Uitnodiging Symposium voor Pathologie Maandag 25 september 2023

## **On-slide controls for diagnostic IHC**

## **IHC Critical Assay Performance Controls**

Søren Nielsen, Director, NordiQC



# Agenda and focus areas

- What is recommended and best practice for IHC controls in diagnostic IHC?
- What are the potentials and limitations for the use of IHC controls ?
- How can IHC controls be used by laboratories and IHC stakeholders?
  - How to use IHC controls to monitor assay consistency.
  - How to use IHC controls to adress inter and intra test accuracy (e.g. EQA).

The role and concept behind ICAPCs -IHC Critical Assay Performance Controls Pros and cons..

### NordiQC

- International IHC proficiency testing program
- Institute of Pathology, Aalborg University Hospital, Denmark
- About 670 participants from >60 countries
- General module: 3 runs/year
  - 15-17 different diagnostic markers (CDs, CKs, Transcrip.,..)
- Breast cancer IHC module: 2 runs/year
  - 3-5 different markers (HER2, ER, PR,..)
- HER-2 ISH module: 2 runs/year
  - BRISH, FISH (breast cancer)
- Companion (CDx) module: 2 runs/year
  - PD-L1 TPS/CPS NSCLC/UC/TNBC (2017-)
  - PD-L1 IC score TNBC/UC (2019-)



www.nordigc.org



All news

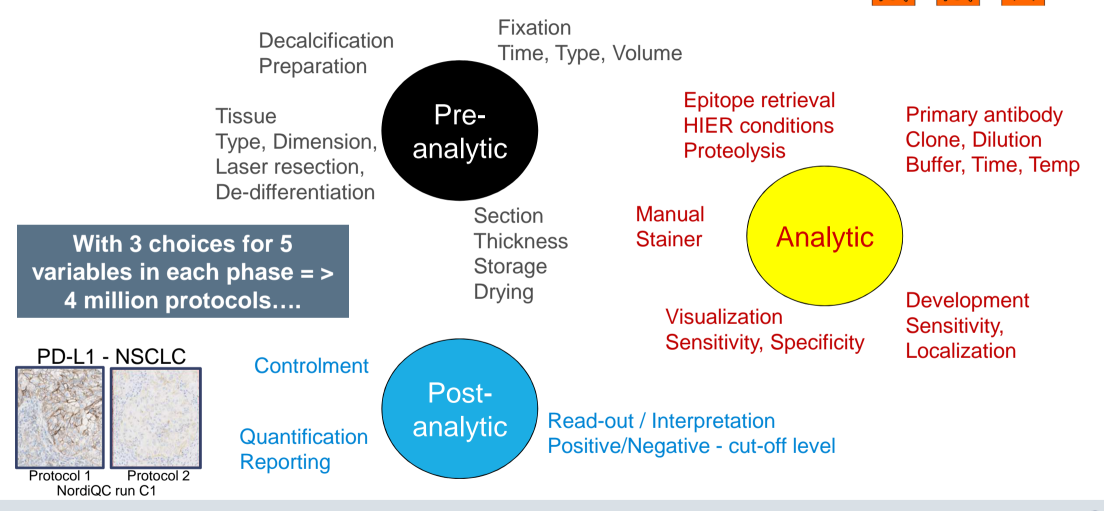




### NordiQC assessment scheme 2024

Module	Winter	Spring	Autum
General*	Run 70 PRAME CGA p53 Bcl-6 GATA3	<b>Run 71</b> <u>INSM1 CD20</u> <u>PMS2 BAP1 Ki67</u> <u>CD117</u>	Run 72 TRPS1 CD8 MSH6 p16 CK-PAN
Breast*	Run B37 HER2 IHC ER PR		Run B38 ER HER2 IHC
HER2 ISH	Run H25 HER2 ISH		Run H26 HER2 ISH
Companion*		Run C15 PD-L1 (TPS/CPS) PD-L1 (IC)	Run C16 PD-L1 (TPS/CPS) PD-L1 (IC)
*Accredited by DANA	Kunder registration nur	mber 616 to proficien	cy testing.

... The IHC biomarker protocol trap – Caution: not for faint-hearted lab personel !!!!!

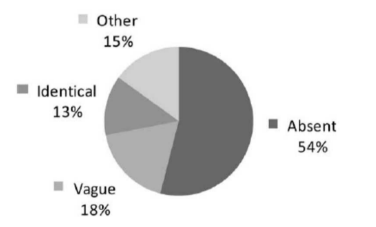


The use of the right controls at the right way can show if an IHC is right or wrong  $\odot$ 

## Importance of IHC controls have been neglected....

### Documentation of Diagnostic Cytopathology, Vol 39, No 4 2011 Immunocytochemistry Controls in the Cytopathologic Literature: ne A Meta-Analysis of 100 Journal Articles

Carol Colasacco, M.L.I.S., S.C.T.(A.S.C.P.), C.T.(I.A.C.),  $\overset{1*}{,}$  Sharon Mount, M.D.,  $^{1,2}$  and Gladv



### ICC Controls in the Literature

Fig. 1. Description of immunocytochemistry controls in articles reviewed.

Absent: Controls were not mentioned.

Vague: Statement such as "appropriate positive and negative controls were included."

Identical: Controls identical to study samples were described.

Other: Controls were dissimilar or partially similar (i.e., tissue control with smears or tissue control with cell block and ThinPrep samples run), or samples were too scant to include controls.

> 70 % of publications based on IHC do not describe controls used to verify data and conclusions....

6

# IHC controls to guide reliability of data...

PAX8 expression in breast cancer – true of false...?

Central for subtyping of unknown primary carcinoma; Ovary, uterine, kidney...

But....

Can PAX8 expression be seen in breast carcinoma??

Right choice, right use and results reported in positive and negative IHC control tissues needed to verify data

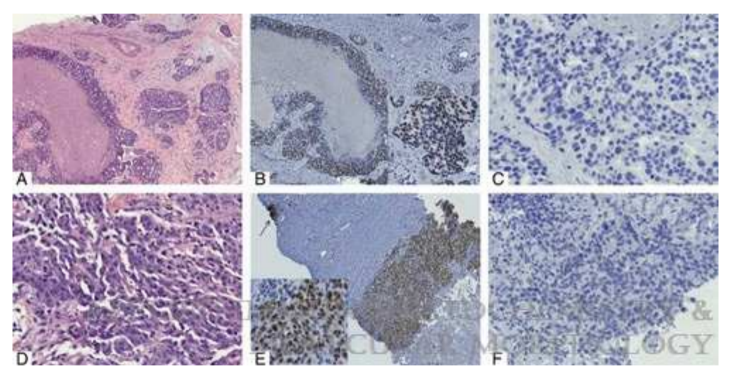
7

## IHC controls to guide reliability of data...

ΗE



BC12



#### Aberrant Immunostaining of Breast Carcinoma by MRQ-50 PAX8 Antibody

Singh, Kamaljeet; Hansen, Katrine; Quddus, M. Ruhul

Applied Immunohistochemistry & Molecular Morphology28(4):e37-e38, April 2020.

doi: 10.1097/PAI.00000000000082

Photomicrographs from 2 breast carcinomas with aberrant PAX8 expression by MRQ-50 clone. On staining with hematoxylin and eosin (A, D) both tumors were high grade with necrosis. Immunohistochemistry for PAX8 with MRQ-50 antibody (B, E) showed nuclear positivity in tumor cells and lymphocytes (arrow). PAX8 IHC with BC12 clone (C, F) did not stain tumor or lymphocytes.

### FIGURE 1

OPYRIGHT © 2021 WOLTERS KLUWER HEALTH, INC. AND/OR ITS SUBSIDIARIES. ALL RIGHTS RESERVED.

# IHC controls to guide reliability of data....

BC12 / SP348

NordiQC Assessments of PAX8 Immunoassays

Rasmus Roge, MD,\*† Ole Nielsen, HT,‡ Michael Bzorek, HT,§ Soren Nielsen, HT,\* and Mogens Vyberg, MD\*†

9

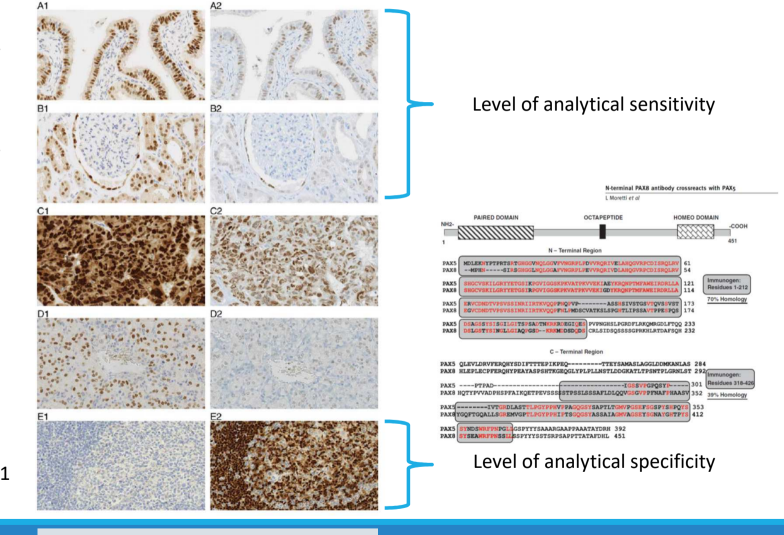
Positive tissue control 1 Fallopian tuba

Positive tissue control 2 Kidney

Tumour type 1 Ovarian carc.

