, Antibody diluents, Detection systems.....



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## HIER buffers used by NordiQC participants

App. 95 % of all pretreatment protocols

In house	Agilent Dako AS/Omnis	Roche Ventana Benchmark	Leica Biosystems Bond	Biocare	Thermo S LAB Vision
<u>Low pH buffers</u>					
Citrate buffer pH 6 / pH6.7	TRS Low pH 6.1	CC2 pH 6	BERS-1 pH 6	Diva Decloaker pH 6.2	
<u>High pH buffer</u>					
EDTA/EGTA pH 8	TRS High pH 9	CC1 pH 8.5	BERS-2 pH 9	Borg Decloaker pH 9.5	HIER buffer H pH 9
Tris-EDTA/EGTA pH 9	TRS High (3-in-1) pH 9				
Tris-HCL pH 9					

### Challenges:

The platform often dictates the choice of HIER buffers



Assessment Run 56 2019

## Epithelial cell-cell adhesion molecule (Ep-CAM)

Table 1. Antibodies and assessment marks for Ep-CAM, run 56

Concentrated antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff. <sup>1</sup>	Suff. OPS <sup>2</sup>
mAb clone <b>BS14</b>	10	Nordic Biosite	9	1	0	0	100%	100%
mAb clone <b>Ber-Ep4</b>	69	Dako	14	13	21	28	36%	93%
	6	Cell Marque						
	1	Diagnostic Biosystems						
mAb clone <b>MOC-31</b>	23	Dako	10	10	7	2	69%	71%
	5	Cell Marque						
	1	Diagnostic Biosystems						
mAb clone <b>VU-1D9</b>	5	Thermo Scientific	9	0	1	0	90%	100%
	3	Merck Millipore						
	1	Immunologic						
	1	Novus Biologicals						
rmAb clone <b>EPR20532-225</b>	1	Abcam	0	0	0	1	-	-
Ready-To-Use antibodies								
mAb clone <b>Ber-Ep4 760-4383</b>	16	Ventana/Cell Marque	1	6	6	3	44%	100%
mAb clone <b>Ber-Ep4 248M-98</b>	49	Cell Marque	5	13	16	15	37%	-
mAb clone <b>Ber-Ep4 IR/IS637</b>	18	Dako	5	9	3	1	78%	87%
mAb clone <b>Ber-Ep4 IR/IS637<sup>3</sup></b>	6	Dako	1	2	2	1	-	-
mAb clone <b>Ber-Ep4 GA637</b>	27	Dako	26	1	0	0	100%	100%

BS14 could be an alternative to Ber-EP4 on platforms excluded from the use of modified low pH buffers e.g., Diva pH 6.2 (Biocare) or TRS pH6.1 (Dako)

HIER in alkaline pH buffers (+/- gentle enzymatic digestion performed after Hier)

OPS based on Hier in mod. Low pH buffers (Dako or Biocare)

OPS based on Hier in alkaline (CC1) or mod. Low pH buffers (Dako)

Omitted from Hier in mod. Low pH buffers (TRS low pH/Dako or Diva Decloaker/Biocare) - these buffers are not part of the reagent portfolio

HIER in mod. Low pH buffers (Dako)



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## EPCAM clone Ber-EP4 or BS14

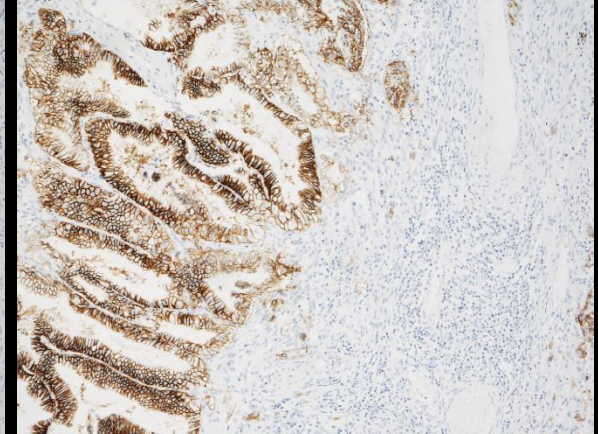
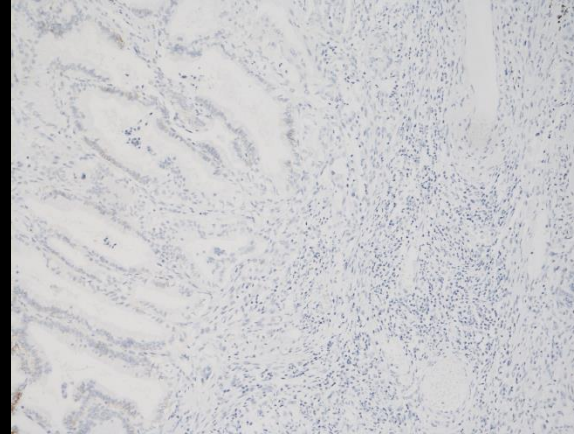
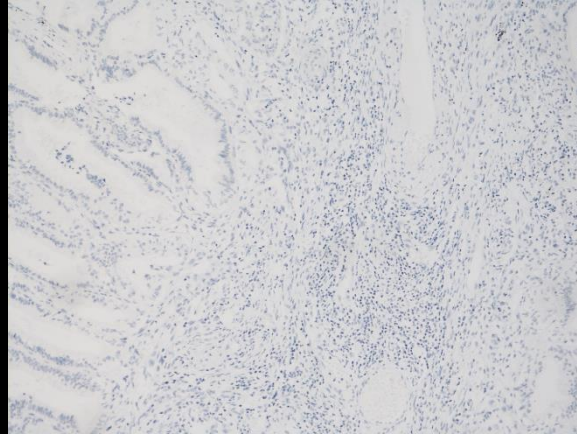
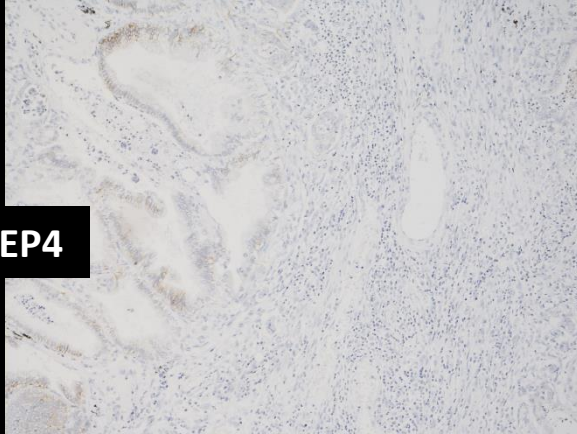
Proteinase K 1:4 (Dako) / 5'

Citrate buffer pH 6 (Dako) / 20'

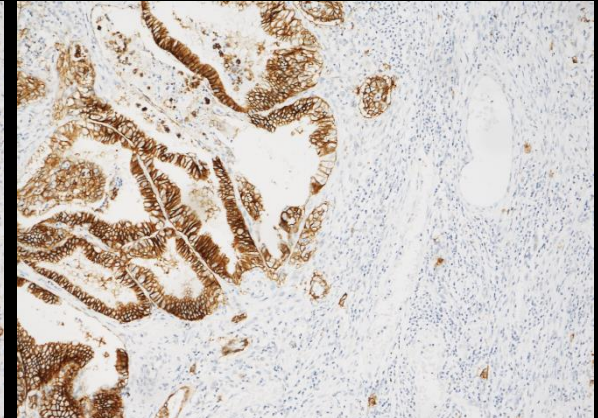
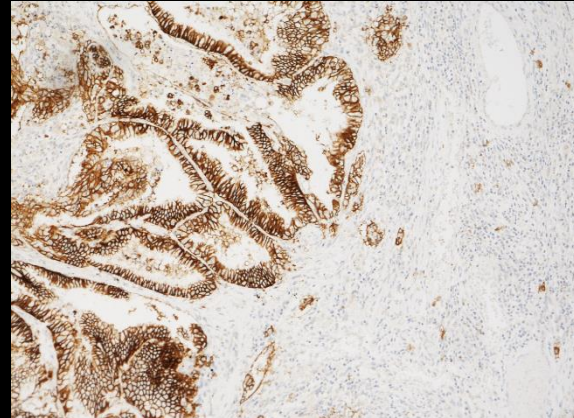
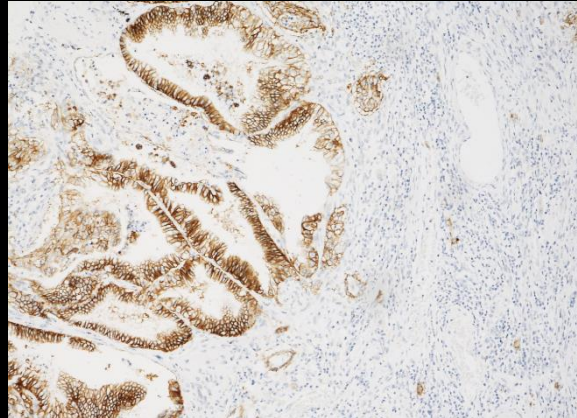
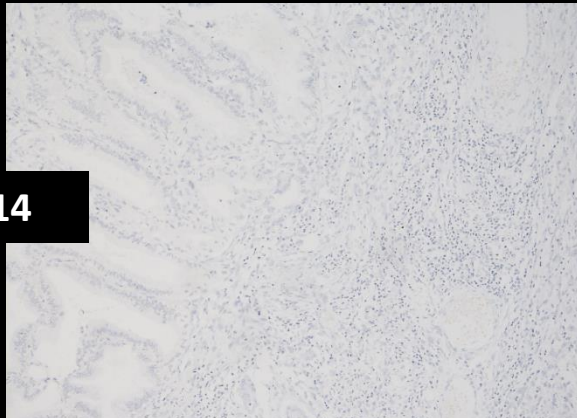
High pH buffer (Dako) / 20'

TRS Low pH (Dako, S1699/S1700) / 20'

Ber- EP4



BS14



EPCAM, BS14 (Nordic Biosite) is a better alternative than EPCAM MOC31 or Ber-EP4 for automated platforms excluded from use of mod. low pH buffers.

Colon tumor

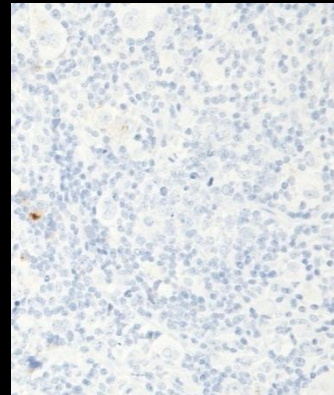


# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

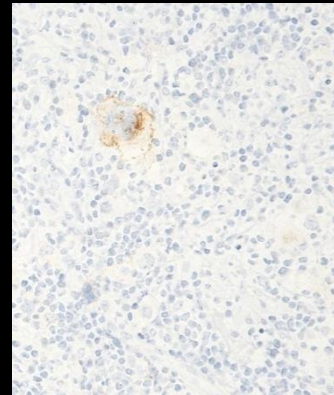
Important questions: Antibody - Antigen retrieval procedure - Automated platform

CD30  
Clone  
ConD6/B5

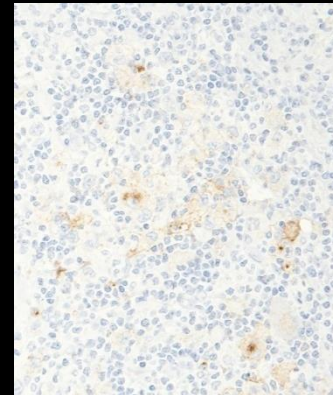
Ci pH 6  
MWO / 20 min



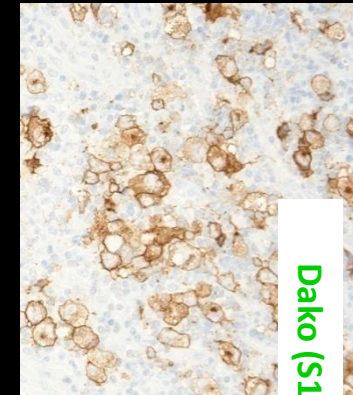
EDTA pH 8  
MWO / 20 min



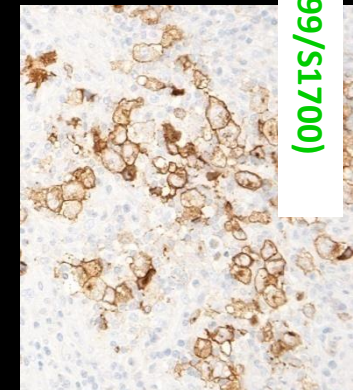
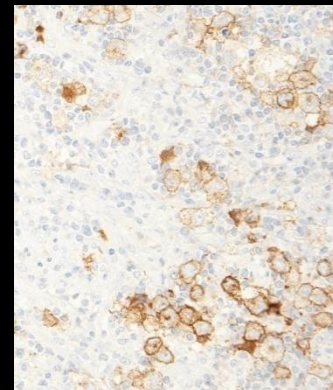
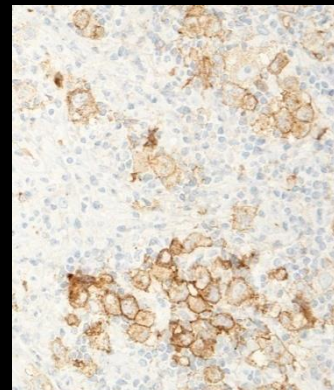
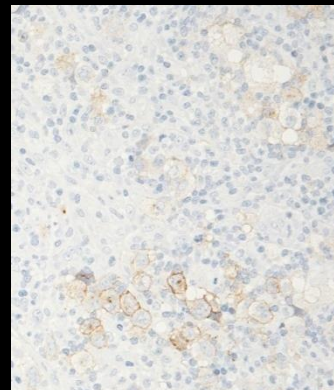
TE pH 9  
MWO / 20 min



TRS pH 6.1  
MWO / 20 min



CD30  
Clone  
Ber-H2



Dako (S1699/S1700)

Hodgkin Lymphoma



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Primary antibodies sensitive to the chosen platform



Implementing a new platform has been a challenge

ALK clone D5F3 or 5A4  
HCL, clone DBA44  
GATA3, clone L50-823  
MART-1/Melan A, clone 103  
PAX 8, clone BC12  
SMAD4, clone B8  
WT1, clone WT49  
MMR  
ASMA, 1A4  
.....

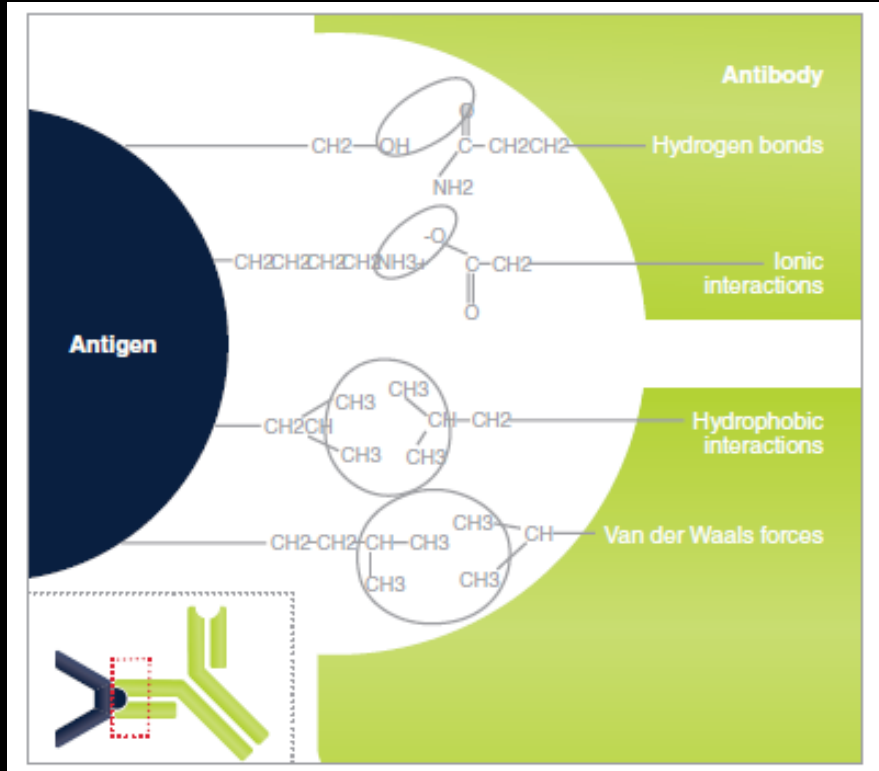
Changing the primary Ab

Changing Ab-Ag reaction microenvironment (Diluent)

Low affinity primary antibodies

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Antibody Diluents: Antigen-Antibody reactions



The strength by which the primary Ab binding site binds to an antigenic epitope is called affinity

Antibodies are attracted initially through electrostatic interactions, and subsequently through weak forces

- Hydrogen bonds
- Hydrophobic interactions
- Van der Waals forces

Antibody diluent formulations can significantly alter stability and binding properties of antibodies affecting both epitope specificity and non-specific interactions



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Applied Immunohistochemistry & Molecular Morphology 9(2): 176–179, 2001

© 2001 Lippincott Williams & Wilkins, Inc., Philadelphia

## Formalin-Fixed and Heat-Retrieved Tissue Antigens: A Comparison of Their Immunoreactivity in Experimental Antibody Diluents

Thomas Boenisch, M.S.