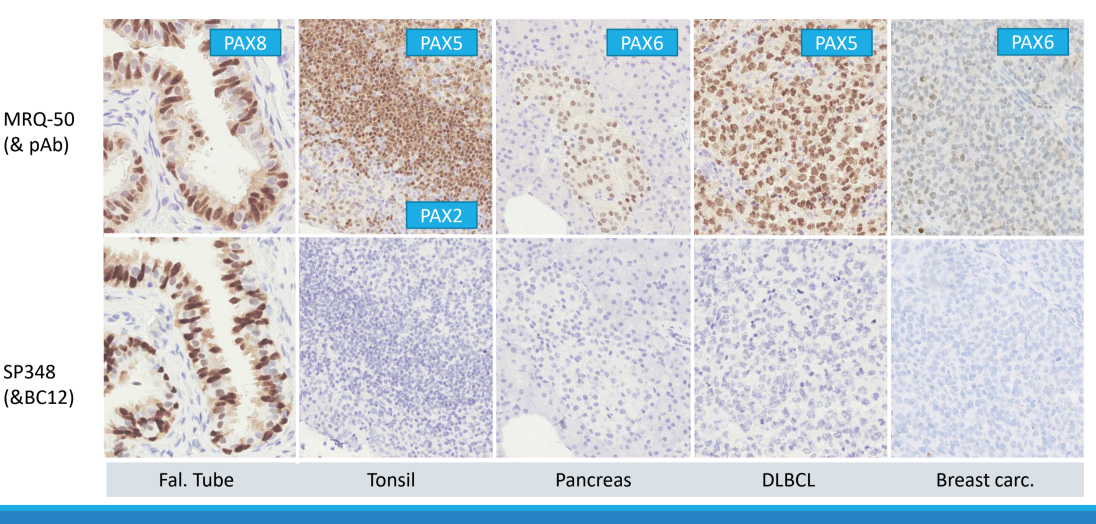
Tumour type 2 Renal cell carc.

Negative tissue control 1 Tonsil



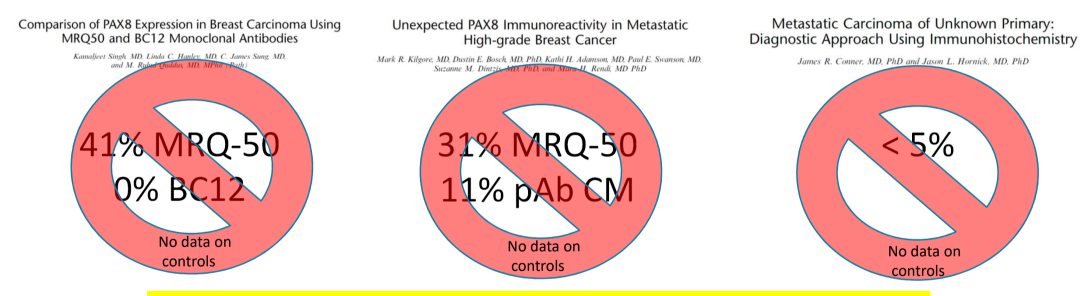
MRQ-50 / pAb

# IHC controls to guide reliability of data...



10

# IHC controls to guide reliability of data... PAX8 expression in breast cancer – true of false...?



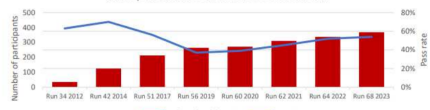
Right choice, right use and results reported in positive and negative IHC control tissues needed to verify data

11

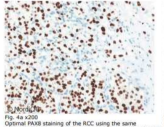
## NordiQC data – PAX8

#### Graph 1. Proportion of sufficient results for PAX8 in the eight NordiQC runs performed

PAX8 performance in NordiQC assessments







protocol as in Figs. 1a-3a. Virtually all the neoplastic cells show a moderate to strong nuclear staining reaction. No background staining is seen. Compare with Fig. 4b.

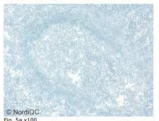


Fig. 5a x100 PAX8 staining without PAX5 cross reactivity. PAX8 staining in tonsil using the same protocol as in Figs. 1a-4a. The rmAb clone SP348 do not cross-react with PAX5, leaving the B-cells unstained. Compare with Fig. 5b.



@NordiQC Fig. 5b x100 PAX8 staining with PAX5 cross reactivity. PAX8 staining in tonsil using the same protocol as in Figs. 1b-4b. The mAb clone MRQ-50 cross-reacts with PAX5 resulting in nuclear staining reaction in virtually all B-cells. Compare with Fig. 5a.

Concentrated antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff,1	0
mAb clone BC12*	9 4	Biocare Zytomed Systems		3	7	3	23%	
mAb clone MRQ-50	16	Cell Marque		8	6	2	50%	
mAb clone PAX8R1	1	Abcam			1	0.00		
mAb clone ZM28	1	Zeta Corporation		1		1045	-	
rmAb clone EP298 <sup>5</sup> *	1	Epitomics <sup>5</sup>		1	1.5			
rmAb clone EP331*	10 4	Cell Marque Epitomics	÷	5	8	1	36%	
rmAb clone <b>SP348</b> *	146	Abcam Gennova Spring Bioscience	102	31	9	4	91%	7
rmAb clone <b>ZR-1</b> *	2	Zeta Corporation BioSite	1		2	1	<u>_</u>	
mAb clone BP6157*	2	Biolynx		1	1	(*)		
rmAb clone QR016*	7	Quartett	3	3	1	1.085	86%	4
pAb, 10336-1-AP	11	Proteintech	•	1	3	7	9%	
pAb, 363A-15	1	Cell Marque		1.1	1			
pAb, CP379 AK	3	Biocare	-		1	2	-	
pAb, <b>RBK047</b>	3	Zytomed Systems Diagomics	-	( <b>a</b> )	3	1.00	~	
Conc total	223		106	54	43	20	72%	4
Ready-To-Use antibodies							Suff. <sup>1</sup>	C
mAb clone MRQ-50, 760-4618 (VRPS) <sup>3</sup>	6	Ventana/Roche	2	-		6	0%	(
mAb clone MRQ-50, 760-4618 (LMPS) <sup>4</sup>	49	Ventana/Roche	-	3	34	12	6%	(
rmAb clone, EP331* 760-6077(VRPS) <sup>3</sup>	3	Ventana/Cell Marque	*	3 <b>1</b>	2	382		
rmAb clone, EP331* 760-6077(LMPS) <sup>4</sup>	11	Ventana/Cell Marque	÷.	4	6	1	36%	(
mAb clone, BC12* API438	6	Biocare Medical	-	2	4	-	33%	0
mAb clone IHC008 PII77R06	3	DCS		200	3	3.00		
rmAb clone ZR-1* Z2202	2	Zeta corporation	-	(a)	1	1		3
rmAb clone SP348* M6481	3	Spring Bioscience	2	1	•	۲		
mAb clone 2774R ANB31	1	Biogenex	•	.•:	1	. es.,	*	
mAb clone GR002* GT210202	1	GeneTech	1			383	*	
rmAb clone QR016* P-P008	2	Quartett	1	1	- 33	- 12 - J	- ¥	
mAb clone EP331* 363M/AC0338	12	Cell Marque		3	7	2	25%	(
rmAb clone SP348* 363R-38	4	Cell Marque	2	1	1	-		
mAb clone MRQ-50, 363M-10/17/18	24	Cell Marque	-	5	13	6	21%	(
pAb clone 363A-17/18 363A17/18	4	Cell Marque		•	3	1	•	
mAb clone MRQ-50, MAD-000550QD mAb clone RM436*	6	Master Diagnostica		4	1	1	67%	(
8257-C010	2	Sakura Finetek	1	<b>1</b>				
mAb clone IHC048*	1	GenomeMe			1			
mAb clone C12A32	1	Celnovte	-	1				
Clone MXR013* RMA-1024	2	Fuzhou Maixin	2				(*) 	
Clone H5A8 DTBL0220101	1	DaTe Bioengineering Technology	1	~		-	1	
Unknown	1					1	1.0	
					77	31	26%	8
RTU total Total	145 368		10 116	27 81	120	51	26%	25

1) Proportion of sufficient stains (optimal or good). (25 assessed protocols). 2) Proportion of Optimal Results (25 assessed protocols). 3) Vendor Recommended Protocol Settings (VRPS) to a specific RTU product applied on the vendor recommended platform(s) (25

assessed protocols). assessed protocols). 4) Laboratory Modified Protocol Settings (LMPS) to a specific RTU product (≥5 assessed protocols). 5) Ab terminated by vendor. \*Clones that do not show cross reactivity with PAX5.

# References central for the area of IHC controls

The "Kick-off" phase for

"Standardization of IHC controls"

Definitions and requirements Usage Potentials / Limitations Perspectives **REVIEW ARTICLE** 

Appl Immunohistochem Mol Morphol . Volume 22, Number 4, October 2014

### Standardization of Negative Controls in Diagnostic Immunohistochemistry: Recommendations From the International Ad Hoc Expert Panel

Emina E. Torlakovic, MD, PhD,\*†‡ Glenn Francis, MBBS, FRCPA, MBA, FFSc (RCPA),§||¶ John Garratt, RT,†‡# Blake Gilks, MD, FRCPC,†‡\*\* Elizabeth Hyjek, MD, PhD,\* Merdol Ibrahim, PhD,†† Rodney Miller, MD,‡‡ Soren Nielsen, HT, CT,§§|| || Eugen B. Petcu, MD, PhD,§ Paul E. Swanson, MD,¶¶ Clive R. Taylor, MD, PhD,## and Mogens Vyberg, MD§§|| ||

### REVIEW ARTICLE

Appl Immunohistochem Mol Morphol • Volume 23, Number 1, January 2015

### Standardization of Positive Controls in Diagnostic Immunohistochemistry: Recommendations From the International Ad Hoc Expert Committee

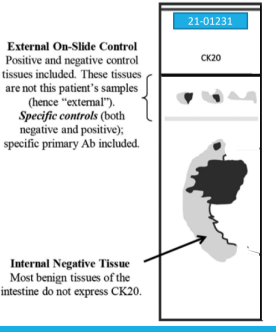
Emina E. Torlakovic, MD, PhD,\*† Søren Nielsen, HT, CT,‡§ Glenn Francis, MBBS, FRCPA, MBA, FFSc (RCPA), || ¶# John Garratt, RT,†\*\* Blake Gilks, MD, FRCPC,†††
Jeffrey D. Goldsmith, MD,‡‡ Jason L. Hornick, MD, PhD,\*§§ Elizabeth Hyjek, MD, PhD,\* Merdol Ibrahim, PhD, || || Keith Miller, FIBMS, || || Eugen Petcu, MD, PhD, ||
Paul E. Swanson, MD, ¶¶## Xiaoge Zhou, MD,\*\*\*††† Clive R. Taylor, MD, PhD,‡‡‡ and Mogens Vyberg, MD‡§

# Tissue controls

Reagent and tissue controls are necessary for the validation/ verification of immunohistochemical staining results.

Tissue controls are the most valueable tool to monitor the specificity and sensitivity for IHC

- Internal positive and negative tissue control
  - Cells/structures within the patient material
- External positive and negative tissue control
  - Slide next to patient material **on-slide optimally**



**Internal Negative Tissue** Most benign tissues of the intestine do not express CK20.

External On-Slide Control

Positive and negative control

are not this patient's samples

(hence "external"). Specific controls (both negative and positive); specific primary Ab included.

## How to use internal tissue controls

Appl Immunohistochem Mol Morphol • Volume 22, Number 4, April 2014

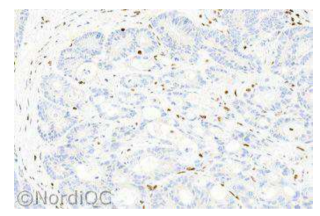
Standardization of Negative Controls

### TABLE 2. Examples of IHC Assays Where Preferential Use of Internal Positive Controls Recommended

IHC Assay	Use	Comments
Cytokeratin 5	Demonstration of basal cells in glandular structures of prostate to differentiate between benign (positive) and malignant (negative) glands	Interpretation of the results in the tumor directly depends on clear demonstration of internal positive control Tested sample may be completely negative if no normal tissue is
Mismatch repair proteins (MLH1, MSH2, PMS2, MSH6)	Absence of expression in the cells of colon or endometrial adenocarcinoma is abnormal; patients referred for molecular testing to rule out Lynch Syndrome	present Interpretation of the results in the tumor directly depends on clear demonstration of internal positive control

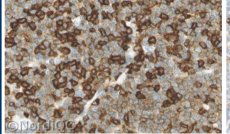
Target analyte	Application	Internal control to confirm "true" loss
BAP1, MTAP	Mesothelioma	Stromal cells
р53	Gynelogical carc.	Stromal cells
PTEN	Lung and gynecological carc.	Stromal and benign cells
MMR (MLH1, MSH2, MSH6, PMS2)	Lynch syndrome	Stromal cells / lymphocytes
SMAD4	Pancreas and GI carc.	Stromal and benign cells

Internal postive tissue controls; Principally ideal as processed identically to patient relevant material / target evaluated



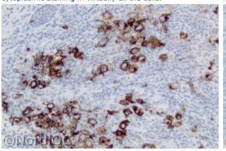


## Limitations of internal tissue controls



staining for CD5 of the B-CLL no. 5 using same protocol as in Figs. 1a - 4a. The majority of the neoplastic cells show a moderate and distinct staining reaction, while the infiltrating normal T-cells normal T-cells are clearly demonstrated. show a strong staining reaction

4b. Insufficient staining for CD5 of the B-CLL using same protocol as in Figs. 1b - 3b - same field as in Fig. The neoplastic cells are virtually negative and only the



2a. Optimal CD15 staining of the Hodgkin lymphoma 2 (NS) using same protocol as in Fig. 1a. The Reed-Sternberg and Hodgkin cells show a strong membranous staining and a dot-like positivity



no Fig. 2b. CD15 staining of the Hodgkin lymphoma no 2 (NS) using same protocol as in Fig. 1b. Only few Reed-Sternberg and Hodgkin cells show a weak staining - same field as in Fig. 2a

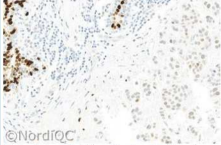


Fig. 3a, Optimal ER staining of the breast ductal carcinoma no. 3 with 60 - 80 % cells positive. A weak but distinct nuclear staining is seen in the appropriate proportion of the protocol as in Figs. 1b and 2b - same field as in Fig. 3a. neoplastic cells. Same protocol as in Figs. 1a and 2a.

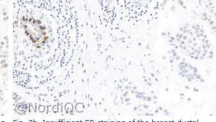


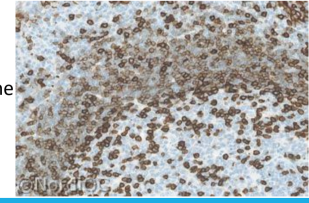
Fig. 3b. Insufficient ER staining of the breast ducta carcinoma no. 3 with 60 - 80 % cells positive using same Only dispersed neoplastic cells show an equivocal staining

Internal positive tissue controls;

In general not applicable as positive controls due to levels of expression may not be relevant for level of test calibration

e.g. CD5, CD15, CD34, CD45, CD56, S100, ER, PD-L1 etc

CD5; Tonsil Mantle zone Critical control

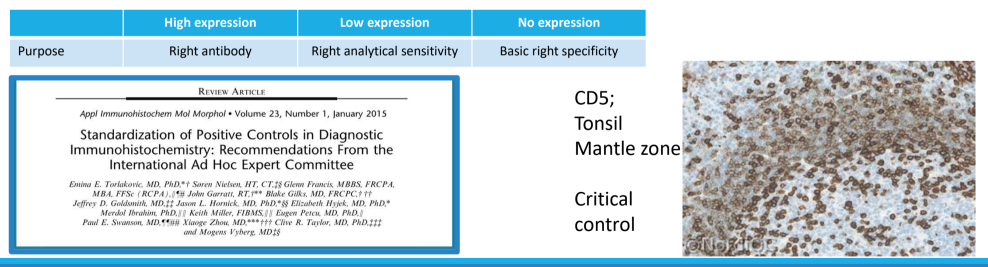


Critical tissue controls = ICAPCs

IHC Critical Assay Performance Controls (ICAPCs)

are basically human positive control tissues with

- clinical relevant range of target analyte (antigen) especially with low limit detection
- well characterized expression pattern preferable normal tissues
- predictable levels and specified cellular and architectural localization



## Test Performance Characteristics - TPCs

Test performance characteristics;

Which staining pattern characterizes an optimally calibrated IHC assay for a specific purpose?

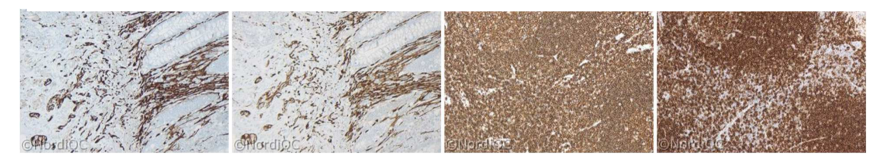
Analytical sensitivity Analytical specificity Precision / reproducibility of IHC assay

Which tissues / cellular structures show the clinical relevant range of the target analyte with focus on required low level of demonstration – <u>CRITICAL CONTROLS - ICAPCs</u>?