### Antibody diluents

#### Demonstrated that:

**pH of the antibody diluent had a high impact of the final IHC result**

**Addition of NaCl (increasing the ionic strength) to the diluent negates most of the sensitivity gained through Antigen Retrieval (Table 3).**

**TABLE 3.** Comparison of staining scores of 13 optimally diluted antibodies as a function of antigen retrieval at pH 9.9, use of 0.05 M Tris (TB), pH 6.0 and 8.6, or Tris-buffered 0.15 M NaCl (TBS) of pH 6.0 and 8.6, and 0.02 M phosphate-buffered 0.15 M NaCl of pH 7.5 (PBS)

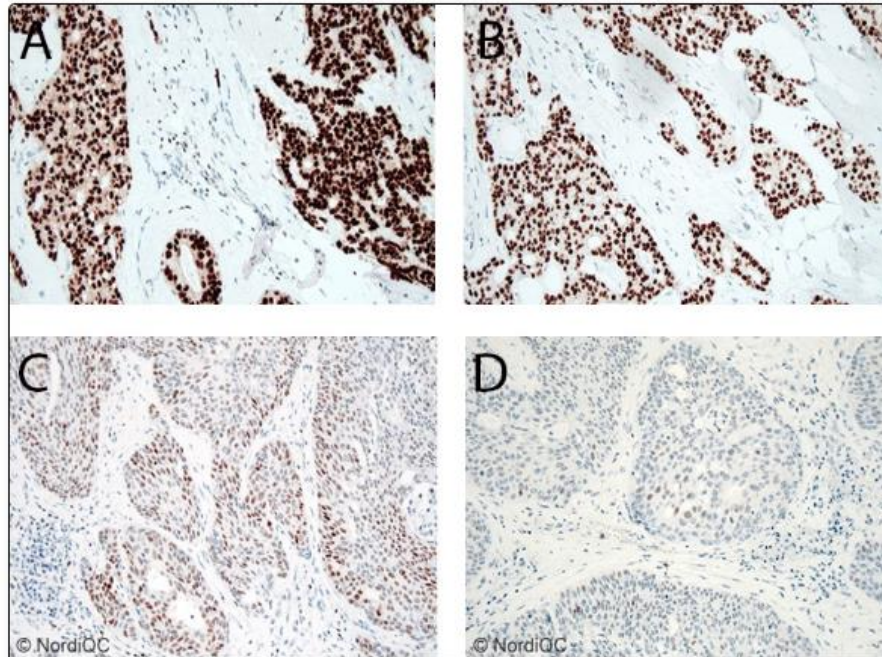
Clone	pH	TB		TBS		PBS
		6.0	8.6	6.0	8.6	7.3
BLA.36		2	4	1	2	1
UCHL1		4	3	2	1	1
L26		4	3	3	3	2
PC10		4	3	4	4	3
N10/2		3	2	1	2	1
V9		4	3	4	4	2
TAL1B5		4	2	3	2	2
ER-PR-8		4	3	2	1	2
Ber-H2		4	3	ND	ND	0
4KB5		4	2	4	2	4
DF-T1		4	2	2	0	1
PD7/26		4	3	ND	ND	3
C3D-1		4	2	ND	ND	1

ND, not done.

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase



Info ▾ Modules ▾ Assessments Protocols Controls Events ▾ [Login](#)



Ductal breast carcinoma (A and B) and urothelial carcinoma (C and D) stained for GATA3 (clone L50-823) in two laboratories. Optimal results in A and C, insufficient in D. The only difference in protocols is the antibody dilution buffer: In A and C the antibody was diluted in Biocare Renoir red (pH 6.2). In B and D the Dako antibody diluent pH 7.3 was used.

Results - Run 54, B26, H14, C4

14-Dec-2018

The results for the runs 54, B26, H14, C4 are now available on the website. Individual results can be seen after logging in.

[All news](#)

## Events

[NordiQC Workshop in Diagnostic Immunohistochemistry](#)  
2-4 Oct 2019: Aalborg, DK

6th Annual Course of Academy of Immunohistochemistry "Diagnostic Immunohistochemistry and Molecular"  
9-11 Oct 2019: Krakow, Poland

4th NordiQC Conference on Applied Immunohistochemistry  
2-5 Jun 2020: Aalborg, Denmark

## Important dates

[Run 55, B27, H15](#)  
Protocol submission deadline  
3 Jan 2019  
Slide circulation  
9 Jan 2019  
Slide return deadline  
13 Feb 2019  
Publication of results  
20 Apr 2019

## Questions

Check out our [FAQ](#) (Frequently asked questions) or [contact us](#)

# GATA3 (L50-823)

# Antibody diluents



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

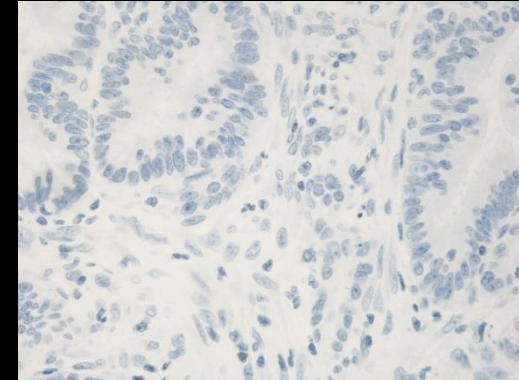
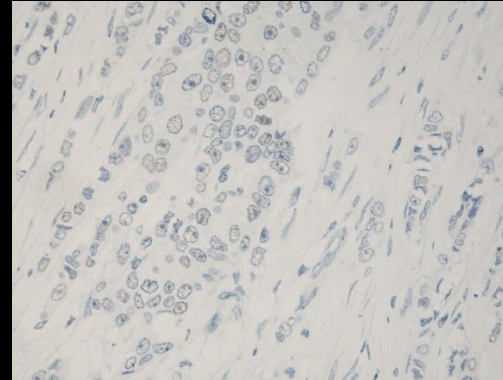
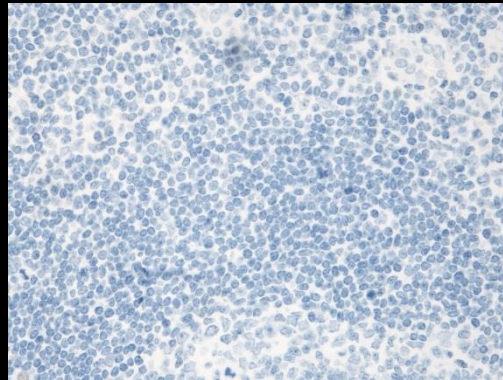
Antibody diluents

**GATA3, L50-823 (Biocare) 1:800**

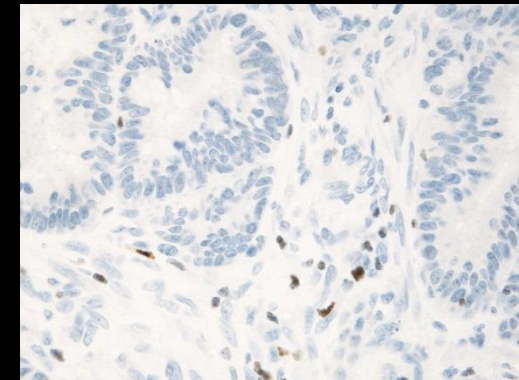
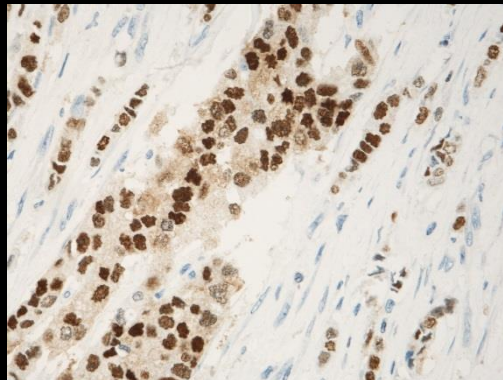
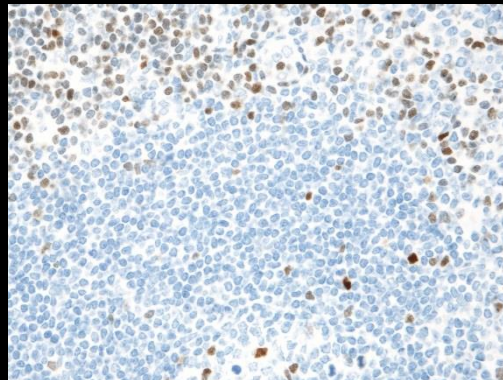
**Tonsil**

**Breast adenocarc.**

**Colon adenocarc.**



**Dako Dil. pH 7.3**



**Biocare Renoir Red pH 6.2**

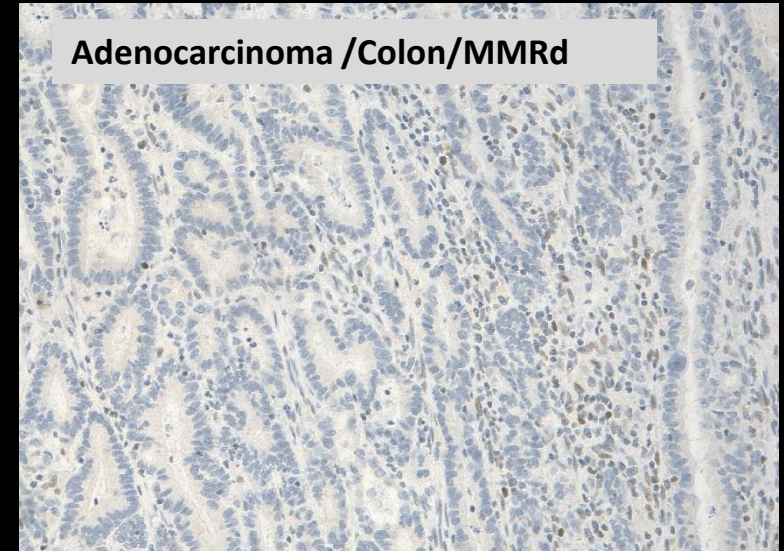
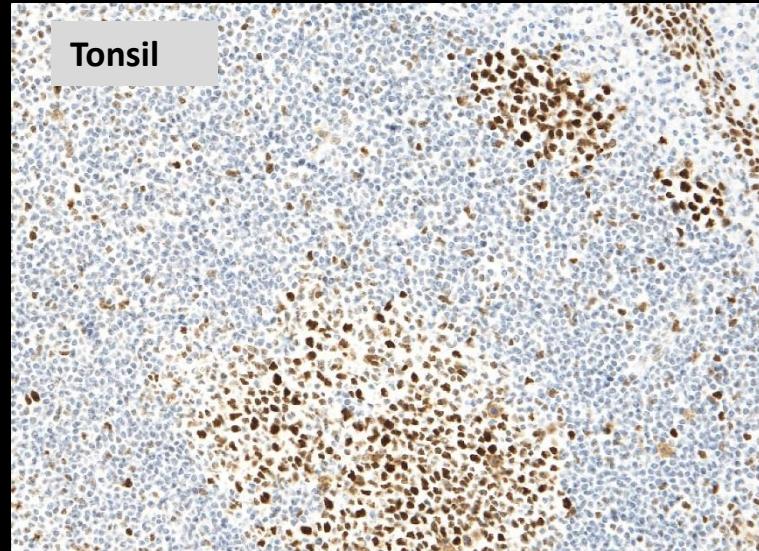
**Omnis: HIER/HIGH pH 24` at 97°C, Flex+ Mouse (10+20`)**



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

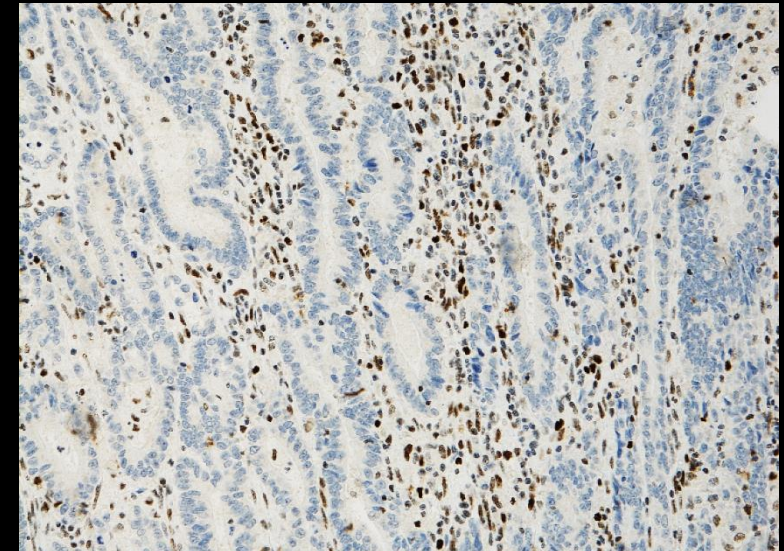
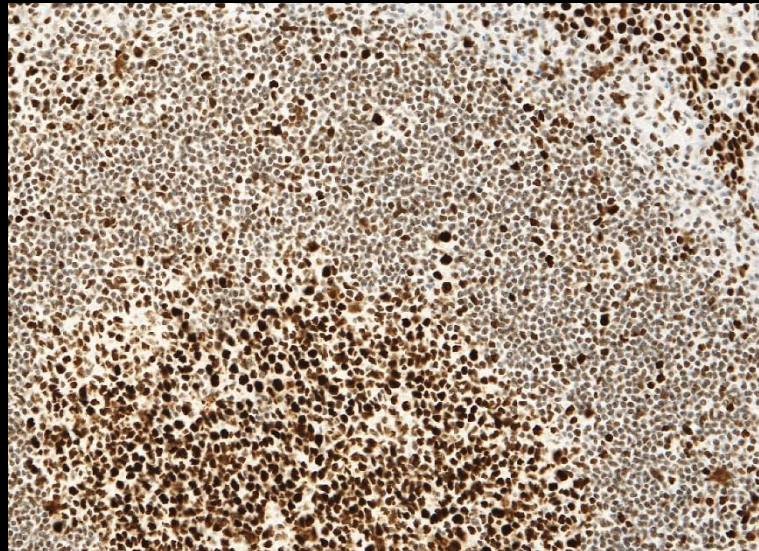
## Antibody diluents

MSH2, FE11 1:50  
Dako dil. pH 7.3



Omnis: HIER/HIGH pH 24', Flex+ Mouse (10+20')

MSH2, FE11 1:50  
Renoir Red pH 6.2

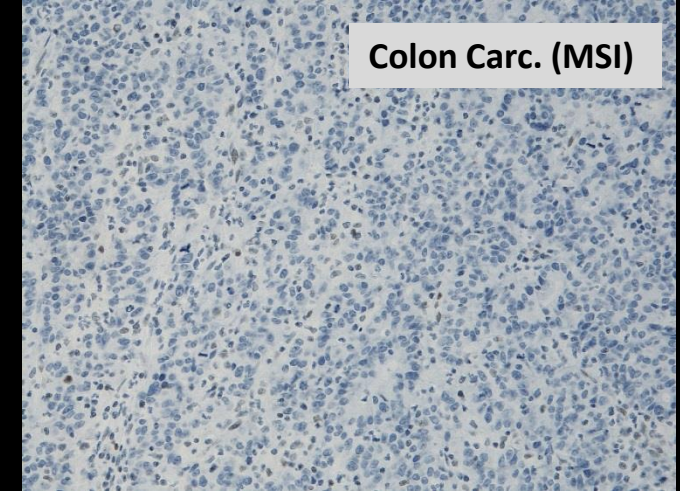
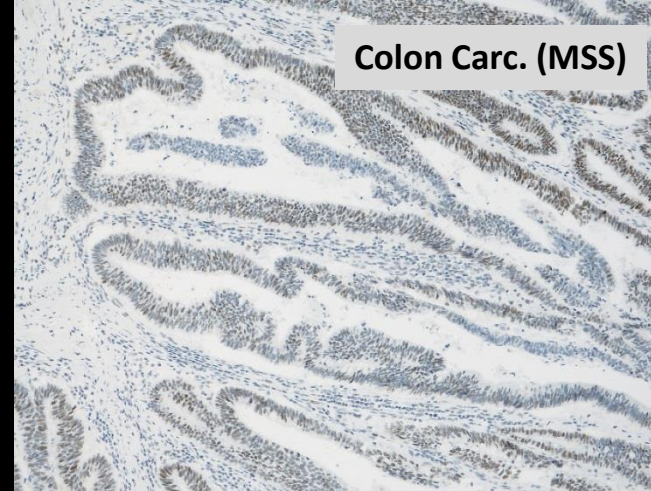
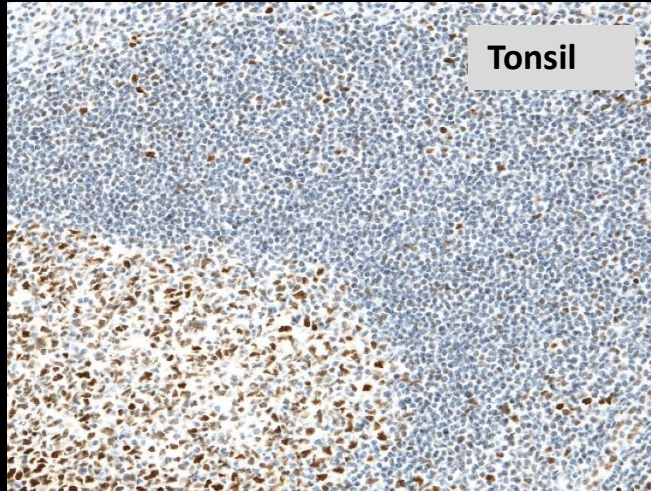




# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

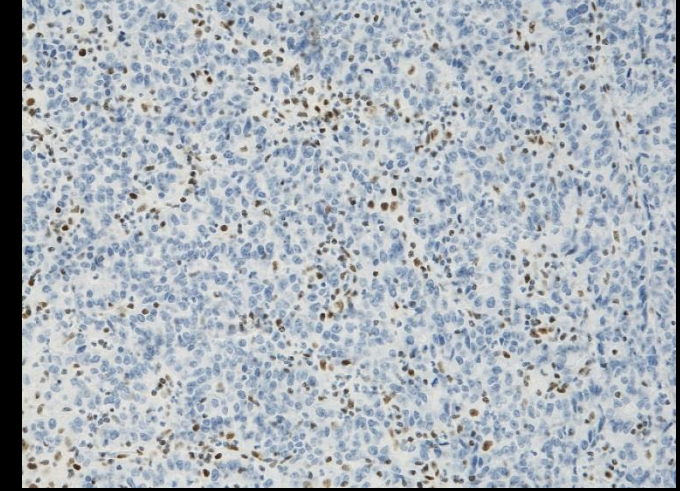
## Antibody diluents

MLH1, BC23 1:40  
Dako dil. pH 7.3



Omnis: HIER/HIGH pH 24`, Flex+ Mouse (10+20`)

MLH1, BC23 1:40  
Renoir Red pH 6.2

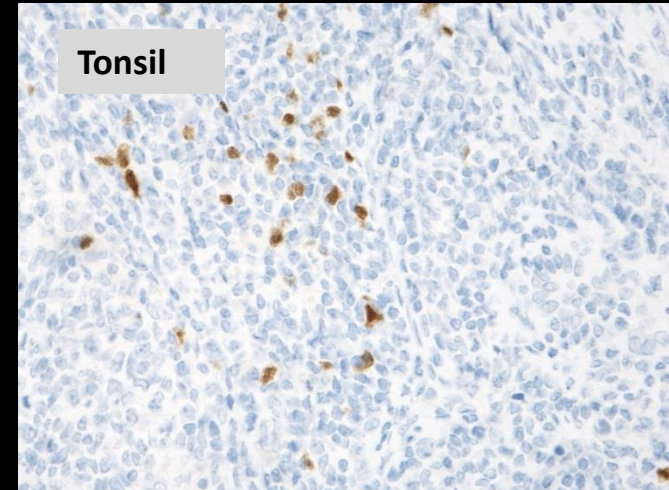
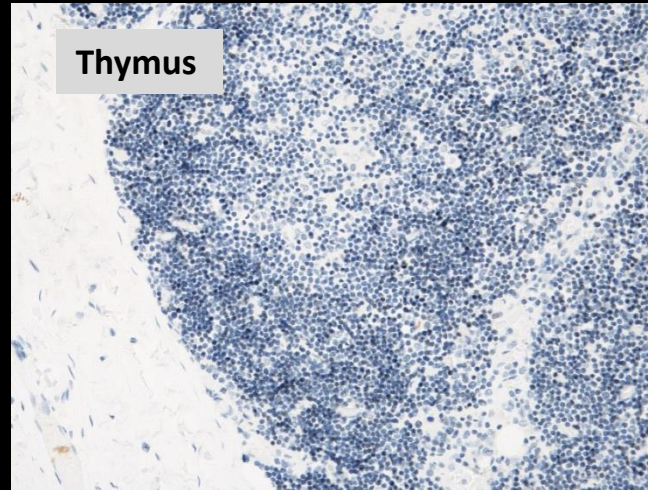




# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

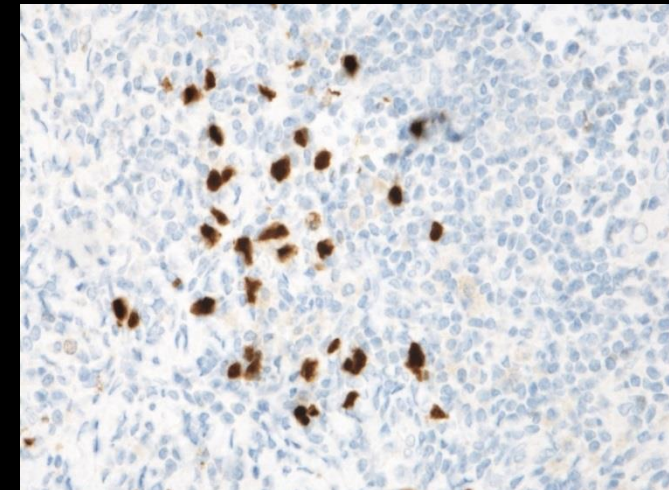
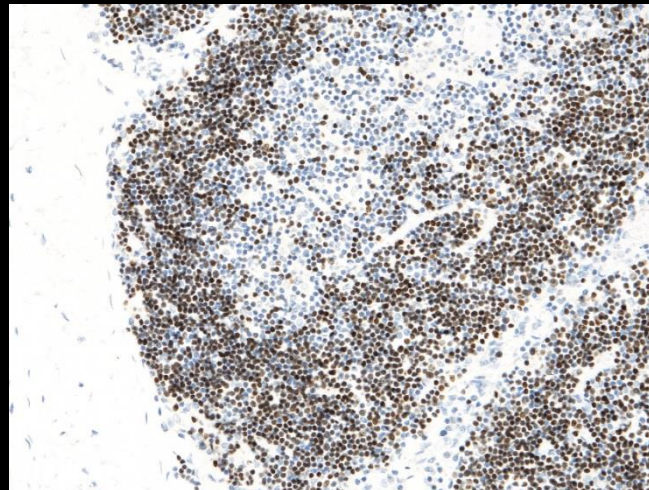
## Antibody diluents

TdT, SEN28 1:50  
Dako dil. pH 7.3



Omnis: HIER/HIGH pH 24`, Flex+ Mouse (10+20`)

TdT, SEN28 1:50  
Renoir Red pH 6.2





# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Problem: Antibody diluent

CD10 clone 56C6 (1:50)/**Dako Diluent pH 7.3**



CD10 clone 56C6 (1:50)/Renoir Red pH 6.2



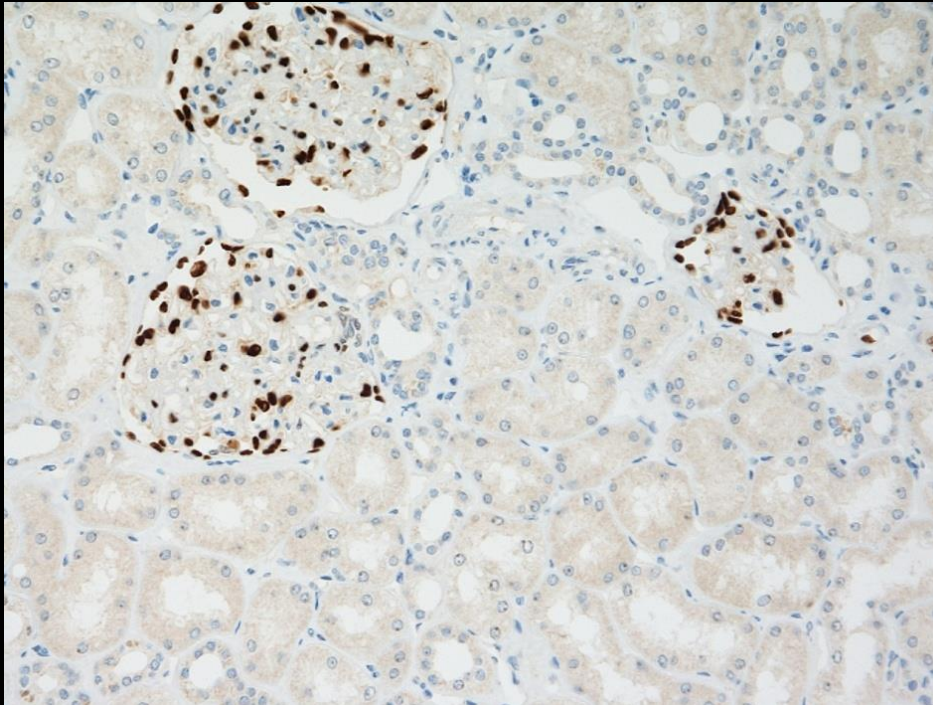
**Renoir Red is not always the best antibody diluent  
Use a “antibody diluent test battery”**



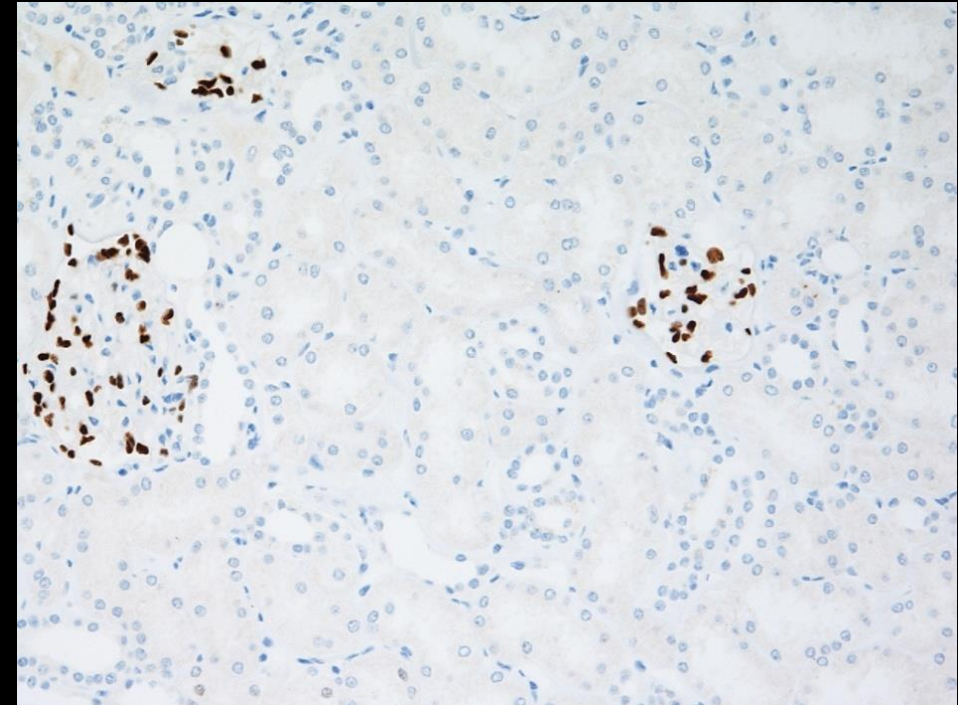
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Antibody diluents

WT1, EP122 1:25  
Renoir Red (Biocare)



WT1, EP122 1:25  
Background Sniper (Biocare)



Kidney

The choice of antibody diluent can suppress unwanted/unspecific background staining

HIER TRS pH9 (24' /97°C) + Pep © (3')



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Antibody diluents (Experience from Dept. of surgical Pathology, Region Zealand, Denmark on the Omnis)

Antibodies benefitting from dilution in Renoir Red pH 6.2 (improving signal):

ALK (1A4), CR (CAL6), CD4 (EP204), CD5 (SP19), CMYC (EP121), **GATA3 (L20-823)**, GPC3 (1G12), **IMP3 (69.1)**, **MLH1 (BC23)**, ES05 & GM011), MSH2 (G219-1129), **MSH2 (FE11)**, MSH6 (EP49), NKX 3.1 (poly), SALL4 (6E3), **PAX8 (ZR1)**, PMS2 (EP51), SOX10 (EP268), SOX11 (C1 & MRQ58), **TdT (SEN28 & EP266)**, UP-II (BC21), WT1 (WT49) and .....

Antibodies not benefitting from dilution in Renoir Red pH 6.2:

BCL2 (124), BCL6 (LN22 & PG-B6p & GI191E/A3), CR (DAK-Calret1), **CD10 (56C6)**, CD163 (MRQ26), CD21 (2G9), CD5 (4C7), ER (SP1), **HHV8 (13B10)**, Mammaglobin (304-1A5), MUC5AC (CLH2), MUC6 (CLH5), and .....

Antibodies benefitting from dilution in Background sniper (reduces background problems):

Spirochete (poly), BORR (poly), WT1 (EP122) and .....

## Proficiency testing in immunohistochemistry—experiences from Nordic Immunohistochemical Quality Control (NordiQC)

Mogens Vyberg<sup>1,2</sup> · Søren Nielsen<sup>1</sup>

Virchows Arch (2016) 468:19–29

**Table 3** Major causes of insufficient staining reactions

1. Less successful antibodies (17 %)
  - a. Poor antibodies<sup>a</sup>
  - b. Less robust antibodies<sup>b</sup>
  - c. Poorly calibrated RTUs
  - d. Stainer platform dependent antibodies
2. Insufficiently calibrated antibody dilutions (20 %)
3. Insufficient or erroneous epitope retrieval (27 %)
4. Error-prone or less sensitive visualization systems<sup>c</sup> (19 %)
- 5 Other (17 %)
  - a. Heat-induced impaired morphology
  - b. Proteolysis induced impaired morphology
  - c. Drying out phenomena
  - d. Stainer platform-dependant protocol issues
  - e. Excessive counterstaining impairing interpretation

**19% insuff.**

<sup>a</sup> Consistently gives false negative or false positive staining or a poor signal-to-noise ratio in one or more assessment runs

<sup>b</sup> Frequently giving inferior staining results, e.g., due to mouse-anti-Golgi reactions or sensitive to standard operations as blocking of endogenous peroxidase

<sup>c</sup> Biotin-based detection kit for cytoplasmic epitopes, use of detection kits providing a too low sensitivity, or use of detection kits and chromogens giving imprecise localization of the staining signals complicating the interpretation

### Problems related to the choice of the detection system:

- Provides low sensitivity
  - 2 versus 3-step multimer/polymer detection systems
- Provides low specificity and sensitivity
  - Biotin based systems

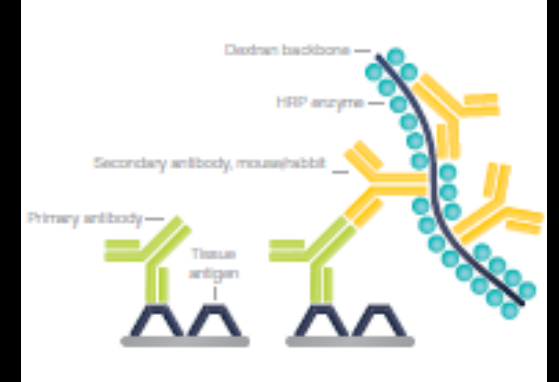
**False positive or false negative results**



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Detection systems (polymer/multimer) used by NordiQC participants

Vendor	Detection System	Detection System	Amplifier	Cat.no
	2- Step	3-step		
Dako	EnVision EnVision /Flex	Envision Flex+	Anti -Ms/Rb	K4001 K8000/10 (K5007) K8002/12 GV800 + GV809/821
Ventana	UltraView	UltraView + Amp OptiView Optiview + Amp	Anti -Ms/Rabbit Anti-Hapten Anti-Hapten + TSA	760-500 760-500 + 760-080 760-700 760-700 + 760-099
Leica		Bond Refine (PowerVision)	Anti-Ms (Rb?)	DS9800 (HRP); DS9390 (AP)
Biocare	MACH 2	MACH 3 MACH 4	Ms/Rb probe Ms probe (Rb?)	M2U522; MHRP520; RHRP520 M3M530; M3R531 M4U534
LAB Vision/TS	UltraVision One	Quanto	?	TL-125-HLJ TL-125-QHD /QHL
Immunologic	BrightVision (PowerVision)	BrightVision+	Anti-Ms/Rat (Rb ?)	DPVM (Anti-Ms)/DPVR (Anti-Rb ) DPVO (Anti-Ms/Rb/Rat) DPVB ((Anti-Ms/Rb/Rat)
Master Diag.		Quanto	?	MAD-021881QK
ZytoMed System		ZytoChem Plus (PowerVision)	Anti-Ms (Rb?)	PolHRP-100
And a few more (Advance, GTVision.....)				

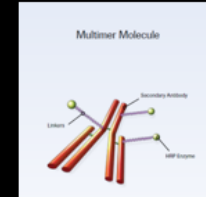


**App. 98-99% of all NordiQC participants use a polymer/multimer based detection systems**

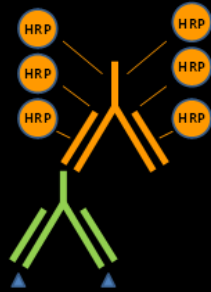
### Considerations related to the choice of detection system:

- ☐ Sensitivity
- ☐ Specificity
- ☐ Enzyme conjugate
- ☐ Blocking of endogenous activity
- ☐ Turn around time (TAT)
- ☐ Automatic platform (open or closed system)
- ☐ Price

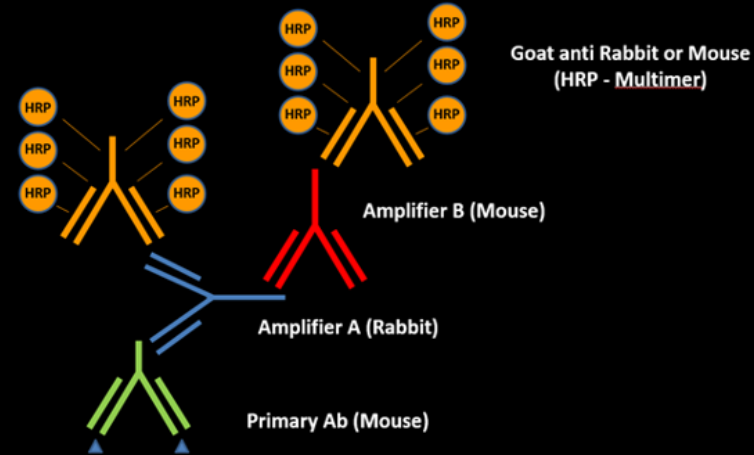
## Multimer detection systems (Ventana/Roche)



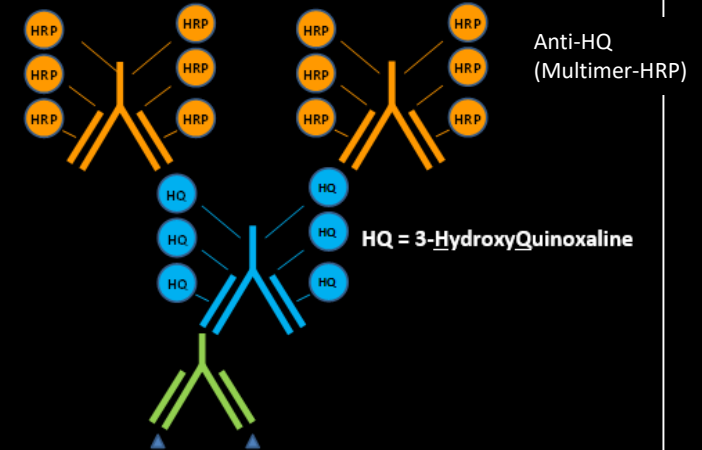
*UltraView*



*UltraView /Amp*



*OptiView*



Sensitivity

HQ-Linker  
Amplifier (A/B)

Linker (Mouse/Rabbit)  
Enhancer  
Universal Linker  
Post Blocking  
.....