CD56

Colon

CD45



Tonsil

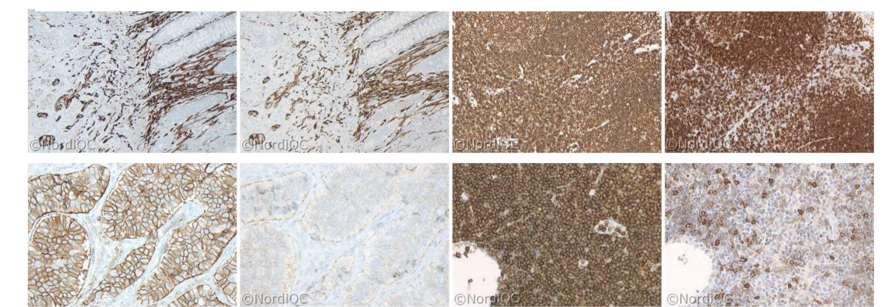
Test A	Test B	Test A	Test B

CD56

CD45

Tonsil

**B-CLL** 





NET

Colon

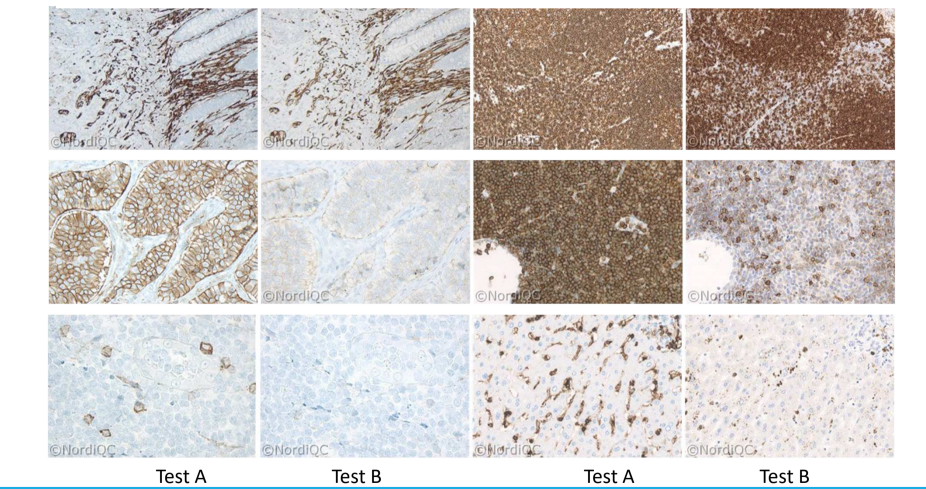
CD56

Colon

NET

**Tonsil** 

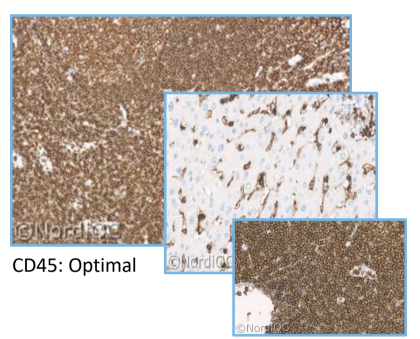
CD45

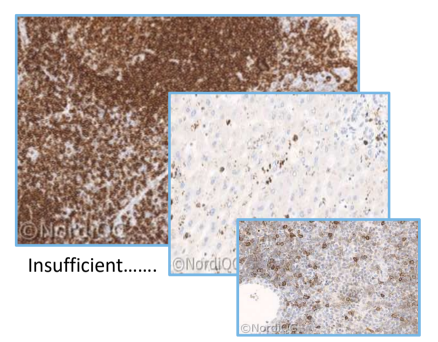


Tonsil

**B-CLL** 

<u>Liver</u>





Tissues/cells with only high expression will not identify:

- 1. A poorly calibrated IHC assay
- 2. A reduced sensitivity in an optimally calibrated IHC assay

If an IHC test is used to identify the target antigen being expressed at different levels, controls must reflect this!

## iCAPCs - concept

IHC Critical Assay Performance Controls (iCAPCs)

Which tissues are recommended ?

What is the expected staining pattern ?

Which tissues / cells are critical ?

Right antibody Appropriate level of sensitivity Guidance level of specificity **REVIEW ARTICLE** 

Appl Immunohistochem Mol Morphol • Volume 23, Number 1, January 2015

### Standardization of Positive Controls in Diagnostic Immunohistochemistry: Recommendations From the International Ad Hoc Expert Committee

Emina E. Torlakovic, MD, PhD,\*† Søren Nielsen, HT, CT,‡§ Glenn Francis, MBBS, FRCPA, MBA, FFSc (RCPA), [[¶]# John Garratt, RT,†\*\* Blake Gilks, MD, FRCPC,††† Jeffrey D. Goldsmith, MD,‡‡ Jason L. Hornick, MD, PhD,\*§§ Elizabeth Hyjek, MD, PhD,\* Merdol Ibrahim, PhD, []] Keith Miller, FIBMS, [] [] Eugen Petcu, MD, PhD, [] Paul E. Swanson, MD, []¶]## Xiaoge Zhou, MD,\*\*\*††† Clive R. Taylor, MD, PhD,‡‡‡ and Mogens Vyberg, MD‡§

### Estrogen and Progesterone Receptor Testing in Breast Cancer: ASCO/CAP Guideline Update

Kimberly H. Allison, MD<sup>1</sup>; M. Elizabeth H. Hammond, MD<sup>2</sup>; Mitchell Dowsett, PhD<sup>3</sup>; Shannon E. McKemin<sup>4</sup>; Lisa A. Carey, MD<sup>5</sup>; Patrick L. Fitzgibbons, MD<sup>6</sup>; Daniel F. Hayes, MD<sup>7</sup>; Suni R. Lakhani, MD<sup>6,9</sup>; Mariana Chavez-MacGregor, MSc<sup>10</sup>; Jane Perlmutter, PhD<sup>11</sup>; Charles M. Perou, PhD<sup>6</sup>; Meredith M. Regan, ScD<sup>12</sup>; David L. Rimm, MD, PhD<sup>13</sup>; W. Fraser Symmans, MD<sup>16</sup>; Emina E. Torlakovic, MD, PhD<sup>14,15</sup>; Leticia Varella, MD<sup>16</sup>; Giuseppe Viale, MD<sup>17,18</sup>; Tracey F. Weisberg, MD<sup>19</sup>;

Lisa M. McShane, PhD<sup>20</sup>; and Antonio C. Wolff, MD<sup>21</sup>

J Clin Oncol 38:1346-1366. © 2020 by American Society of Clinical Oncology

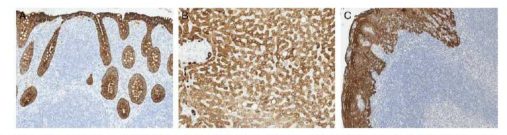


FIGURE 1. Pan-keratin iCAPC. A, Appendix: virtually all columnar epithelial cells must show a moderate to strong predominantly cytoplasmic staining reaction (a membranous accentuation will typically be seen). B, Liver: the vast majority of hepatocytes must show at least weak to moderate cytoplasmic staining reaction with a membranous accentuation (LLOD). C, Tonsil: all squamous epithelial cells must show a moderate to strong cytoplasmic staining reaction. Cytokeratin (CK)-positive interstilal reticulum cells (CIRCs) with dendritic/reticular pattern can show a weak to moderate cytoplasmic staining reaction (LLOD). iCAPC indicates immunohistochemistry critical assay performance controls: LLOD. low limit of detection.



FIGURE 7. TTF-1 iCAPC. A, Thyroid: virtually all epithelial cells must show a strong nuclear staining reaction. B, Lung: virtually all pneumocytes and basal cells of terminal bronchi must show a moderate to strong nuclear staining reaction. Columnar epithelial cells of terminal bronchi must show an at least weak nuclear staining reaction (LLOD). C, Tonsil: no staining reaction must be seen. iCAPC indicates immunohistochemistry critical assay performance controls; LLOD, low limit of detection.



FIGURE 8. CDX-2 iCAPC. A, Appendix: virtually all epithelial cells must show a strong nuclear staining reaction. A weak cytoplasmic staining reaction in addition to strong nuclear staining is often present. B, Pancreas: the majority of epithelial cells of intercalated ducts must show a weak to moderate nuclear staining reaction (LLOD). C, Tonsil: no staining reaction must be seen. iCAPC indicates immunohistochemistry critical assay performance controls; LLOD, low limit of detection.

Examples for 17 markers Generel expected patterns

High expression (Right antibody)

Low expression (Appropriate sensitivity)

No expression (Appropriate specificity)

Which tissue Which cells Which extension Which intensity

### NordiQC IHC tissue control atlas – open from 05.2022

NordiQC

Info • Modules • Assessments Protocols Controls Events • <u>SN</u>

### **Recommended controls**

		Search:
Epitope 🔺	Tissues	Actions
ALK (lung)	Appendix/colon, Tonsil	See controls
AMACR	Kidney, Prostate	See controls
ASMA	Appendix/colon, Liver	See controls
BcI-2	Tonsil	See controls
Bcl-6	Tonsil	See controls
BSAP	Hodgkin lymphoma, Tonsil	See controls
C-MYC	Appendix/colon, Tonsil	See controls
CD3	Appendix/colon, Tonsil	See controls
CD4	Liver, Tonsil	See controls
CD5	Tonsil	See controls
CD8	Appendix/colon, Tonsil	See controls
CD10	Kidney, Tonsil	See controls
CD15	Kidney, Tonsil	See controls
CD19	Appendix/colon, Tonsil	See controls
CD20	Appendix/colon, Tonsil	See controls
CD23	Tonsil	See controls
CD30	Tonsil	See controls
CD31	Appendix/colon, Liver, Tonsil	See controls

Available for NordiQC participants
Tissues
Purpose
Reaction patterns
Online scans accessible

### NordiQC IHC tissue control atlas – open from 05.2022



Info • Modules • Assessments Protocols Controls Events • <u>SN</u>

Available for NordiQC participants

Tissues

Purpose

Reaction patterns

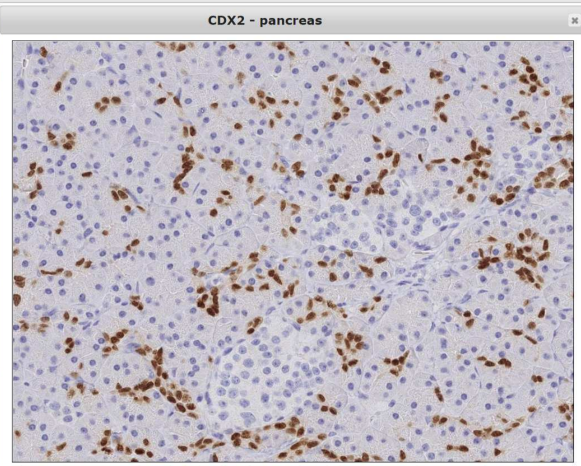
Online scans accessible

Control type	Positive tissue control High expression level	Positive tissue control Low expression levels	Negative tissue control
Tissue	Appendix/colon	Pancreas	Tonsil
Description	All epithelial cells must show a strong nuclear staining reaction. Note, a weak cytoplasmic staining reaction in CDX2 positive cells can be seen and should be accepted if signal-to- noise ratio otherwise is acceptable.	The vast majority of epithelial cells of intercalated ducts must show a weak to moderate nuclear staining reaction.	No staining reaction should be seen. Note, dispersed lymphocytes can show a faint nuclear staining reaction.
Example	Click to enlarge	Click to enlarge	•     •       •

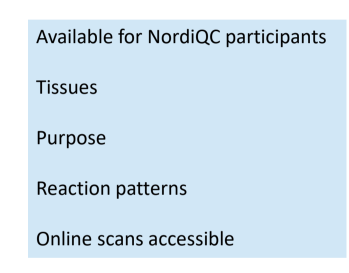
### CDX2 - CDX2

Back

### NordiQC IHC tissue control atlas – open from 05.2022



The vast majority of epithelial cells of intercalated ducts must show a weak to moderate nuclear staining reaction.