**Increases sensitivity**

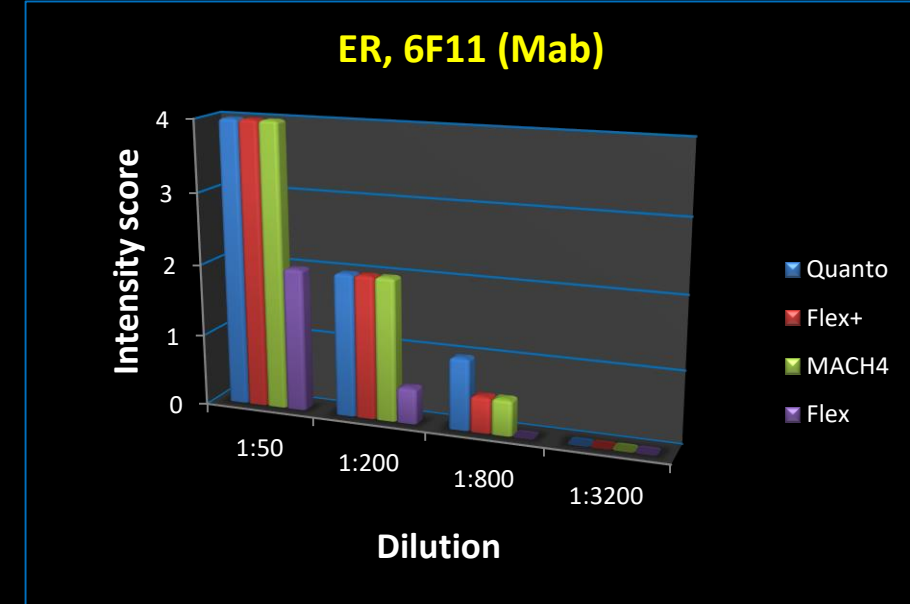
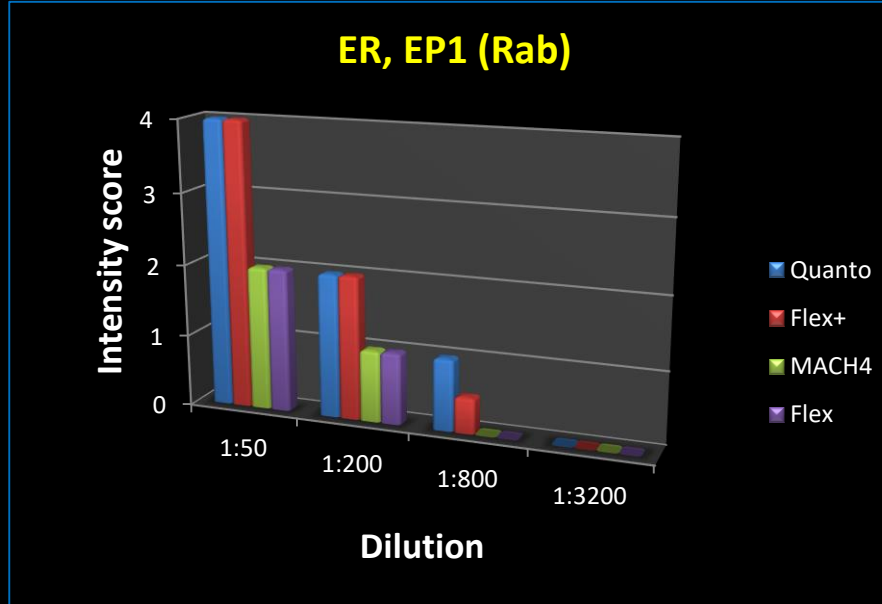
Know your detection system

Strength and weakness



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Detection systems - Performance Testing



### ER - Endpoint titration (some general remarks and important issues):

- ☐ The 3-step polymer detection systems Quanto and Flex+ - produced the overall highest intensity.
- ☐ The 3-step polymer detection system MACH4 only enhances reactions with mouse monoclonal Abs (ER,6F11).
- ☐ “Optimal staining” was highly influenced by the concentration of the primary Abs and the nature of detection system.

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Performance testing of detection systems (Vendor recommended protocol settings)

Breast tumor

ER, EP1 (Rab)

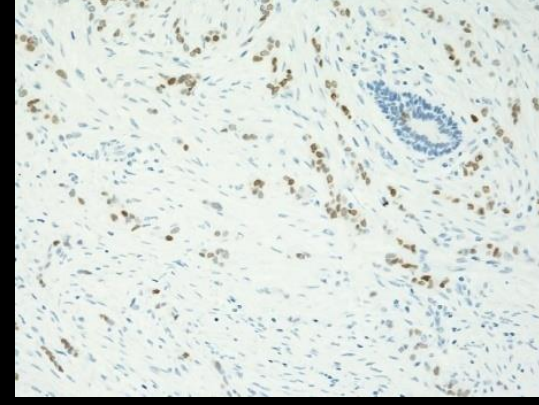
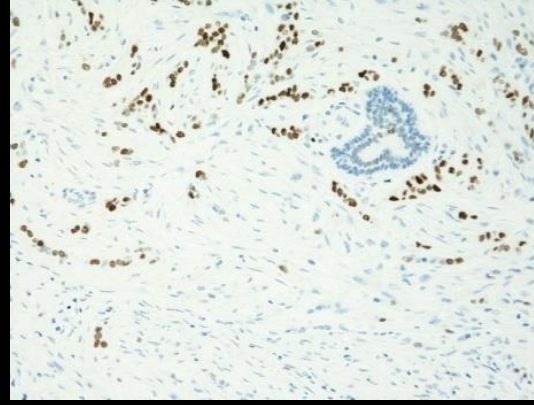
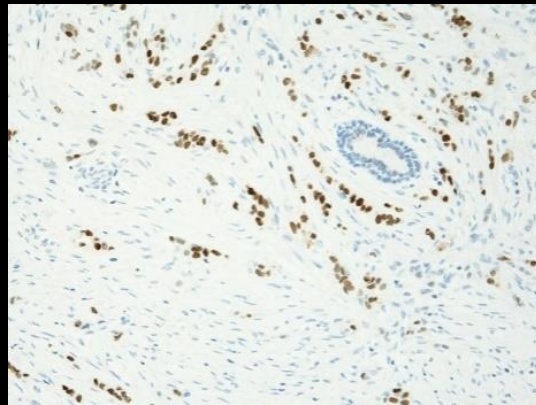
Quanto

Flex+

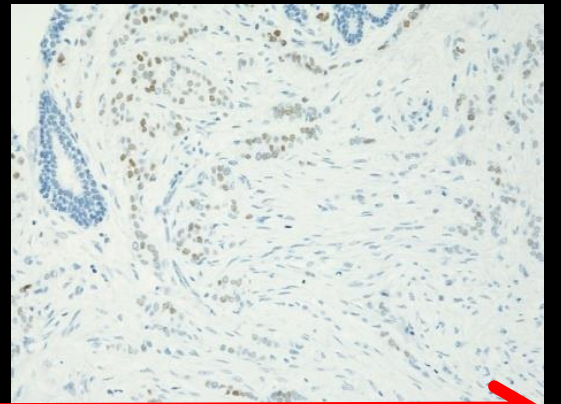
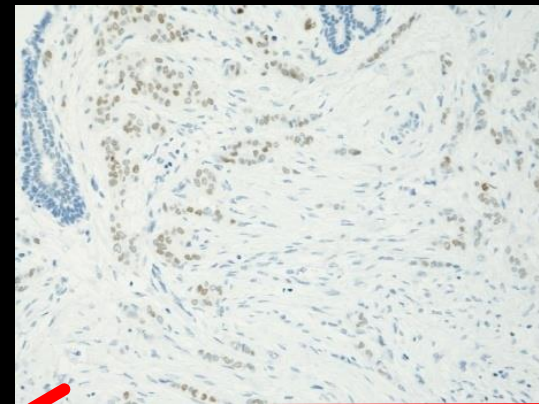
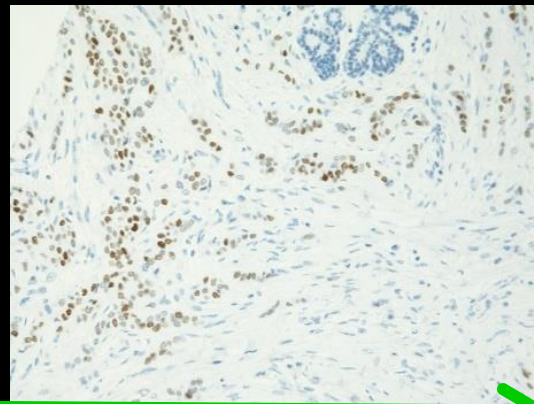
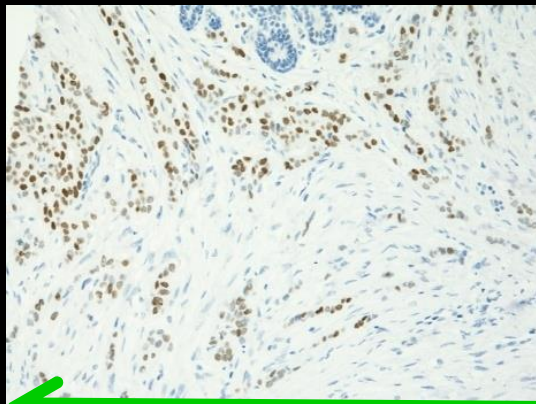
MACH4

Flex

1:50



1:200



High Intensity (3-step detection systems)

Low Intensity (2-step detection systems)



# The technical test approach – Analytical phase

Performance testing of detection systems (Vendor recommended protocol settings)

Breast tumor

ER, 6F11 (Mab)

Quanto

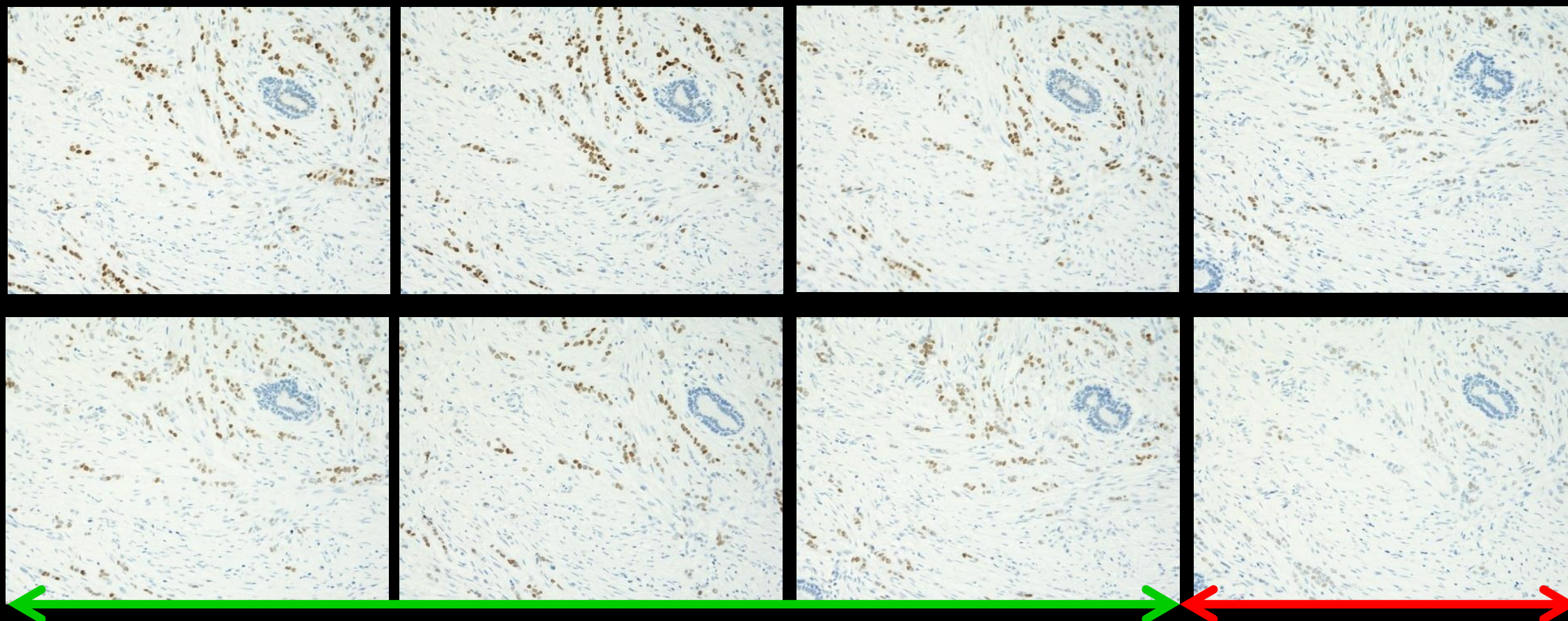
Flex+

MACH4

Flex

1:50

1:200

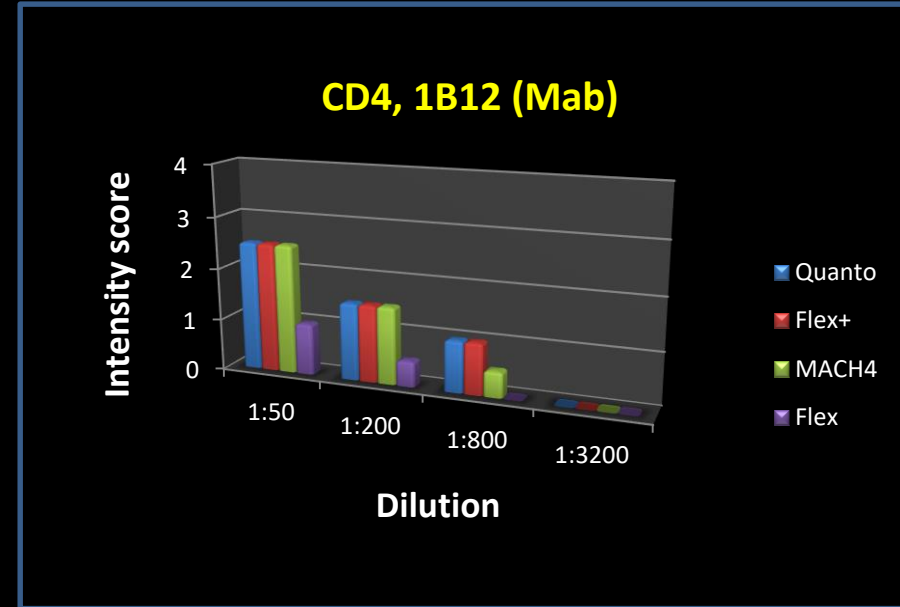
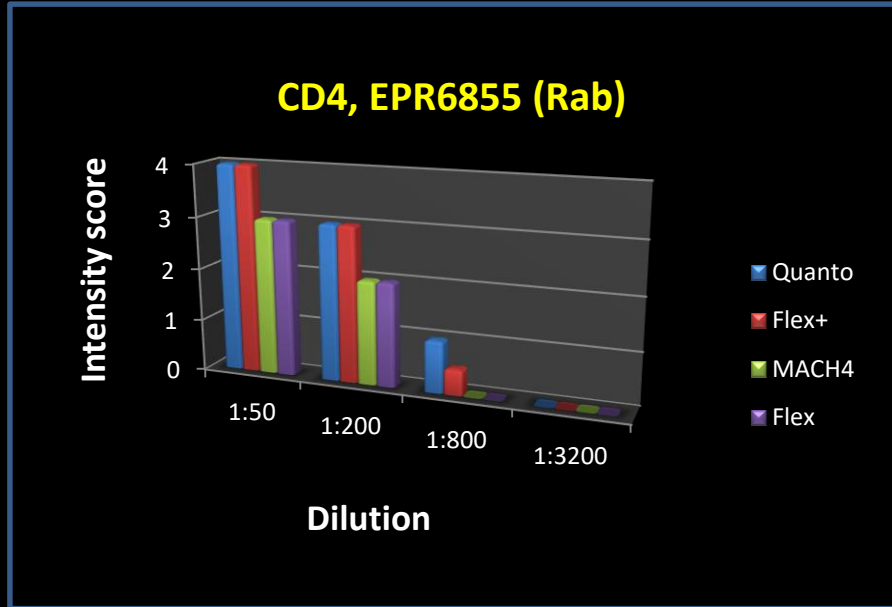


High Intensity (3-step DS)

Low Intensity (2-step DS)

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Detection systems - Performance Testing



### CD4 – Endpoint titration (some general remarks and important issues):

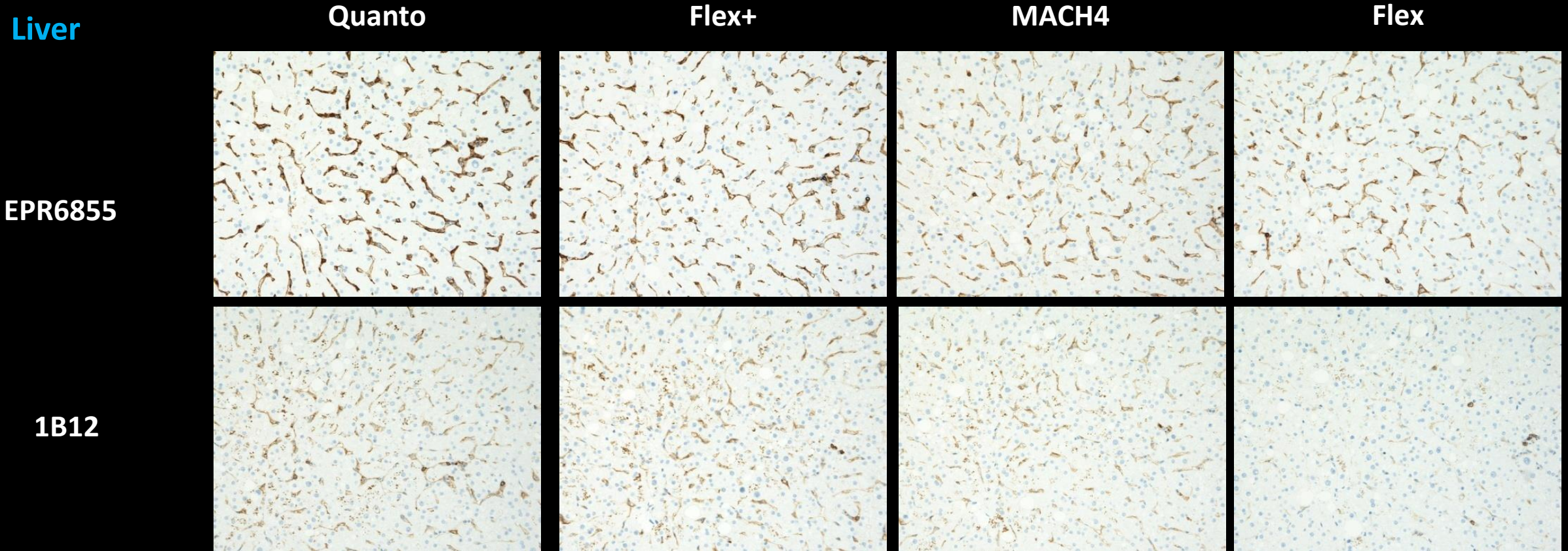
- ☐ The 3-step polymer detection systems Quanto and Flex+ - produced the overall highest intensity.
- ☐ Intensity was highly influenced by the nature of primary Ab and “optimal” staining could only be obtained with the Rab (CD4, EPR6855) used in combination with the 3- step polymer detection systems Quanto or Flex+.
- ☐ Intensity was higher with the Rab (CD4, EPR6855) at 1:50 with all of the detection systems tested compared to any intensity obtainable with the Mab (CD4, 1B12) even with the use of a 3-step polymer system ( e.g. Quanto).



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Performance testing of detection systems (Vendor recommended protocol settings)

CD4, EPR6855 (Rab, 1:50) and 1B12 (Mab, 1:50)



Strong staining of hepatic sinusoidal endothelial cells and kupffer cells using the Rab (CD4, EPR6855) with all the detection system tested (2-step or 3-step polymer systems) . Intensity is significantly reduced using the Mab (CD4, 1B12).

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Performance testing of detection systems (Vendor recommended protocol settings)

CD4, EPR6855 (Rab, 1:50) and 1B12 (Mab, 1:50)

Brain

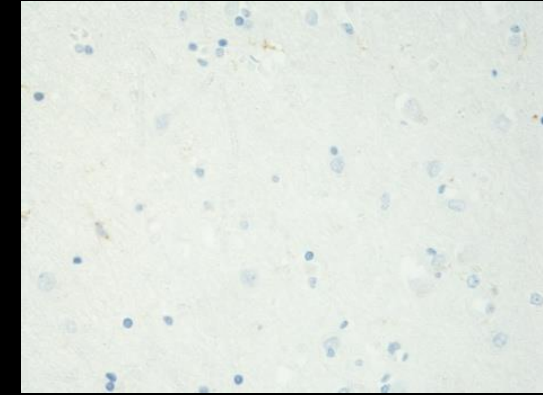
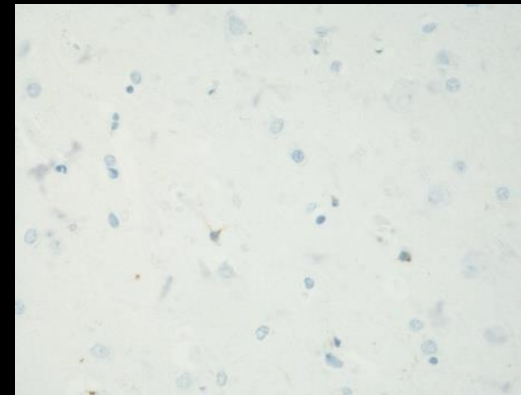
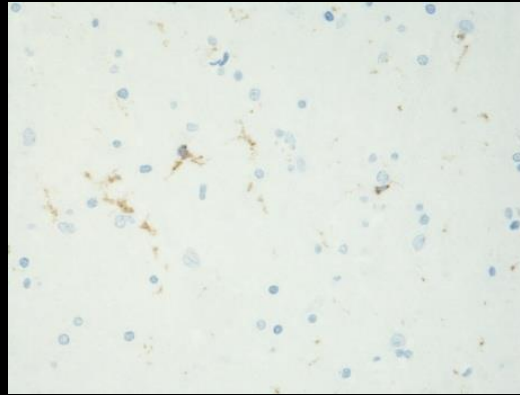
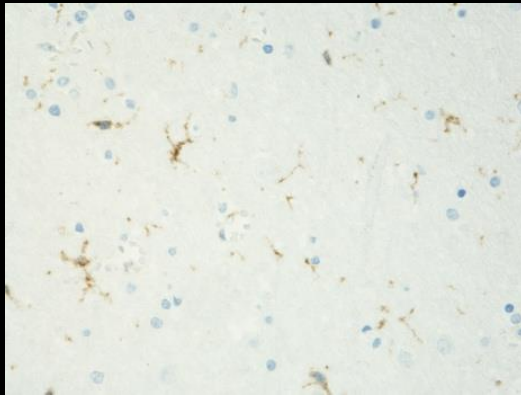
Quanto

Flex+

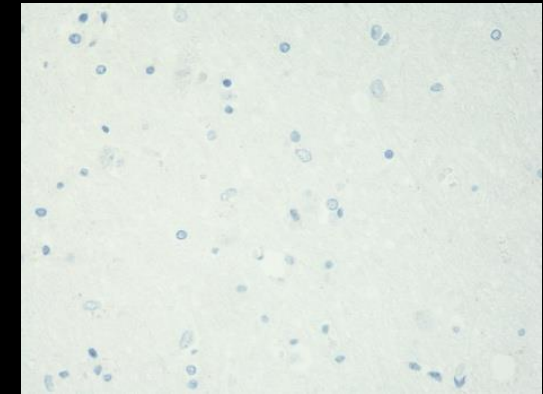
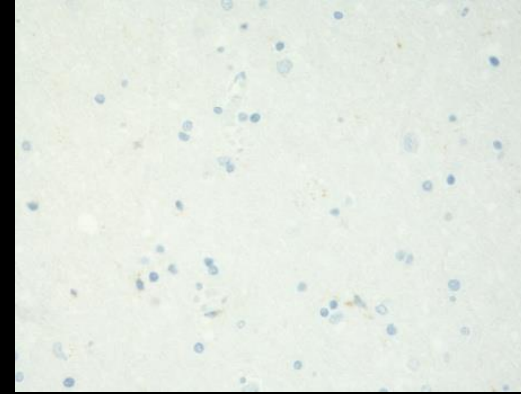
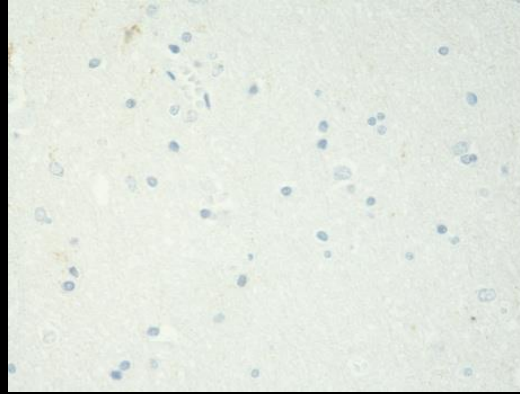
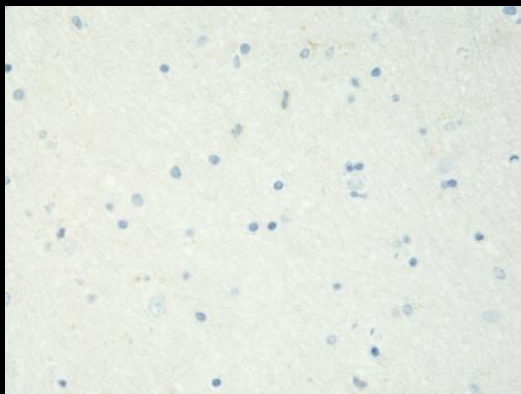
MACH4

Flex

EPR6855



1B12



Staining of microglia cells is only obtainable using the Rab (CD4, EPR6855) and the 3-step polymer detection systems Quanto or Flex+.



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Less sensitive detection systems

CYCD1, RUN 47 (2016)

Table 3. Proportion of optimal results for CyD1 for the most commonly used antibodies as concentrate on the 3 main IHC systems\*

Concentrated antibodies	Dako		Ventana		Leica	
	Autostainer / Omnis		BenchMark XT / Ultra		Bond III / Max	
	TRS pH 9.0	TRS pH 6.1	CC1 pH 8.5	CC2 pH 6.0	ER2 pH 9.0	ER1 pH 6.0
rmAb clone EP12	4/5** (80%)	-	3/5 (60%)	-	1/2	-
rmAb clone SP4	20/41** (64%)	0/1	11/31 (49%)	-	2/15 (13%)	0/1

\* Antibody concentration applied as listed above, HIER buffers and detection kits used as provided by the vendors of the respective systems.

\*\* (number of optimal results/number of laboratories using this buffer)

### Bond™ Polymer Refine Detection

Catalog No: DS9800

#### Intended Use

This detection system is for *in vitro* diagnostic use.

Bond Polymer Refine Detection is a biotin-free, polymeric horseradish peroxidase (HRP)-linker antibody conjugate system for the detection of tissue-bound mouse and rabbit IgG and some mouse IgM primary antibodies. It is intended for staining sections of formalin-fixed, paraffin-embedded tissue on the Bond™ automated system.

The clinical interpretation of any staining or its absence should be complemented by morphological studies and proper controls. They should be evaluated within the context of the patient's clinical history and other diagnostic tests by a qualified pathologist.

The Bond Polymer Refine Detection Kit must be used with laboratory best practice in the use of tissue controls. For assurance, laboratories should stain each patient sample in conjunction with positive, negative, and other tissue specific controls as needed.

#### Summary and Explanation

Immunohistochemical techniques can be used to demonstrate the presence of antigens in tissue and cells (see "Using Bond Reagents" in your Bond user documentation).

Bond Polymer Refine Detection utilizes a novel controlled polymerization technology to prepare polymeric HRP-linker antibody conjugates. The detection system avoids the use of streptavidin and biotin, and therefore eliminates non-specific staining as a result of endogenous biotin.

Bond Polymer Refine Detection works as follows:

- The specimen is incubated with hydrogen peroxide to quench endogenous peroxidase activity.
- A user-supplied specific primary antibody is applied.
- Post Primary IgG linker reagent localizes mouse antibodies.
- Poly-HRP IgG reagent localizes rabbit antibodies.
- The substrate chromogen, 3,3'-Diaminobenzidine tetrahydrochloride hydrate (DAB), visualizes the complex via a brown precipitate.
- Hematoxylin (blue) counterstaining allows the visualization of cell nuclei.

Using Bond Polymer Refine Detection in combination with the Bond automated system reduces the possibility of human error and inherent variability resulting from individual reagent dilution, manual pipetting and reagent application.

The detection system Bond Refine acts by nature as a 2-step polymer system for detection of rabbit polyclonal or rabbit monoclonal primary antibodies

Only enhances reactions with mouse primary antibodies due to the Post Primary IgG linker (Rabbit antibody)

# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Less sensitive detection systems

### CD10, RUN 39 (2013)

Table 2. Optimal results for CD10 using concentrated antibodies on the 3 main IHC systems\*

Concentrated antibodies	Dako		Ventana		Leica	
	Autostainer Link / Classic		BenchMark XT / Ultra		Bond III / Max	
	TRS pH 9.0	TRS pH 6.1	CC1 pH 8.5	CC2 pH 6.0	ER2 pH 9.0	ER1 pH 6.0
mAb clone 56C6	64 % 14/22**	0 % 0/1	67 % 35/52	-	95 % 19/20	0 % 0/1

\* Antibody concentration applied as listed above, HIER buffers and detection kits used as provided by the vendors of the respective platforms.

\*\* (number of optimal results/number of laboratories using this buffer)

Protocols based on 2-step polymer/multimer detection systems, provided a sufficient staining result in 91% (n=42/46), and in 52% (n=24/46) an optimal result was obtained. If the same protocol settings were applied with a 3-step polymer/multimer based detection system (EnVision FLEX+ (Dako), Optiview (Ventana) or Bond Refine (Leica)), sufficient staining results were seen in 100% (n=58/58), and 86% (n=50/58) was evaluated as optimal. This is also reflected in the high proportion of optimal results (95 %) using the mAb clone 56C6 as concentrate on the Leica IHC platforms (table 2), on which a 3-step polymer based detection system is used as standard.

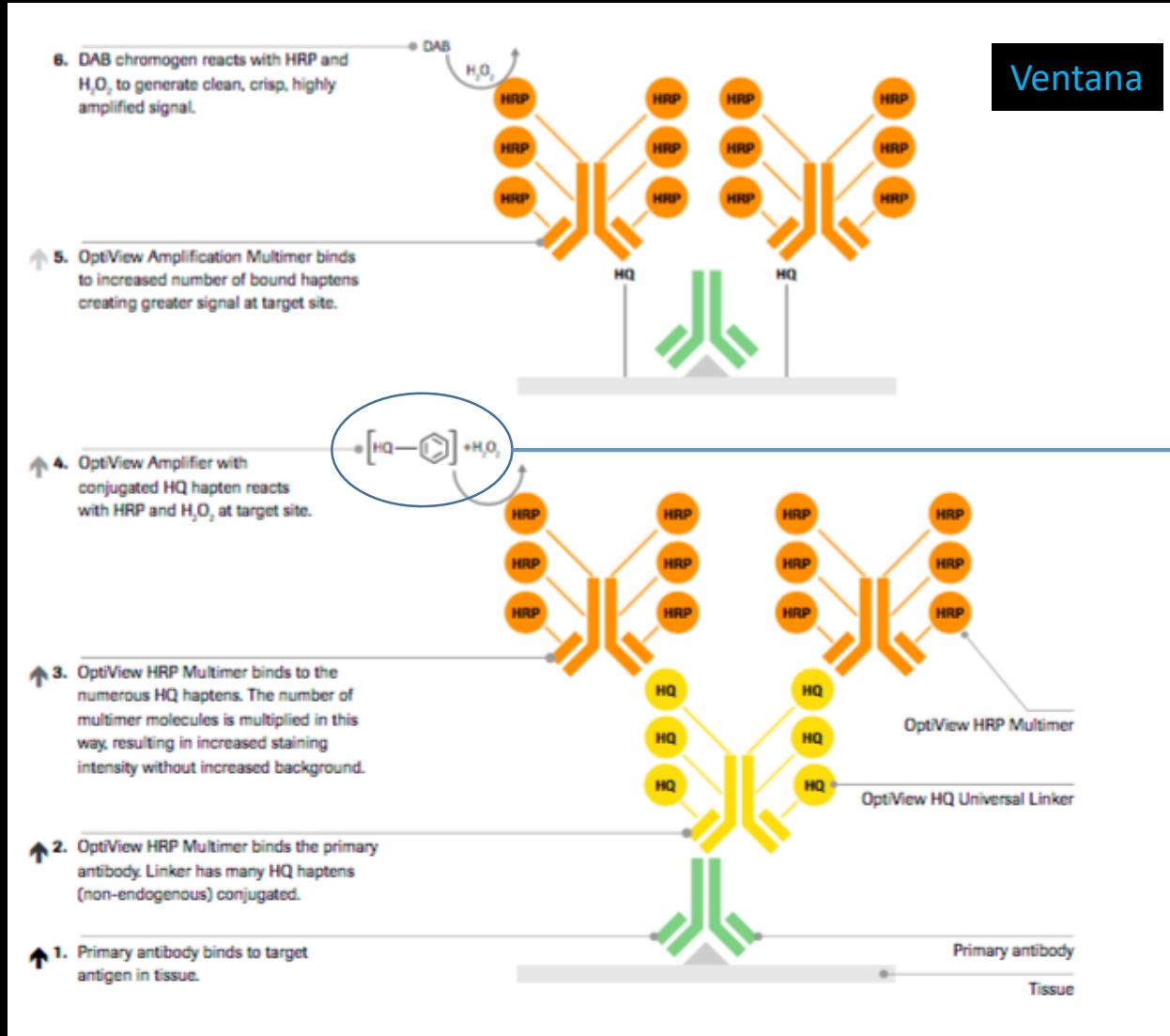
2 & 3-step detection system (UV/OV or Flex/Flex+)

BOND refine (3-step detection system)

LD assay ( mAb clone 56C6) HIER in alkaline buffer and optimal dil. range	Detection system	Pass Rate`s (%)	Optimal (%)
2-step polymer/multimer system	Flex (Dako) or UltraView (Ventana)	91 (42 of 46)	52 (24 of 46)
3-step polymer/multimer system	Flex+ (Dako), OptiView (Ventana) or BOND Refine (Leica)	100 (58 of 58)	86 (50 of 58)