# Challenges for ICAPCs / Critical Controls

- Limited access to relevant tissues rare incidences
   ALK (lung), ROS1, Myogenin..
- New markers not described in details no data on test performance characteristics
   SATB2, Claudin-4, PRAME, TRPS1....
- Limited access to reference material and/or critical expression levels
   PD-L1, HER2, ER...

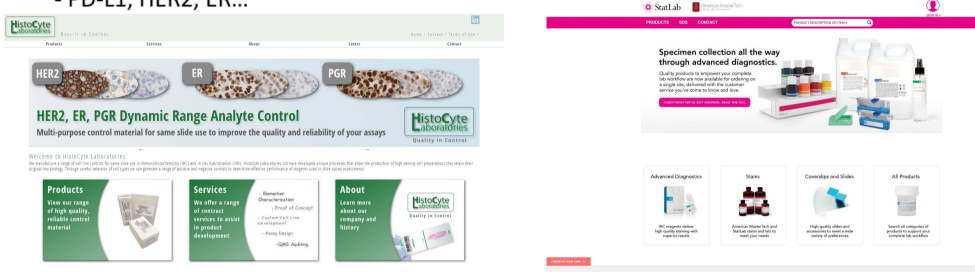


# Role of cell lines & histoids for IHC test development

- 1. Limited access to relevant tissues rare incidences
  - ALK (lung), ROS1, Myogenin..
- Limited access to reference material and/or critical expression levels

   PD-L1, HER2, ER...

Starting help to guide development – validation still required....



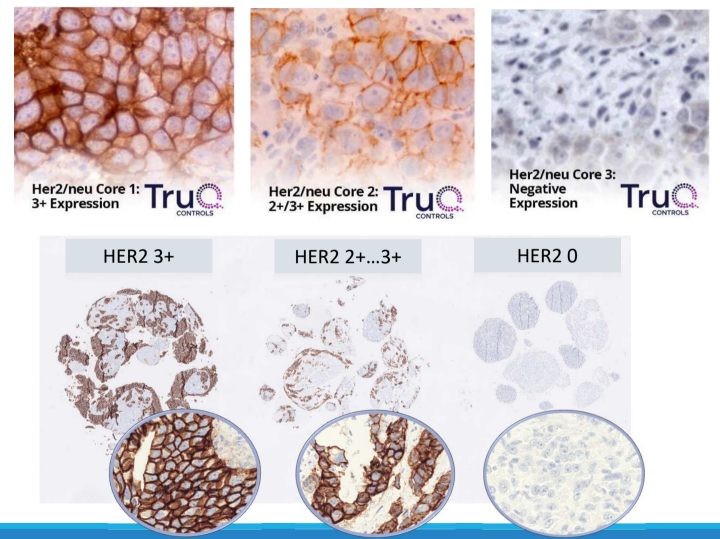
### www.histocyte.com

Cell lines ALK and ROS1 being +/-HER2, ER, PR and PD-L1 with dynamic range

### www.statlab.com

Histoids / Faux tissue ALK +/-HER2, PD-L1 with dynamic range

## Histoids / Faux tissue – TruQ IHC controls



Tissue core with IHC 3+ and IHC 2+ almost identical concerning expression levels.

No IHC 1+ tissue

Design seems less adequate for "precision testing" for HER2 IHC both "classical" and HER2 low.

www.statlab.com

# Role of cell lines for IHC test development

#### HER2 Analyte Control<sup>DR</sup>

Cell line controls for immunohistochemistry and in situ hybridization.

#### **Research Use Only**

#### PRODUCT AVAILABILITY

Product Code	Product Description
HCL026	X2 Cut slides
HCL027	X5 Cut slides
HCL028	X1 Cell microarray block

#### APPLICATION

This product is suitable for use in immunohistochemistry and in situ hybridization.

#### MATERIALS

Four formalin fixed paraffin embedded cell lines with a dynamic range (DR) of expression for Human Epidermal growth factor Receptor 2 (HER2).

Cell line A: Breast adenocarcinoma Cell line B: Breast adenocarcinoma Cell line C: Gastric adenocarcinoma Cell line D: Breast adenocarcinoma

Cells are fixed in 10% neutral buffered formalin and paraffin wax embedded. Sections are cut at 4 $\mu$ m, mounted on positively charged slides

and baked overnight at 37°C.



Cell microarrays (CMA) contain cores that are 1.5-2mm in diameter and 3-3.5mm in length. It is possible to obtain over 300 sections depending on thickness.



### Expression Profile

Cell Line	IHC for HER2	amplification
A	0	Non-amplified
В	1+	Non-amplified
с	2+	Equivocal
D	3+	Amplified

#### Storage and Handling

Store at 2-8°C. Do not freeze (for expiration date please see the product label)

#### WARNINGS AND PRECAUTIONS

- The product is intended for research use only. It is the responsibility of the end user to determine suitability with their reagents and procedures within their laboratory.
- Do not use after expiration date printed on product labels. The user must validate any storage conditions other than those specified in the package insert.

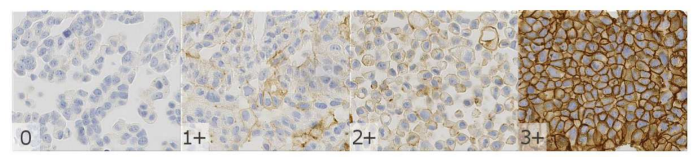
#### TROUBLE SHOOTING

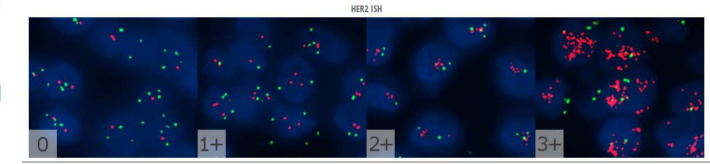
2

For further help please feel free to contact HistoCyte Laboratories Ltd at info@histocyte.com or call +44 (0)191 603 1007.

For updates and additional product information please visit: www.HistoCyte.com

In NordiQC run B34 10% of the participants used cell lines as onslide control





Still need evidence/proof (VALIDATION) how to correlate any change in staining pattern in cell lines for accuracy in tissues of breast carcinoma.

Tissue and cell line expression robustness ( too fragile or too stabile)? What expression levels characterizes a successful vs insuccessful test? Impact on section thickness? Pattern on different assays?

31

### Correlation of IHC for HER2 – accurate PATHWAY – cell lines and tissues

2+ 1+ 2+ AMP 3+ 1+ 0

Cell lines (HCL028 Histocyte)

Breast carcinomas

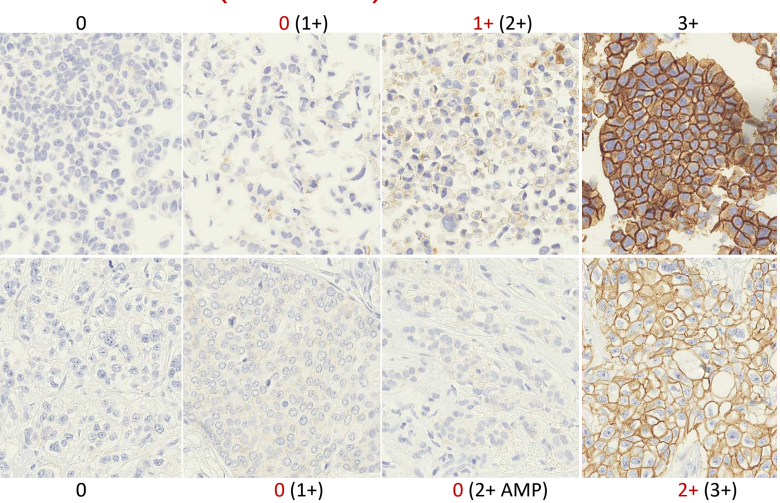
NordiQC run B34

### Correlation of IHC for HER2 – (inaccurate) PATHWAY – cell lines and tissues

Cell lines (HCL028 Histocyte)

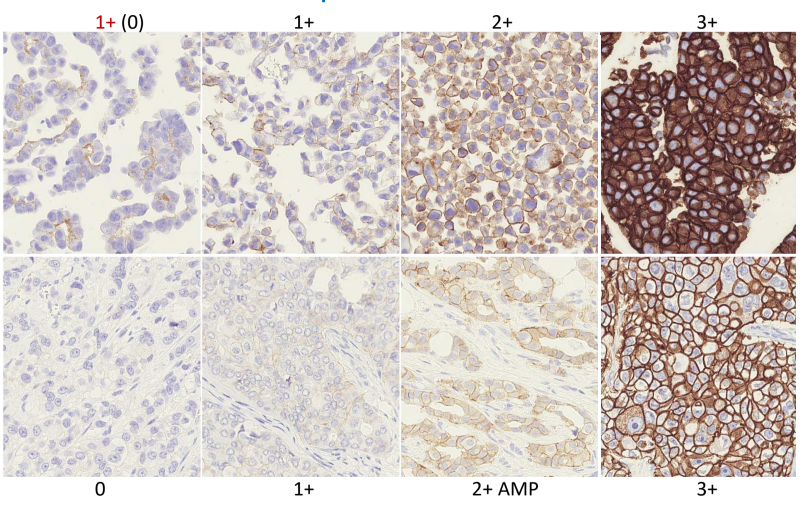


NordiQC run B34



### Correlation of IHC for HER2 – HercepTest 2' Gen – cell lines and tissues

Cell lines (HCL028 Histocyte)