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase



## Tyramide Signal Amplification

### Mechanism of Tyramide amplification:

- Introducing HRP (Optiview)
- Incubation with HQ-labelled Tyramide +  $H_2O_2$

**Tyramide, a phenolic compound, is converted into an short-lived extremely reactive intermediate**

- Deposit HQ molecules in close vicinity of Ab/Ag reactions

**Intermediates covalently binds rapidly to electron rich regions of adjacent proteins ( esp. tyrosine)**

- Detection of HQ with anti-HQ / HRP Multimer
- Visualization with DAB

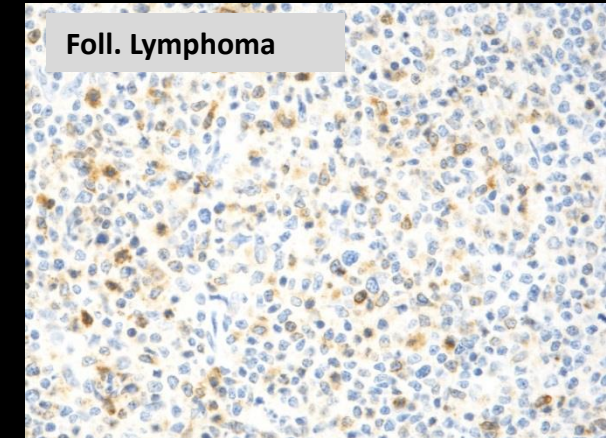
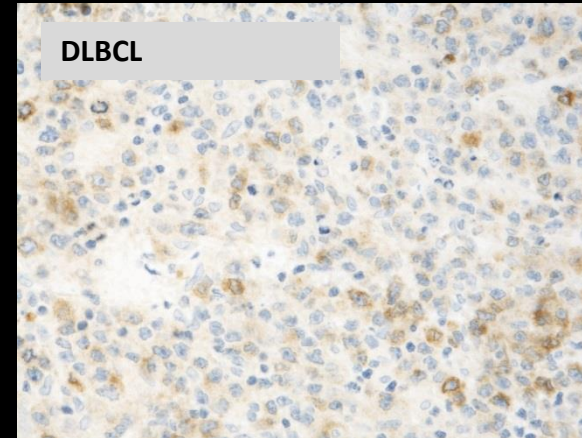
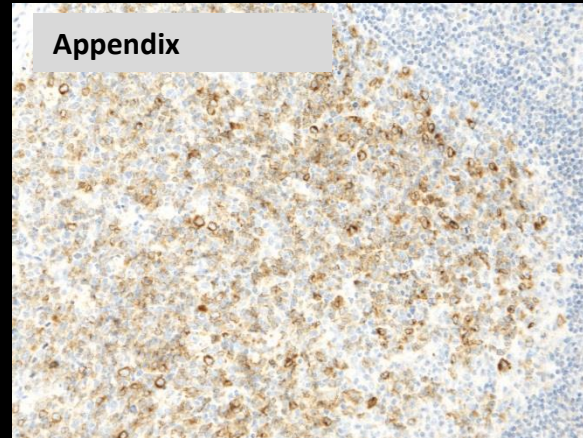
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Problem: Detection systems

**Serpin A9/GCET1 (clone 585302/1:2000RR or RAM341/1:200)**

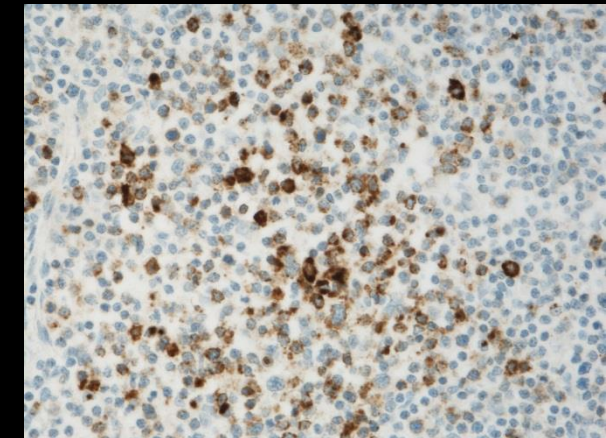
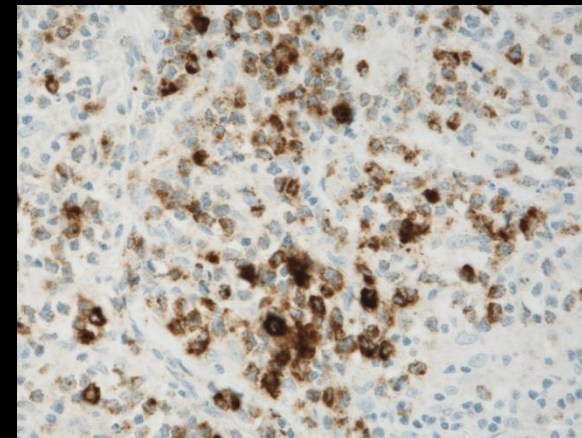
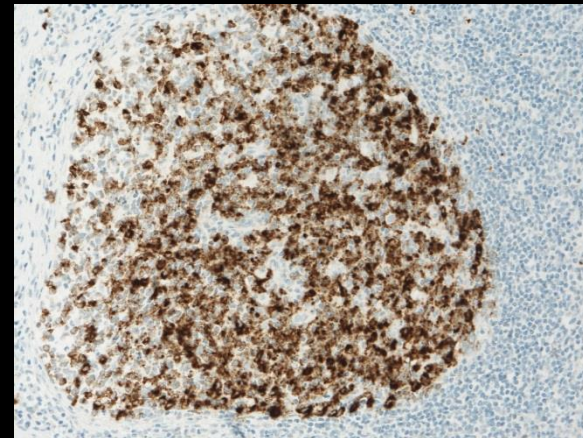
**Omnis**

HIER High pH (24`), Flex++  
(30-10-10-20`)



**Benchmark Ultra**

P3 (4`), CC1 (32`), OV+A  
(16-(8-8)-(4-4`))

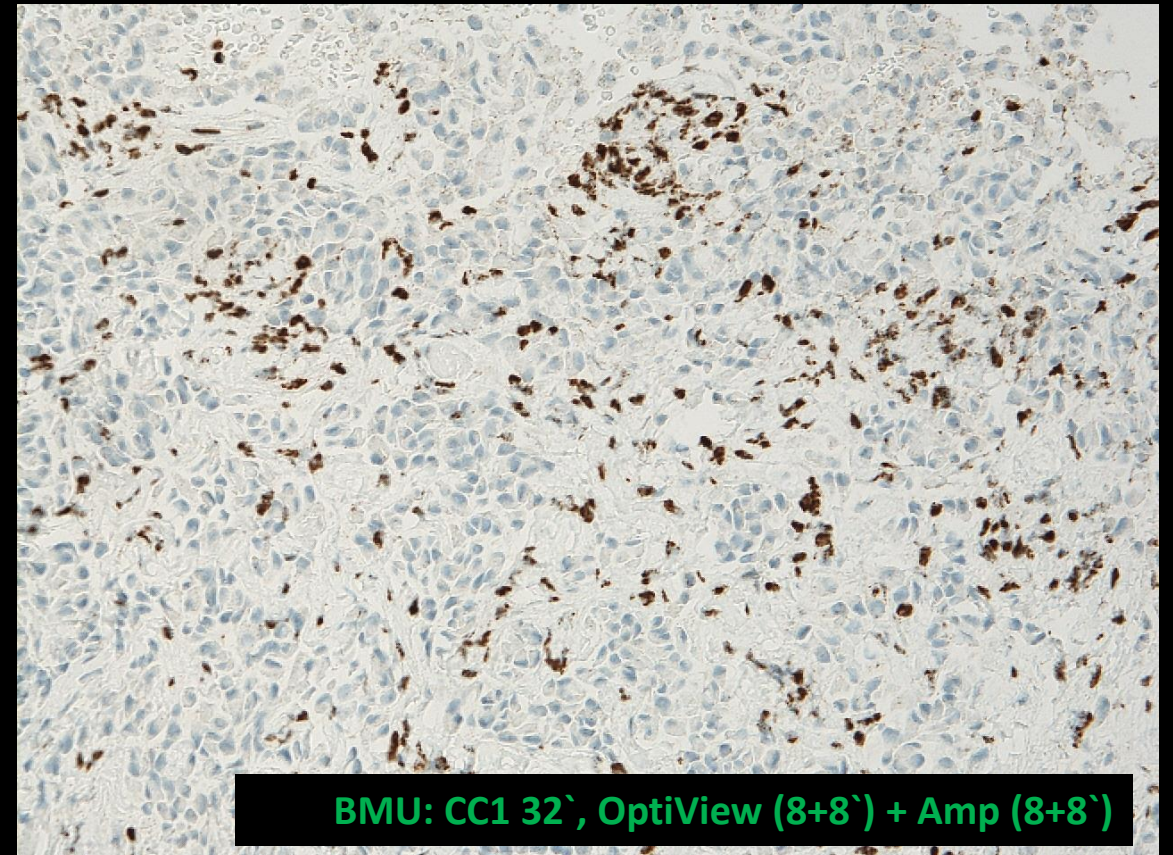
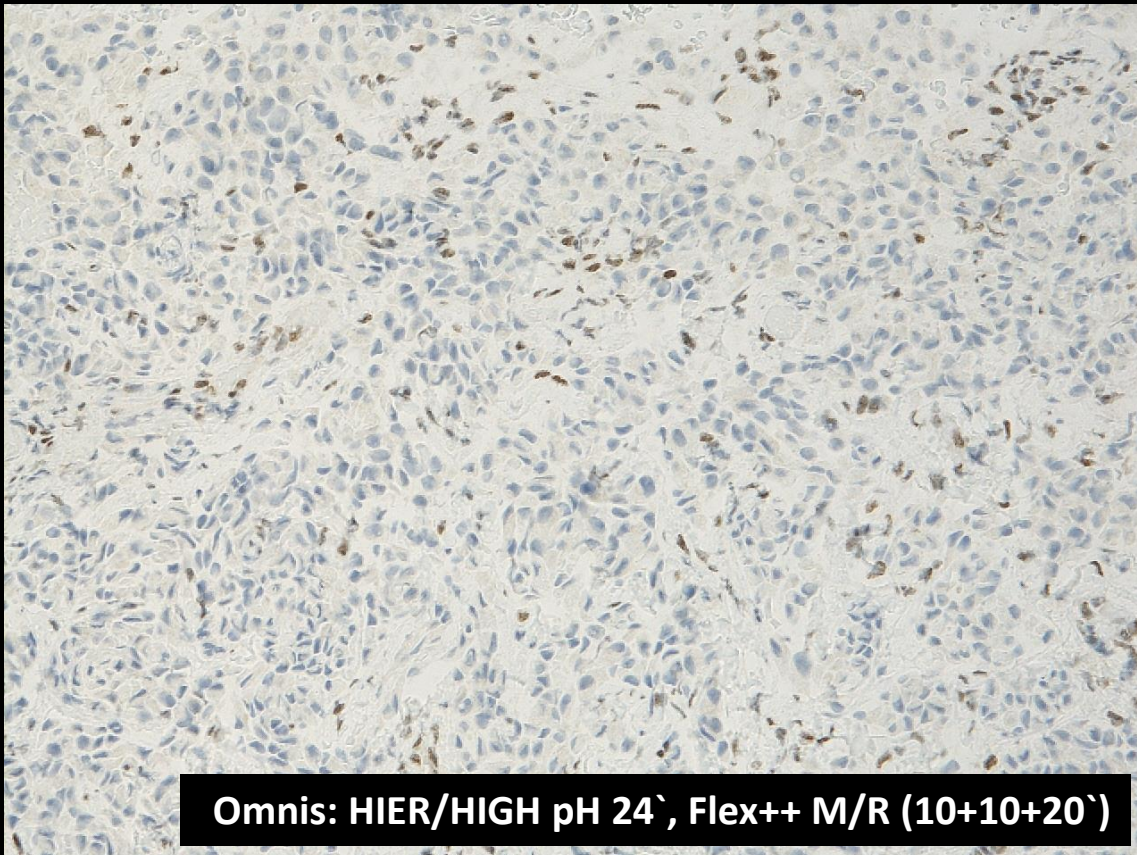




# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Problem: Detection System

## BAP1 clone C-4 (1:100) / Mesothelioma





# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Tyramide Signal Amplification (TSA)

Table 1. Antibodies and assessment marks for lu-ALK, run 45

Concentrated antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff. <sup>1</sup>	Suff. OPS <sup>2</sup>
mAb clone <b>5A4</b>	46	Leica/Novocastra						
	3	Thermo/NeoMarkers						
	2	Monosan						
	1	Abcam	24	16	13	1	74%	81%
	1	Biocare						
	1	Zytomed						
mAb clone <b>ALK1</b>	8	Dako	0	0	3	5	0%	-
mAb clone <b>OTI1A4</b>	5	ORIGENE	4	1	0	0	100%	100%
rmAb clone <b>D5F3</b>	21	Cell Signaling	18	2	1	1	91%	95%
	1	PrimeBioMed						
rmAb clone <b>SP8</b>	2	Thermo/NeoMarkers	0	0	1	1	-	-
Ready-To-Use antibodies								
mAb clone <b>5A4 PA0306</b>	3	Leica/Novocastra	0	1	2	0	-	-
mAb clone <b>5A4 API3041</b>	1	Biocare	1	0	0	0	-	-
mAb clone <b>5A4 MAB-0281</b>	1	Maixin	1	0	0	0	-	-
mAb <b>5A4 MAD-001720QD</b>	1	Master Diagnostica	0	0	0	1	-	-
mAb <b>ALK1 IR641</b>	15	Dako	0	0	4	11	0%	-
mAb clone <b>ALK1 790/800-2918</b>	10	Ventana	0	1	6	3	10%	-
mAb clone <b>ALK1 204M-18</b>	1	Cell Marque	0	0	0	1	-	-
mAb clone <b>ALK1 GA641</b>	1	Dako	0	0	0	1	-	-
rmAb clone <b>D5F3 790-4794</b>	47	Ventana	41	4	2	0	96%	96%
rmAb clone <b>D5F3 790-4843 (CDx assay)</b>	4	Ventana	3	0	1	0	-	-
Unknown	1	Unknown	1	0	0	0	-	-
Total	176		93	25	33	25	-	
Proportion			53%	14%	19%	14%	67%	

1) Proportion of sufficient stains (optimal or good).

2) Proportion of sufficient stains with optimal protocol settings only, see below.

## Lu-ALK

For certain type of markers, the TSA system provides optimal results but.....

### Ready-To-Use antibodies and corresponding systems

mAb clone **5A4**, product no. **API3041**, Biocare, IntelliPATH: One protocol with an optimal result was based on HIER using Borg Decloaker pH 9.5 (Biocare) in a pressure cooker (efficient heating time 15 min. at 110°C), 30 min. incubation of the primary Ab and MACH 4 Universal HRP-Polymer (M4U534) as detection system.

mAb clone **5A4**, product no. **MAB-0281**, Maixin, manual staining:

One protocol with an optimal result was based on HIER using Citrate buffer pH 6 (efficient heating time 2 min. at 120°C), 60 min. incubation of the primary Ab. and KIT-5230 (Maixin) as detection system.

rmAb clone **D5F3** product no. **790-4794**, Ventana, BenchMark GX, XT and Ultra:

Protocols with optimal results were typically based on HIER using Cell Conditioning 1 (efficient heating time 32-92 min.), 16-44 min. incubation of the primary Ab. and OptiView (760-700) + amplification kit (760-099) as detection system. Using these protocol settings, 45 of 47 (96%) laboratories produced a sufficient staining result.

rmAb clone **D5F3** product no. **790-4843**, CDx Assay, Ventana, BenchMark XT and Ultra:

Protocols with optimal results were typically based on HIER using Cell Conditioning 1 (efficient heating time 72-92min.), 16-36 min. incubation of the primary Ab. and OptiView (760-700) + amplification kit (760-099) as detection system. Using these protocol settings, 3 of 3 (100%) laboratories produced a sufficient staining result.

TSA



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

*The Histochemical Journal* 31: 195–200, 1999.  
© 1999 Kluwer Academic Publishers. Printed in the Netherlands.

## Concentration dependent and adverse effects in immunohistochemistry using the tyramine amplification technique

Michael Mengel<sup>1</sup>, Martin Werner<sup>2</sup> & Reinhard von Wasielewski<sup>1,\*</sup>

<sup>1</sup>*Institut für Pathologie der Medizinischen Hochschule Hannover, Carl Neuberg-Str. 1, D-30625 Hannover, Germany*

<sup>2</sup>*Institut für Pathologie der Technischen Universität München, Germany*

*\*Author for correspondence*

Received 8 September 1998 and in revised form 1 December 1998

### Summary

Although the tyramine amplification technique to enhance sensitivity in immunohistochemistry has been described in numerous methodological papers, it has not yet gained access to diagnostic immunohistochemistry. This is mainly due to problems and pitfalls occurring in adaptation of this method to routine application.

In this study a monoclonal antibody and a polyclonal antiserum (pan-cytokeratin and anti-myoglobin) were tested in tissues with different amounts of epitopes, using a checkerboard table and testing a total of 133 different dilution combinations of both the tyramide solution and the primary antibodies.

The specific tissue investigated, i.e. the amount of accessible epitope to be detected and the applied concentration of the tyramide solution mainly influenced the staining reaction. Several pitfalls such as an uneven distribution of the staining or dramatic overstaining (paradoxical overstaining) must be considered to achieve optimal results.

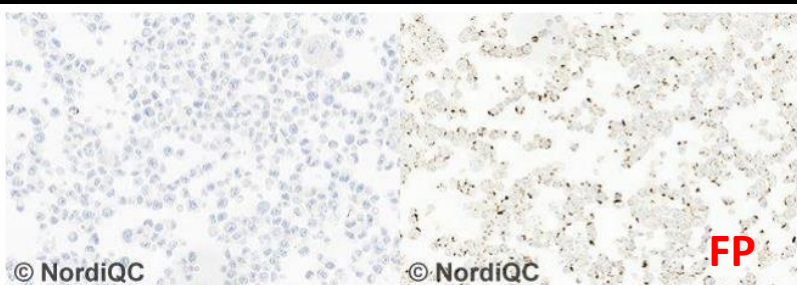
In conclusion, our data confirm methodological studies that the tyramine amplification technique is a powerful method to enhance immunohistochemical sensitivity. However, for reliable daily practice several pitfalls of the technique have to be circumvented.

## TSA enhancement

Difficult to control causing:

- False positive staining
- Weak or false negative staining, unbalanced reaction of primary Ab and target epitopes giving a yes or no answer
- Uneven distribution of the TSA reaction product





© NordiQC

Fig. 6a  
Optimal ALK staining of the cell line without ALK rearrangement using same protocol as in Figs. 1a – 5a. All cells are unstained.



© NordiQC

Fig. 6b  
Aberrant ALK staining of the cell line without ALK rearrangement using the Ventana Ready-To-Use system based on the rmAb clone D5F3, prod. No. 790-4794. The vast majority of cells show an intracytoplasmic dot-like staining reaction.

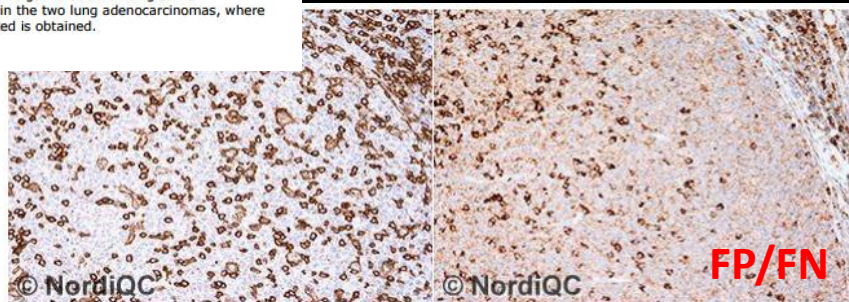
This aberrant result was seen in a high number of protocols based on this system (31 of 51). As the system otherwise provided the results expected in all the histological specimens tested, the unexpected result in the cell line was not encountered in the final assessment score.

The positive staining reaction most likely was due to the tyramide based amplification step interacting with an unknown sequence in the cell lines. As such negative reagent controls omitting the primary antibody revealed same reaction in both cell lines included. Also compare with Figs. 7a and 7b using same protocol/system in the two lung adenocarcinomas, where the result expected is obtained.

FP

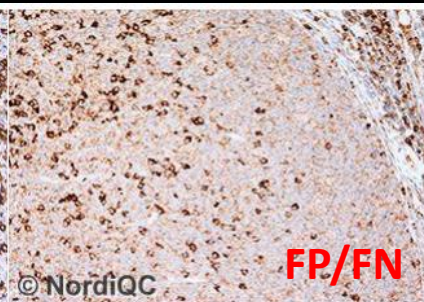
## OptiView + TSA

Lu-ALK



© NordiQC

Fig. 1a (x200)  
Optimal CD4 staining of the tonsil using the rmAb clone SP35 as a concentrate, HIER in an alkaline buffer (CC1 pH 8.5) and a 3-step multimer based detection system (OptiView, Ventana) - same protocol used in Figs. 2a - 4a. The inducer/helper T-cells show a strong staining reaction, while the germinal centre macrophages show a moderate and distinct membranous staining reaction - compare with Fig. 1b.

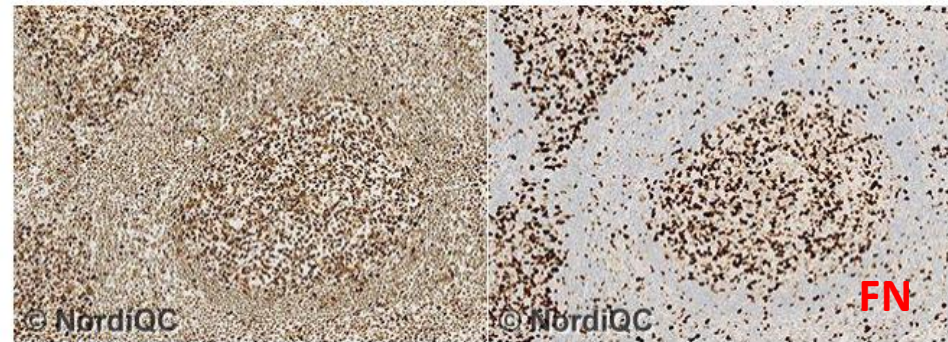


© NordiQC

Fig. 1b (x200)  
Insufficient CD4 staining of the tonsil using the mAb 4B12 as a concentrate, HIER in an alkaline buffer (CC1 pH 8.5) and a multimer based detection system (OptiView with Tyramide amplification, Ventana) - same protocol used in Figs. 2b-4b. The protocol provided a too low sensitivity, but also poor signal-to-noise ratio and false positive staining. No staining of germinal centre macrophages is seen and simultaneously B-cells are labelled. The pattern of too weak staining reaction was observed with all protocols based on the mAb 4B12 performed on the Ventana BenchMark platform - compare with Fig. 1a (same field).

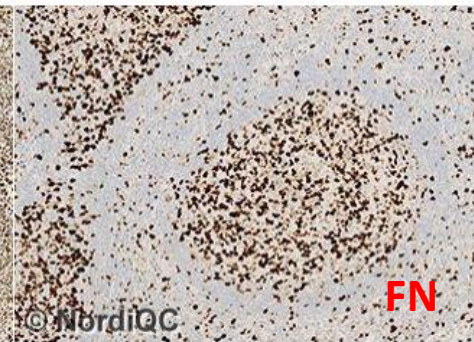
FP/FN

CD4



© NordiQC

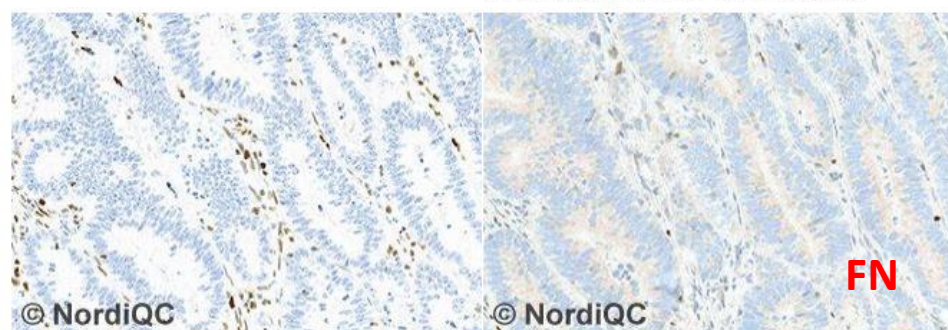
Fig. 1a (X100)  
Optimal MSH6 staining reaction of the tonsil using the rmAb clone EP49, optimally calibrated, HIER in an alkaline buffer and a 3-step multimer based detection system (OptiView, Ventana). Virtually all mantle zone B-cells show a moderate and distinct nuclear staining reaction, while the germinal centre B-cells show a strong nuclear staining reaction. Also compare with Figs. 2a - 4a, same protocol.



© NordiQC

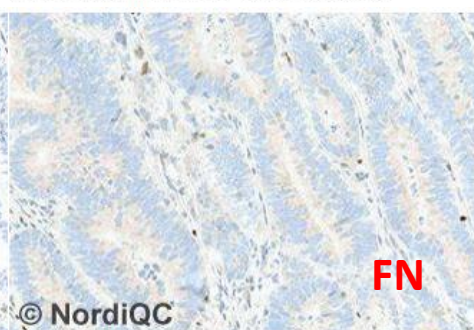
Fig. 1b (X200)  
Insufficient MSH6 staining reaction of the tonsil using the rmAb clone EP49 with a protocol providing a too low sensitivity- same field as in Fig. 1a. Only the germinal centre B-cells are distinctively demonstrated, while mantle zone B-cells expressing low level MSH6 virtually are unstained. The protocol was based on OptiView + amplification (Ventana) and the combination of a too low titre of the primary Ab and the use of tyramide based amplification provided an inadequate balance of the staining reaction. Also compare with Figs. 2b - 4b, same protocol.

FN



© NordiQC

Fig. 4a (X200)  
Optimal MSH6 staining reaction of the colon adenocarcinoma no. 4 with loss of MSH6 expression using same protocol as in Figs. 1a - 3a. The neoplastic cells are negative, while stromal cells show a distinct nuclear staining reaction serving as internal positive tissue control.



© NordiQC

Fig. 4b (x200)  
Insufficient MSH6 staining reaction of the colon adenocarcinoma no. 4 with loss of MSH6 expression using same protocol as in Figs. 1b - 3b - same field as in Fig. 4a. No nuclear staining reaction in the neoplastic cells is seen, but as only an equivocal nuclear staining reaction in the normal stromal cells is present, the staining pattern cannot reliably be interpreted. In addition a weak aberrant cytoplasmic staining reaction is seen.

FN

MSH6

The TSA detection system is not without problems and may provided either false positive or negative results.

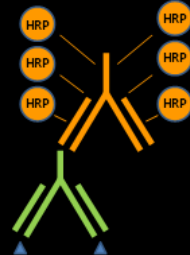
All parameters should be careful calibrated to provide optimal staining result - always possible ?



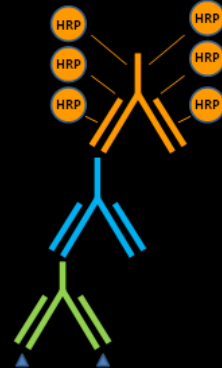
# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

## Polymer detection systems (Agilent/Dako)

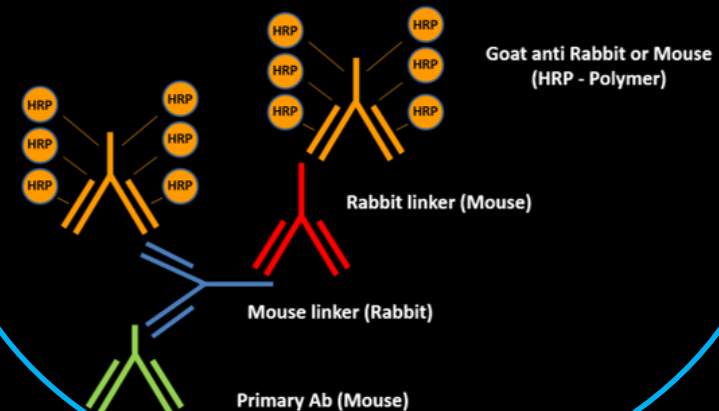
*EnVision/Flex*



*EnVision/Flex+*



*EnVision/Flex++*



Sensitivity

“New option” on the Omnis

In general, works well with rabbit primary Abs but less efficient with primary mouse primary Abs



Flex

Flex+

Flex++

# Omnis: PMS2 clone EP51 (1:300 RR) /High pH 48'

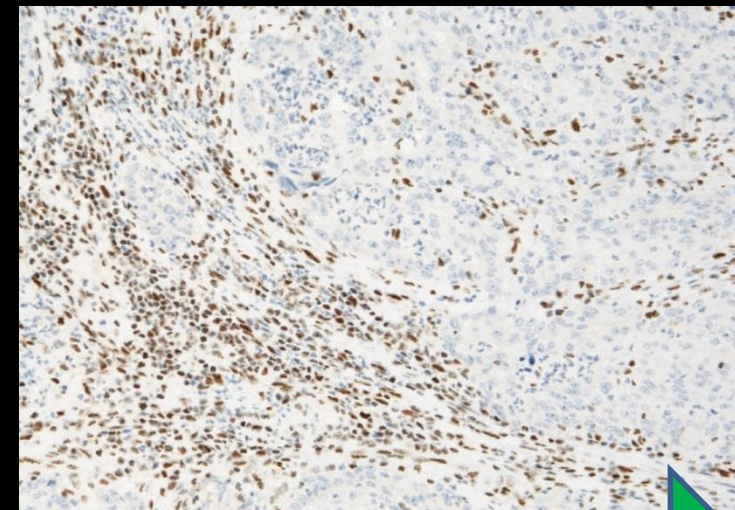
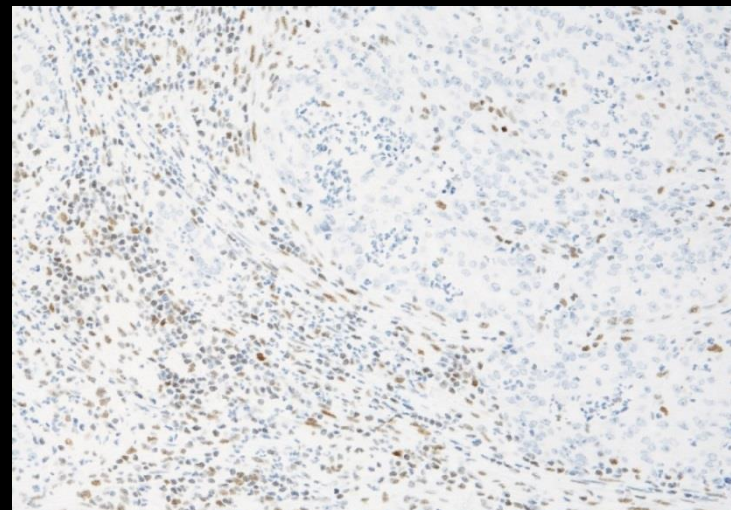
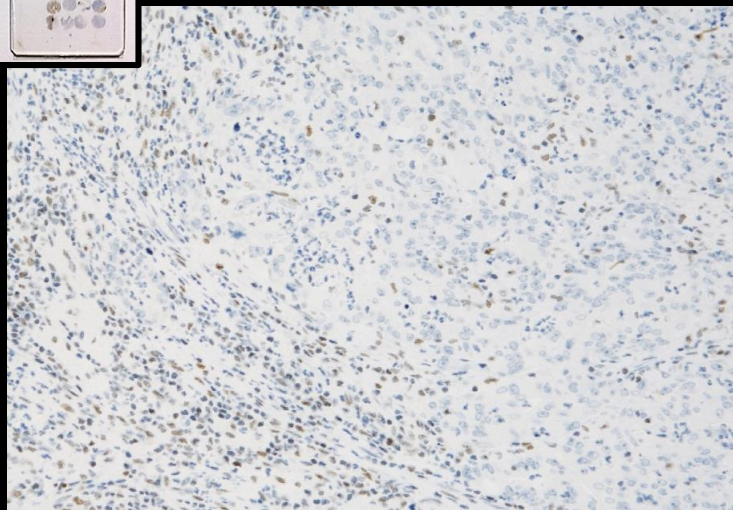
FLEX

FLEX+

FLEX++

Colon tumor

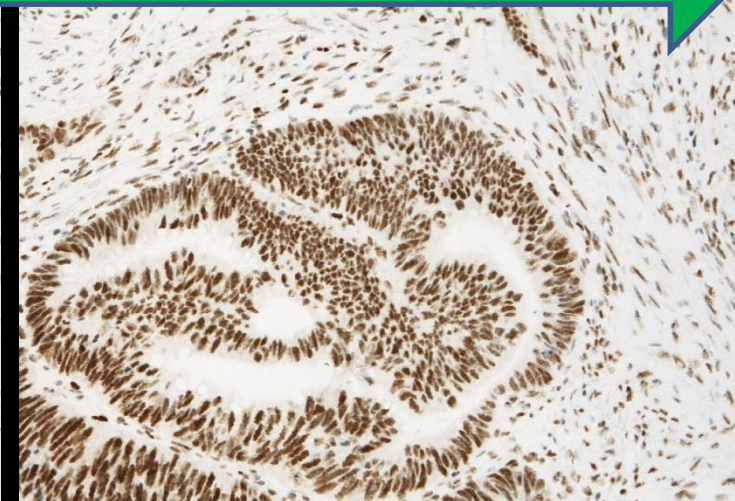
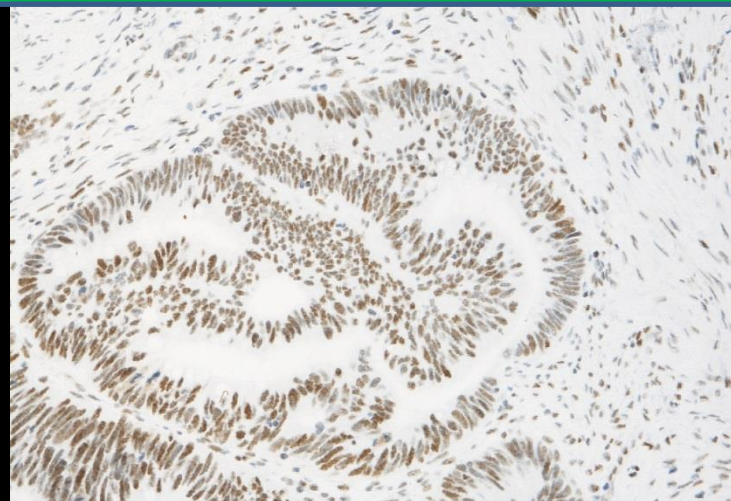
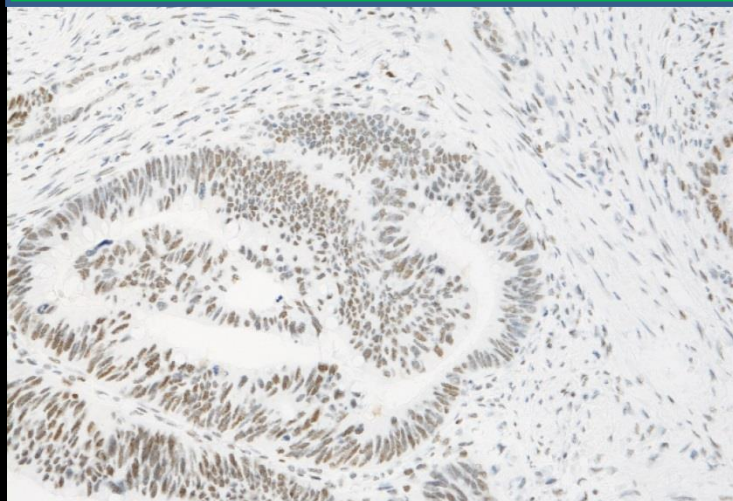
MSI