Breast carcinomas

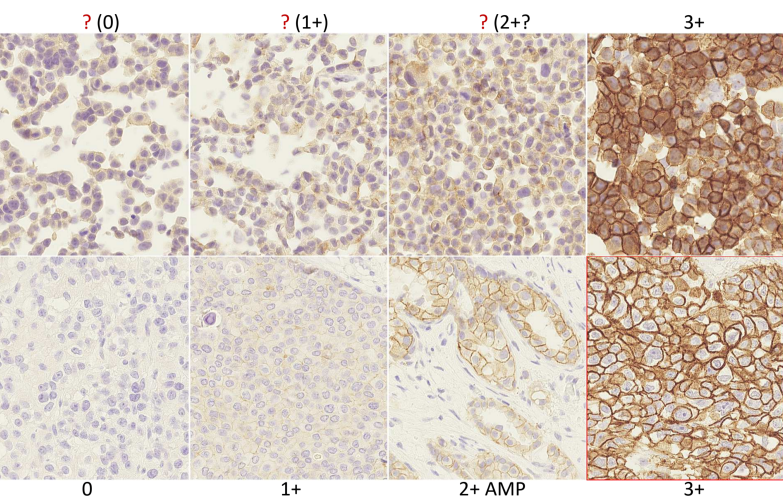
NordiQC run B34

### Correlation of IHC for HER2 – SP3 – cell lines and tissues

Cell lines (HCL028 Histocyte)

Breast carcinomas

NordiQC run B34



9/27/2023

### Correlation of IHC for HER2 – cell lines and scoring

2+ 2+ 2+ weak 1+ 1+ 0

Cell lines (HCL028 Histocyte)

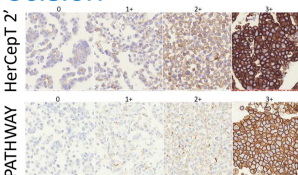
Breast carcinoma 1+

## The needs for cell lines as Quality tool for Accuracy/Precision

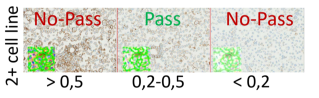
- Need to map staining characteristics for most commonly used IHC assays
  - The different assays will provide different patterns
- Need to identify change in patterns being critical with risk of false negative / false positive results
  - Each assay most likely will have different patterns / tresholds
- Need to integrate software as digital image analysis (DIA) or artificial intelligence (AI) to secure reproducibility
  - Identification of DIA/AI QC-score for successful versus insuccessful test

- The DIA/AI QC-scores must be validated for each IHC assay both with focus on expected level and critical levels
  - Large scale testing on e.g. breast carcinomas with the dynamic and critical range of the target analyte
  - Both to identify e.g. "classical" HER2 overexpression and the novel HER2 low category

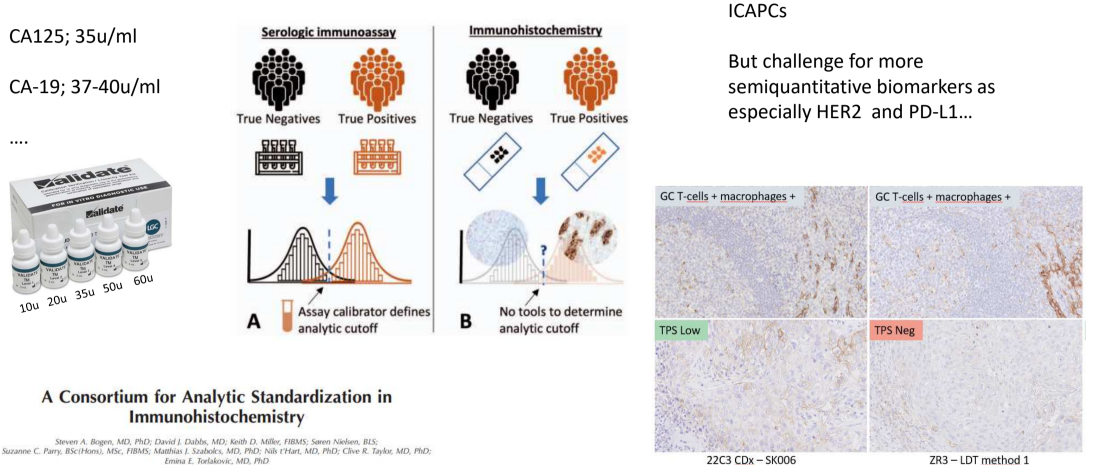








## Analytical standards – IHC versus clinical chemistry; Calibrators



Emina E Torlakovic MD PhD

(Arch Pathol Lab Med. doi: 10.5858/arpa.2022-0031-RA)

....

## Analytical standards – IHC versus clinical chemistry; Calibrators

Developmental and validatation phase to correlate LOD\*/analytical sensitivity in microbeads versus diagnostic accuracy and sensitivity for;

ER, HER2, PD-L1 and p53

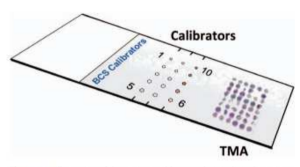


Figure 5. Illustration of the survey tool for correlating clinical accuracy (from the tissue microarray data) with analytic sensitivity (from the calibrator data). The calibrators are at up to 10 different concentrations, for example levels 1–10. The middle row depicts negative controls. Abbreviations: BCS, Boston Cell Standards; TMA, tissue microarray.

#### A Consortium for Analytic Standardization in Immunohistochemistry

Steven A. Bogen, MD, PhD; David J. Dabbs, MD; Keith D. Miller, FIBMS; Søren Nielsen, BLS; Suzanne C. Parry, BSc(Hons), MSc, FIBMS; Matthias J. Szabolcs, MD, PhD; Nils t'Hart, MD, PhD; Clive R. Taylor, MD, Ph Emina E. Torlakovic. MD, PhD

(Arch Pathol Lab Med. doi: 10.5858/arpa.2022-0031-RA)

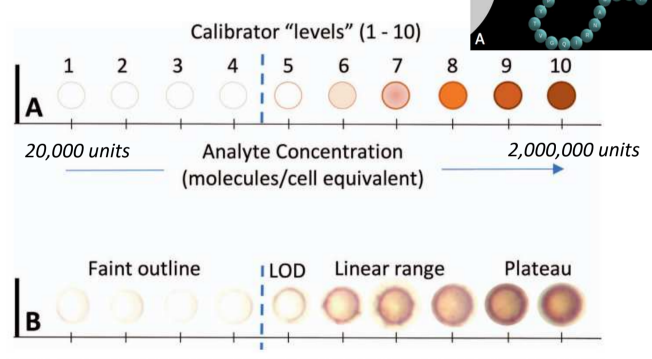


Figure 2. Illustration of a series of immunohistochemistry calibrators after staining. The numbers refer to calibrator levels, from low (1) to high (10) analyte concentrations. A, The illustration shows that rim staining is stronger than central staining because the analyte is attached to the microbead surface. In this example, level 5 represents the lower limit of detection (LOD). B, Images of microbeads from calibrators with an LOD at level 5.

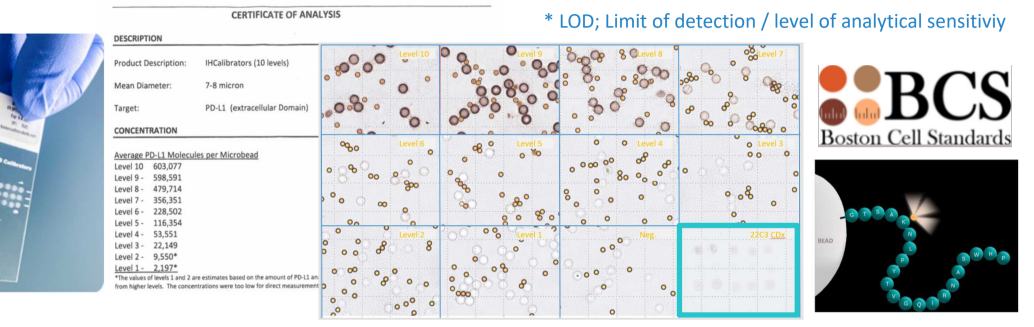
**Boston Cell Standards** 

Micro

bead

## Reference standard materials for IHC; Calibrators – LOD\* - PD-L1





Bogen, SA. 2019. A root cause analysis into the high error rate in clinical immunohistochemistry. Appl. Immunohistochem. Mol. Morphol. 27(5) 329-338.

Sompuram, SR, K Vani, AK Schaedle, A Balasubramanian, & SA Bogen. 2019. Selecting an optimal positive IHC control for verifying retrieval. J. Histochem. Cytochem. 67(4):273-283.

Sompuram, SR, K Vani, AK Schaedle, A Balasubramanian, & SA Bogen. 2018. Quantitative assessment of immunohistochemistry laboratory performance by measuring analytic response curves and limits of detection. Arch Pathol Lab Med. 142 (7):851-862.