Sensitivity

Colon tumor

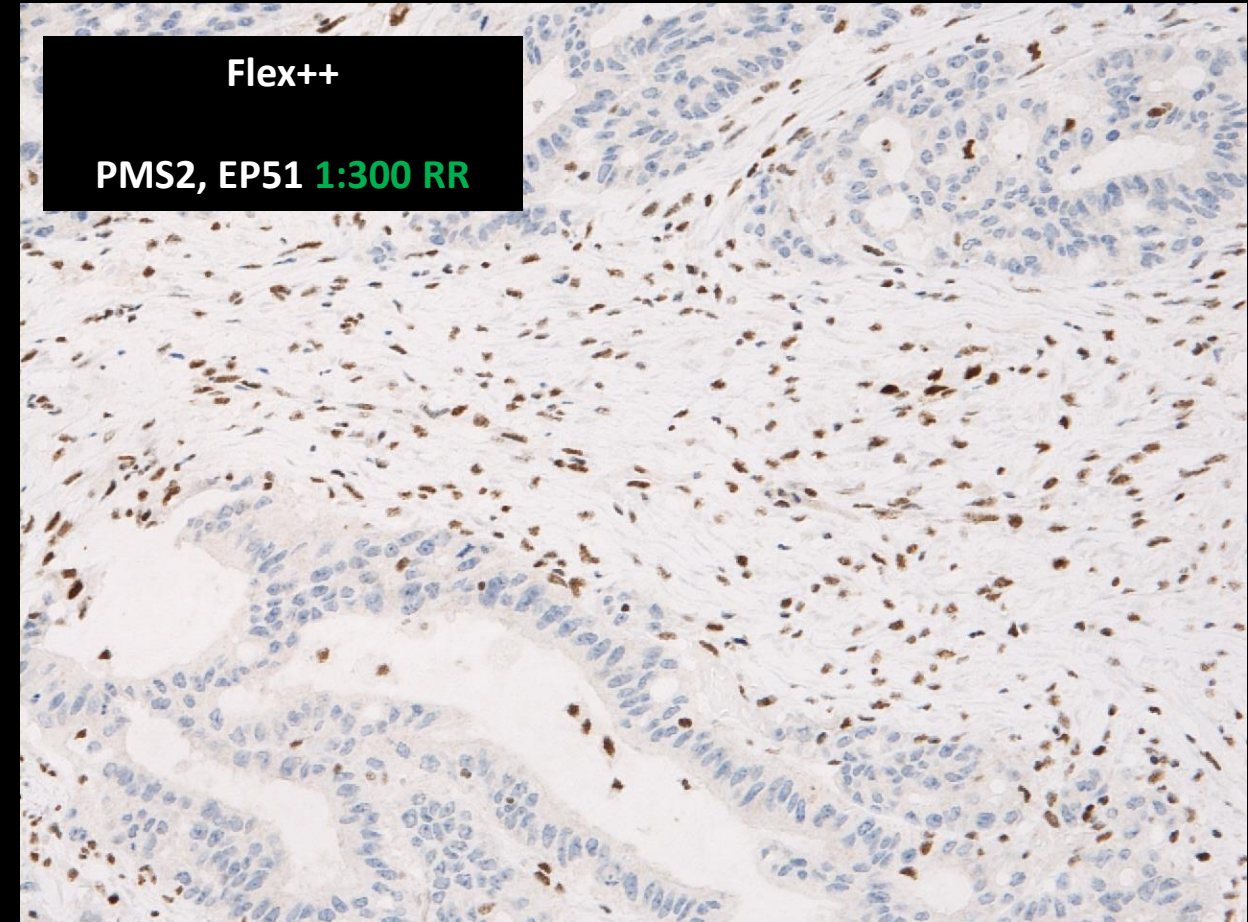
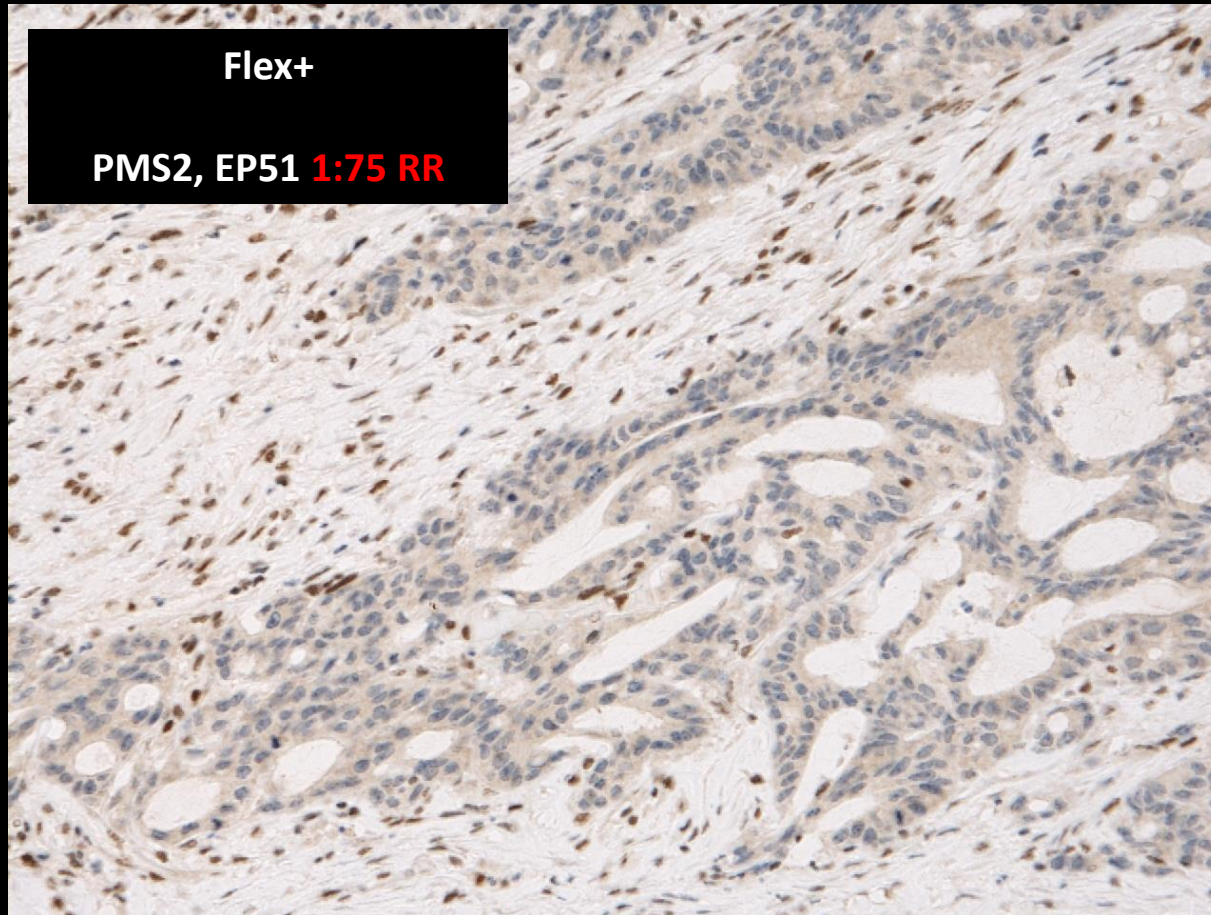
MSS





# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Colon tumor with loss of PMS2: Improves staining quality due to reduced background/noise

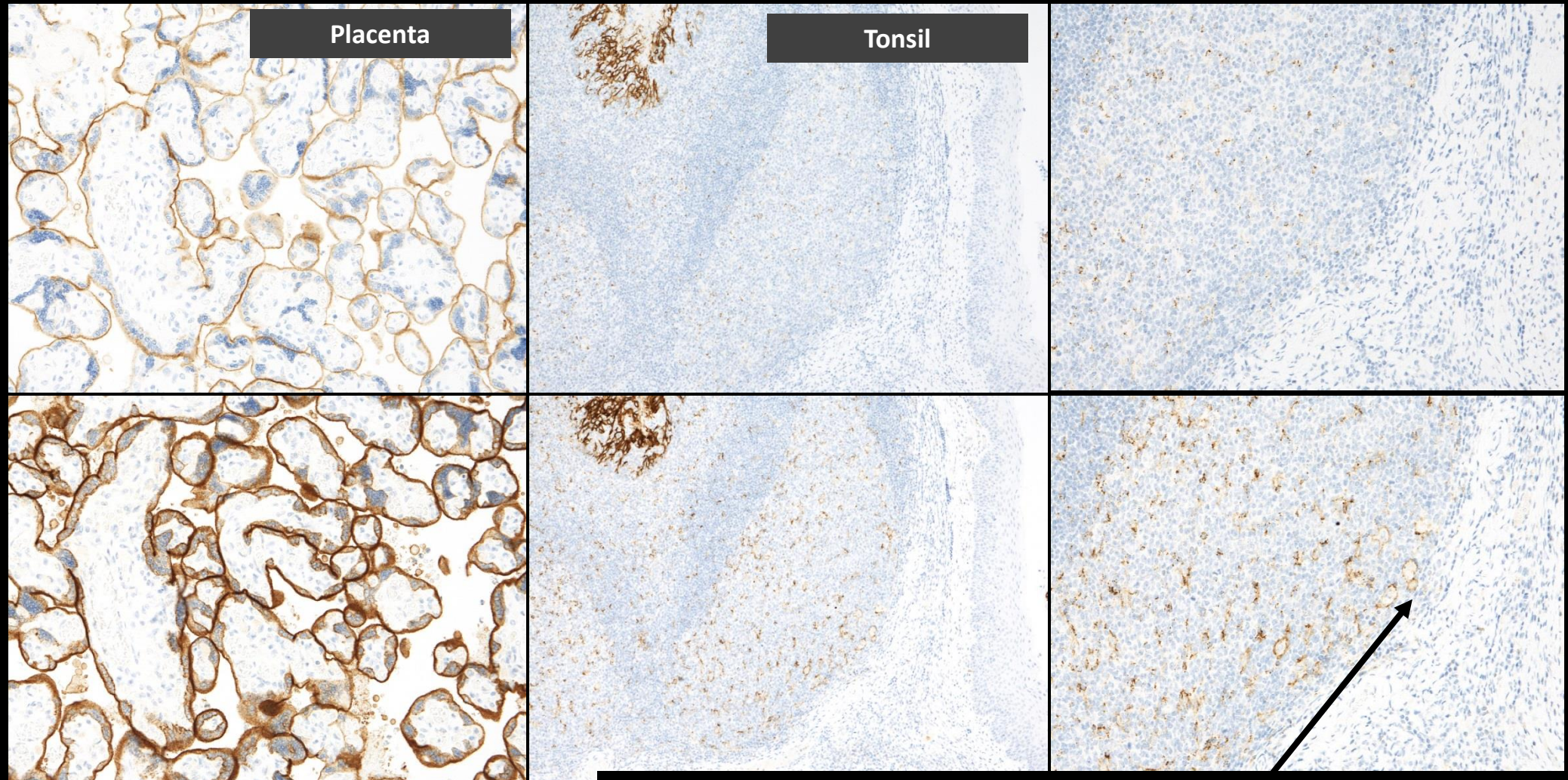


Omnis: HIER High pH 48'



## Flex++ : 4-step polymer detection system (30-10-10-20`/Omnis)

rmAb PD-L1, CAL10 (rmAb 1:30RR); HIER in High pH 48`



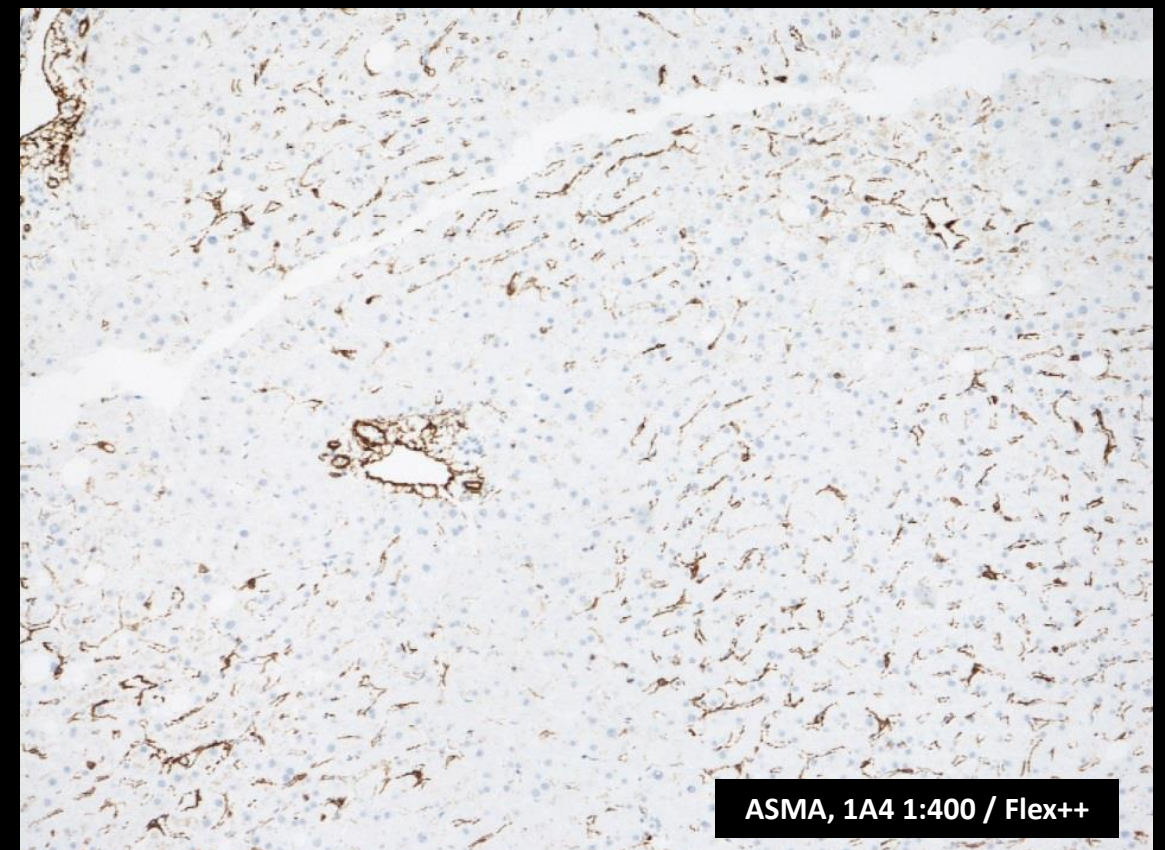
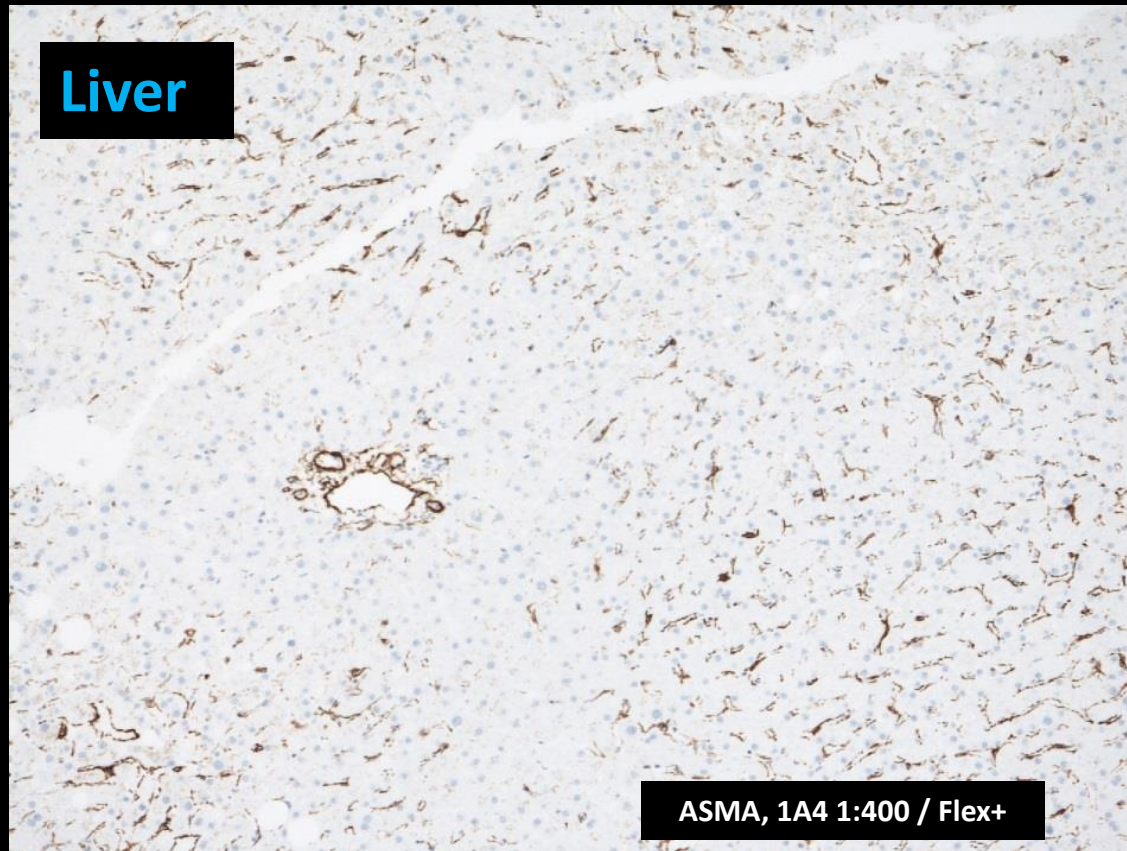
rmAbs: Enhances the sensitivity of the assay



# Technical aspects of immunohistochemistry & pitfalls - Analytical phase

Flex++ : 4-step polymer detection system (Omnis)

mAb ASMA clone 1A4; HIER in High pH 24'



Mouse monoclonal antibodies: In most cases, no improvement in signal intensity using Flex++ compared to Flex+

# The basal fundament for a technical optimal performance is :

## ❑ **Appropriate tissue fixation and processing**

## ❑ **Appropriate and efficient epitope retrieval**

- 95-98% of the Abs require HIER and app. 85-90% prefer high pH (alkaline) retrieval buffers.
- Use efficient HIER temperature and time (app. 100°C for 20 - 40min).

## ❑ **Appropriate choice of antibody / clone, diluent and dilution**

- Compare different clones / Abs against the selected antigen of interest before implementation
- Calibrate the Ab concentration carefully

## ❑ **Appropriate and specific / sensitive detection system**

- Use of a 3-step multimer/polymer system is preferable to a 2- step multimer/polymer system
- Don't use biotin-based detection systems

## ❑ **Appropriate choice of control material**

- Focus on Immunohistochemistry Critical Assay Performance Controls (iCAPCs)



Thank you for your attention