### Reference standard materials for IHC; Calibrators – LOD – PD-L1 22C3

22C3 LOD

356.351 mol. pr microbead

Average PD-L1 Molecules per Microbead

603.077

598.591

479 714

356 351

228.502

116.354

53.551

22,149

Level 10

Level 9

Level 8

Level 7

Level 6

Level 5

Level 4 -

Level 3 -

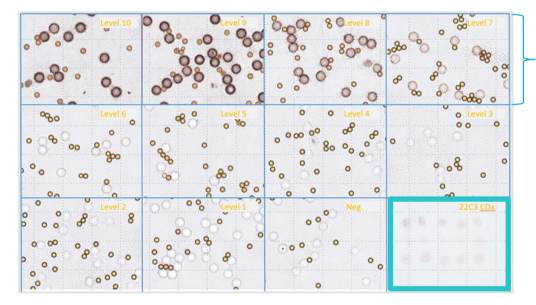
Level 2 - 9,550\*

Level 1 - 2,197\*

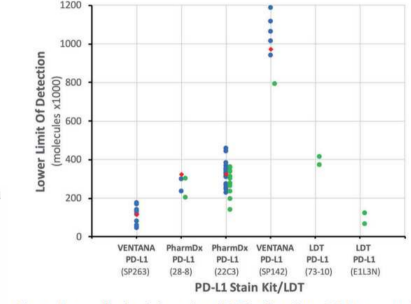
#### ARTICLE OPEN Quantitative comparison of PD-L1 IHC assays against NIST standard reference material 1934

Seshi R. Sompuram<sup>1</sup>, Emina E. Torlakovic<sup>2,3</sup>, Nils A. 't Hart<sup>4</sup>, Kodela Vani<sup>1</sup> and Steven A. Bogen<sup>1⊠</sup>

© The Author(s), under exclusive licence to United States & Canadian Academy of Pathology 2021, corrected publication 2021



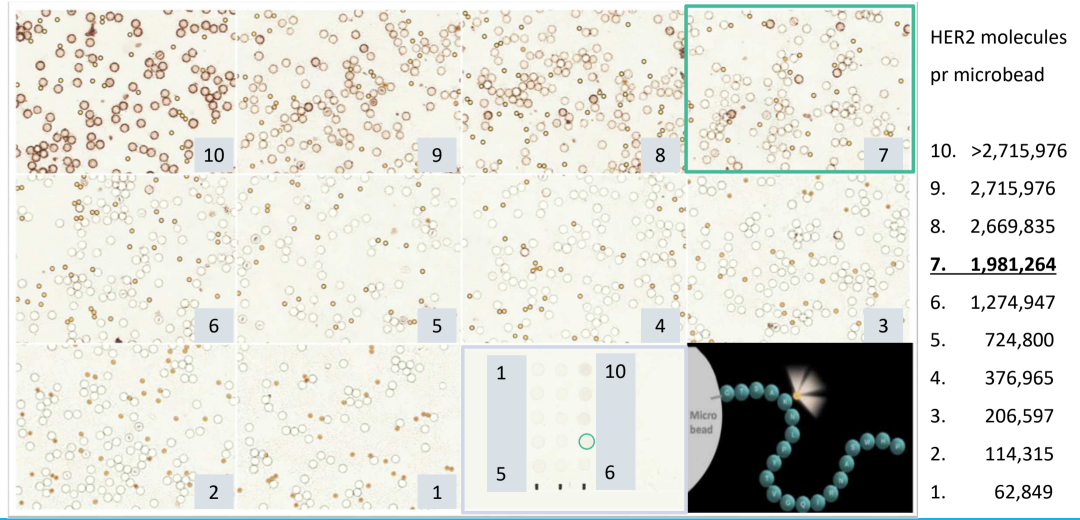




**Fig. 2** Lower limit of detection (LOD) of various PD-L1 assays (x axis). Lower numbers (on the y axis) equate to greater sensitivity. Each dot represents a separate IHC laboratory test. Blue dots depict FDA-cleared assays in clinical laboratories, green dots for laboratory-developed tests (LDTs), and red diamonds for FDA-cleared assays as performed by a reference laboratory. Tissue staining in Fig. 2 was performed by these reference labs. For enhanced clarity, the LDT data are positioned slightly to the right of the vertical lines.

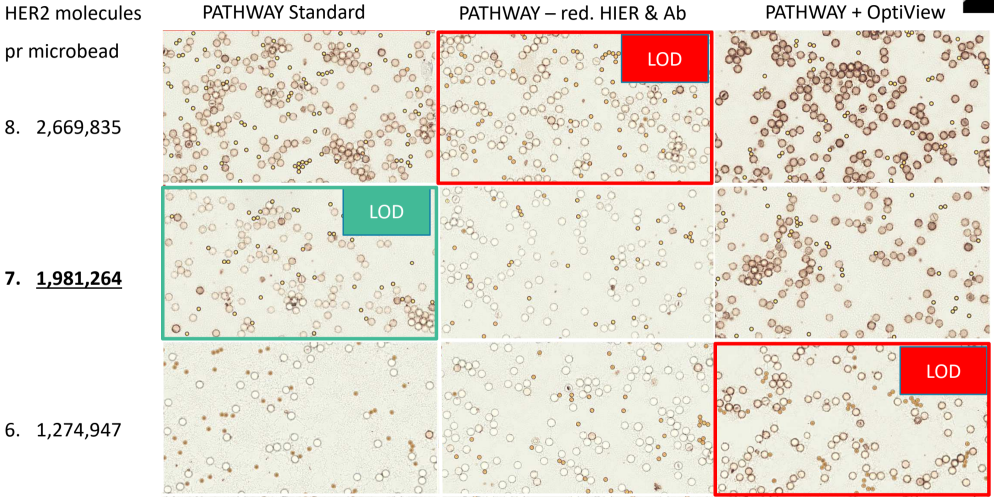


### IHC Calibrator 10 levels HER2 – Boston Cell Standards - PATHWAY

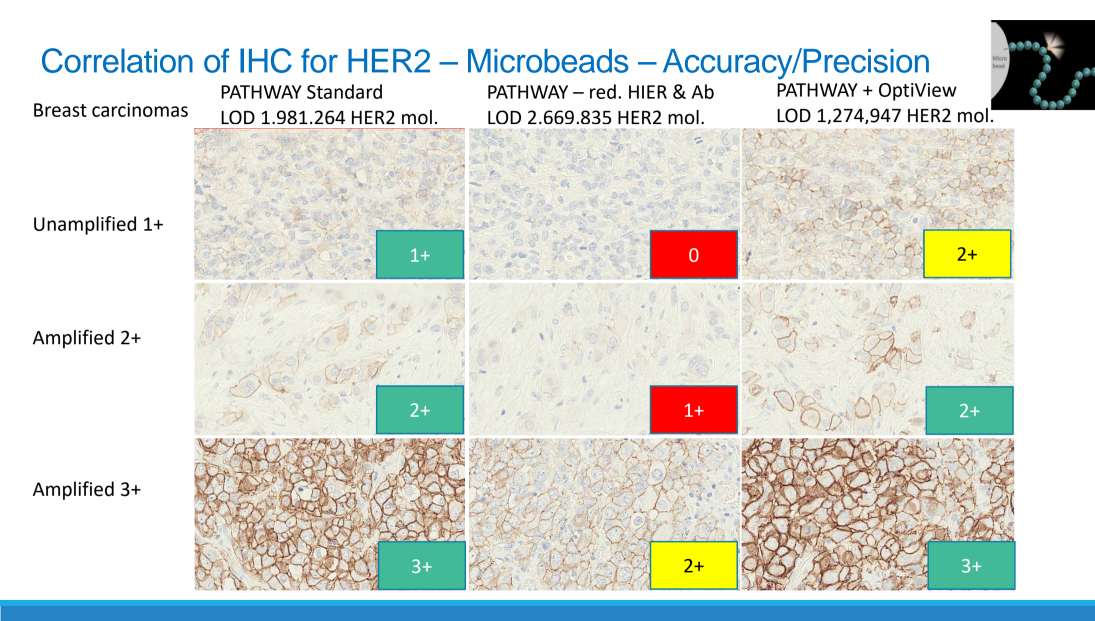


## Correlation of IHC for HER2 – Microbeads – Accuracy/Precision





9/27/2023



## Correlation of IHC for HER2 – Microbeads – Accuracy/Precision



Breast carcinomas

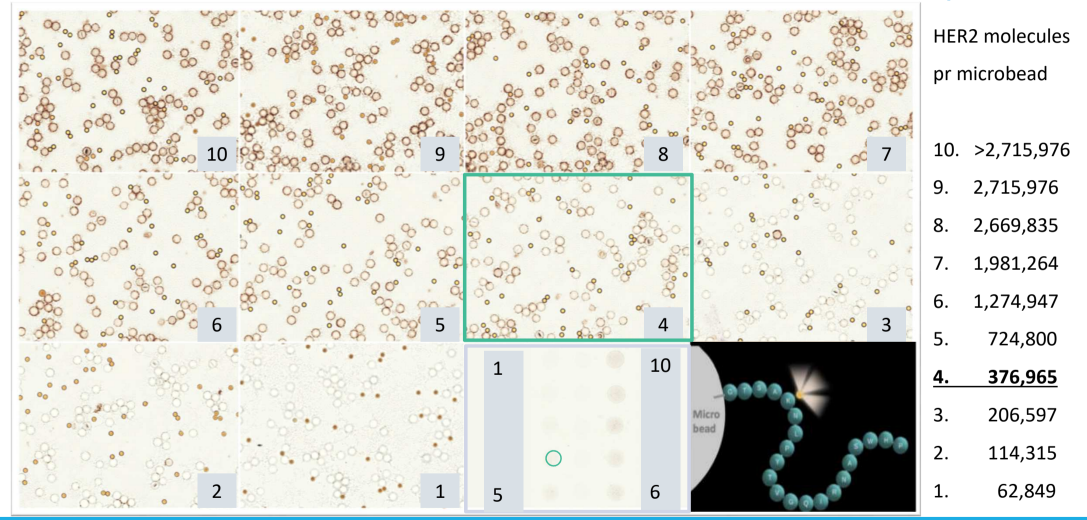
N=15 (NordiQC runs B31, B32, B33)

			PATHWAY Standard LOD 1,981,264 HER2 mol.	PATHWAY – red. HIER & Ab LOD 2,669,835 HER2 mol.	PATHWAY + OptiView LOD 1,274,947 HER2 mol.
HER2 classical	HER2 Low	0	2	5	0
		1+	3	3	3
		2+ Unamplified	1	2	3
		2+ Amplified	3	1	3
		3+ Amplified	6	4	6

Reduced analytical sensitivity (LOD) provided a less accurate HER2 result for both classical overexpression and HER2 low

Increased analytical sensitivity (LOD) provided a less accurate HER2 result for HER2 low

### IHC Calibrator 10 levels HER2 – Boston Cell Standards – HercepTest Mo.



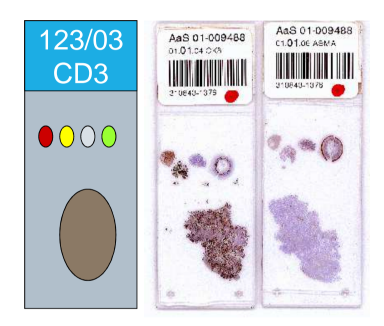
## Standardized controls for Immunohistochemistry

- Precision testing for precision medicine needs precision IHC controls
- At present no "golden standard IHC controls" to fit all IHC biomarkers
- A mixture of carefully selected external tissue controls and non-tissue based controls as cell lines and/or microbeads seem to be best practice
- Cell lines and microbeads have potential to monitor IHC test precision and accuracy, <u>BUT</u> still require extensive documentation and data how to use these

Different performances related to IHC assays Different tresholds for adequate vs inadequate result Software DIA/AI QC-tools to be developed and verified

Daily IHC control for the majority of routine markers:

Appendix Liver Pancreas Tonsil



Each slide stained and evaluated has essential information of the obtained sensitivity and specificity In contrast only using 1 external tissue run control, no information is available for the single slide evaluated

	TMA On-slide control	TMA Run / batch control	Remarks
Missing reagent FN in patient test	Yes	No – only control slide	Potential internal pos. control only indicator of protocol performed
Wrong antibody FP in patient test	Yes	No – only control slide	
Inappropriate protocol performance - Drying out etc FN / FP in patient test	Yes	No – only control slide	Potential internal pos. control only indicator of protocol performed

Errors seen for all IHC automated and semi-automated IHC platforms

49

## On-slide controls....

**REVIEW ARTICLE** 

(Appl Immunohistochem Mol Morphol 2015;23:1-18)

#### Standardization of Positive Controls in Diagnostic Immunohistochemistry: Recommendations From the International Ad Hoc Expert Committee

Emina E. Torlakovic, MD, PhD,\*+ Soren Nielsen, HT, CT. \$\$ Glenn Francis, MBBS, FRCPA, MBA, FFSc (RCPA), ## John Garratt, RT, +\*\* Blake Gilks, MD, FRCPC, ++ Jeffrey D. Goldsmith, MD, 22 Jason L. Hornick, MD, PhD, \* SS Elizabeth Hyjek, MD, PhD,\* Merdol Ibrahim, PhD, III Keith Miller, FIBMS, III Eugen Petcu, MD, PhD, II Paul E. Swanson, MD, ¶## Xiaoge Zhou, MD,\*\*\*††† Clive R. Taylor, MD, PhD, \*\*\* and Mogens Vyberg, MD18

#### **RESEARCH ARTICLE**

(Appl Immunohistochem Mol Morphol 2017;25:308-312)

An Audit of Failed Immunohistochemical Slides in a Clinical Laboratory: The Role of On-Slide Controls

Carol C. Cheung, MD, PhD, JD,\*† Clive R. Taylor, MD, DPhil,2 and Emina E. Torlakovic, MD, PhD†

#### ASCO Estrogen and Progesterone Receptor Testing in Breast Cancer: ASCO/CAP Guideline Update

Kimberly H. Allison, MD<sup>1</sup>; M. Elizabeth H. Hammond, MD<sup>2</sup>; Mitchell Dowsett, PhD<sup>3</sup>; Shannon E. McKernin<sup>4</sup>; Lisa A. Carey, MD<sup>5</sup>: Patrick L, Fitzgibbons, MD<sup>6</sup>: Daniel F, Haves, MD<sup>7</sup>: Sunil R, Lakhani, MD<sup>8,9</sup>: Mariana Chavez-MacGregor, MSc<sup>10</sup>: Jane Perlmutter, PhD<sup>11</sup>: Charles M. Perou, PhD3: Meredith M. Regan, ScD12: David L. Rimm, MD, PhD13: W. Fraser Symmans, MD10: Emina E. Torlakovic, MD, PhD<sup>14,15</sup>; Leticia Varella, MD<sup>36</sup>; Giuseppe Viale, MD<sup>17,18</sup>; Tracey F. Weisberg, MD<sup>39</sup>; Lisa M. McShane, PhD<sup>20</sup>; and Antonio C. Wolff, MD<sup>21</sup>

ticle J Clin Oncol 38:1346-1366. © 2020 by American Society of Clinical Oncology

qs

0

Cla

ar

"even for automated stainers, where it cannot be guaranteed that every slide in fact receives identical treatment".



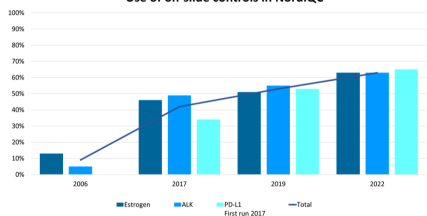
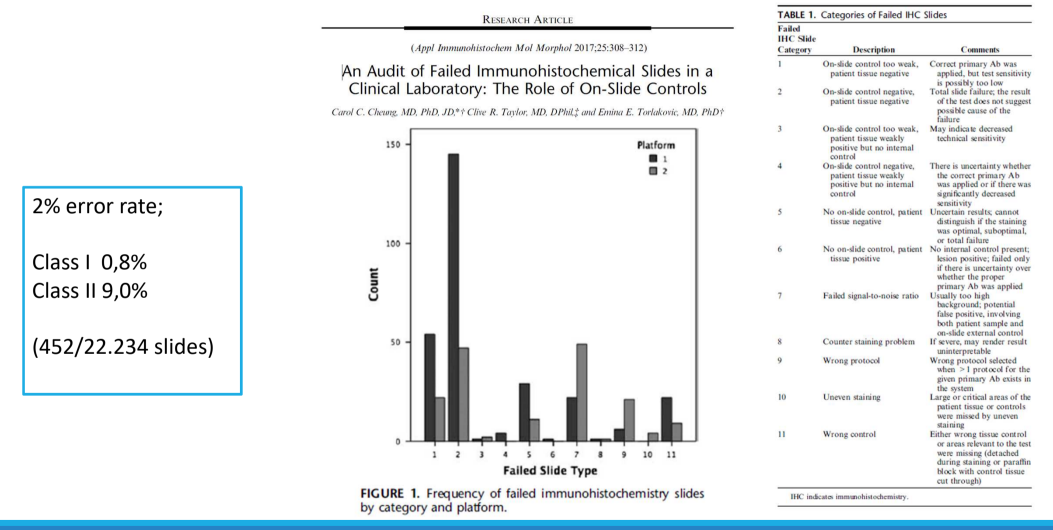


Fig. 5 Evolution of use of on-slide controls in NordiQC

Evolution in the Use of On-Slide Controls for Diagnostic Immunohistochemistry in the Era of Precision Testing Heidi Lykke Kristoffersen, Rasmus Røge, Søren Nielsen. NordiQC, Aalborg Universityhospital, Denmark. **USCAP 2023** 

#### Use of on-slide controls in NordiQC



### A: On-slide controls

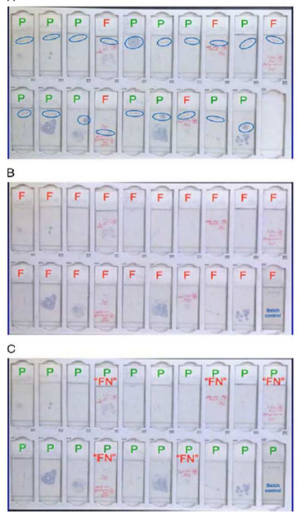
IHC slides stained for ALK (Class II),same run, same instrument, same protocol14/19 passed5/19 failed (5 x 150 USD)

### B: Batch-control - Theoretically:

Batch control <u>failed</u> by same conditions as above 0/19 passed 19/19 failed (no consistent internal control...) (20 x 150 USD)

#### C: Batch-control - Theoretically:

Batch control **passed** by same conditions as above 19/19 passed 0/19 failed (the 5 failed slides not identified....) (Cost...???)



## Conclusions

Focus on external tissue controls is central to standardize and optimize IHC:

- On-slide TMA controls are preferable to 1 bacth control
- Internal tissue controls are of limited value
- Need to generate consensus guidelines on ICAPCs for all IHC tests which tissues, which staining pattern. Interaction of industry, EQA and pathology organisations and societies required.
- Need to identify best practice controls tissues, beads, cell lines.. for type 2 IHC

# Questions and/or comments



Thank You for the attention and.....

54