

Lung tumours

Optimization of antibodies, selection, protocols and controls

NQC Workshop 2019

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Lung markers in NordiQC assessments:

- Iu-ALK (NQC in 2017)
- PD-L1 (NQC in 2019)
- p63 (NQC in 2016)
- p40 (NQC in 2016)
- Napsin A (NQC in 2015)
- TTF-1 (NQC in 2016)
- SYP (NQC in 2018)

- WT1 (NQC in 2019)
- CEA (NQC in 2018)
- Calretinin (NQC in 2018)
- CGA (NQC in 2018)
- Podoplanin (NQC in 2018)
- CD56 (NQC in 2013)



Important prerequisites for an optimal staining result:

- The use of efficient/optimized epitope retrieval protocols (most often HIER in alkaline buffer)
- Careful calibration of primary antibodies with high analytical sensitivity
- The use of highly sensitive detection systems (3-step polymer/multimer systems)



Target	High scoring clones*	Low scoring clones*
lu-ALK	rmAb: D5F3, mAb: OTI1A4	mAb: ALK1 mAb: 5A4
PD-L1	mAb: 22C3, rmAb: 28-8 and rmAb SP263	
p63	mAb: DAK-p63 and 4A4	mAb: 7JUL
p40	mAb: BC28 and rmAb: ZR8	Many pAbs
Napsin A	mAb: IP64 and MRQ-60	pAb: 760-4446 and 352A-7x
TTF1	mAb: SPT24 and SP141	mAb: 8G7G3/1
SYP	mAb: 27G12, rmAb MRQ-40 and DAK-SYNAP	mAb: SY38
WT1	mAb: WT49 and 6F-H2, rmAb: EP122 and D8I7	
CEA	mAb: CEA31 and COL-1	mAb: TF3H8-1 and II-7
CGA	pAb: A0430§ / IR502§, mAb: LK2H10	mAb DAK-A3 and 5H7
Calretinin	rmAb: SP65, mAb DAK-Calret1, pAb 18-0211	rmAb: SP13
Podoplanin	mAb: D2-40	mAb: D2-40 #
CD56	rmAb: MRQ-42, mAb: CD564 and 123C3	mAb: 123C3 #
# Ventana plat	form § Products discontinued * on the	basis of the assessments in NordiQC

Recommended protocols - p63



				Search:	
Epitope 🔷	Staining Platform	Clone name	Clone format	• Version date •	View 🔷
p63	Autostainer, LabVision	DAK-p63	CONC	23 Sep 2015	<u>PDF</u>
p63	Dako Autostainer Link 48 +	DAK-p63	CONC	13 Sep 2015	<u>PDF</u>
p63	Dako Autostainer Link 48 +	4A4	CONC	29 Aug 2016	<u>PDF</u>
p63	Dako Omnis	DAK-p63	CONC	05 Oct 2016	<u>PDF</u>
p63	DBS Montage 360 system	DBR16.1	Other	23 Aug 2016	<u>PDF</u>
p63	Gene Stainer, Gene Tech	4A4	CONC	23 Sep 2015	<u>PDF</u>
p63	Gene Tech Genestainer	4A4	Other	19 Aug 2016	<u>PDF</u>
p63	Leica BOND III	4A4	CONC	12 Sep 2015	<u>PDF</u>
p63	Leica BOND III	4A4	Other	25 Aug 2016	<u>PDF</u>
p63	Leica BOND III	DAK-p63	CONC	30 Aug 2016	<u>PDF</u>
p63	Thermo Autostainer 36/48/72	DAK-p63	CONC	05 Sep 2016	<u>PDF</u>
p63	Ventana Benchmark Ultra	DAK-p63	CONC	16 Sep 2015	<u>PDF</u>
p63	Ventana Benchmark Ultra	4A4	CONC	16 Sep 2015	<u>PDF</u>
p63	Ventana Benchmark Ultra	DAK-p63	CONC	29 Aug 2016	<u>PDF</u>
p63	Ventana Benchmark Ultra	4A4	CONC	02 Sep 2016	<u>PDF</u>

Recommended protocol for p63

Obtained in run 48 29 Aug 2016

Immunostainer

Type:

Ventana Benchmark Ultra

Primary antibody

Clone: Producer: Product no. / lot no.: Diluent: Dilution factor: Incubation time / temperature:

Epitope retrieval, HIER

Device: Buffer: Heating time at max. temp.: Maximum heating temp.:

Visualization system

Producer: Product / no: Incubation time linker: Incubation time polymer: Incubation temperature:

Chromogen

Producer: Product / no: Incubation time / temperature: Enhancement: DAK-p63 Dako M7317 / 20032413 Da Vinci Green 1:100 32 min. / 36°C

On Board / On Machine Ventana Ultra CC1 56 min. 100°C

Ventana

OptiView DAB IHC Detection Kit / 760-700 8 min. 8 min. 36°C

Ventana OptiView DAB IHC Detection Kit / 760-700 8 min. / 22°C CuSO4





Target	Controls, positive	Controls, negative
lu-ALK	Colon/appendix (LE*), lung carc. (lu-ALK pos)	Lung carcinoma (lu-ALK neg)
PD-L1	Tonsil (HE** and LE), placenta (HE) and kit controls	Kit controls
p63	Tonsil (HE and LE) or prostate (LE)	Prostate and tonsil
p40	Tonsil (HE) and placenta (LE)	Tonsil
Napsin A	Kidney (LE) and lung (HE)	Colon/appendix
TTF1	Lung terminal bronchioles (HE and LE)	Liver
SYP	Colon/appendix (HE and LE)	Liver "Onslide" control
WT1	Fallopian tube (LE and HE) and kidney (HE)	Kidney
CEA	Colon/appendix (HE and LE)	Liver
CGA	Colon/appendix (HE and LE) and pancreas (HE)	Liver
Calretinin	Adrenal gland (LE) and appendix (HE and LE)	Appendix
Podoplanin	Tonsil (HE and LE)	
CD56	Tonsil (LE) colon/appendix (HE)	Tonsil
	*Low Expresser **High Expresser	LE = LLOD (Low limit of detection)



Target	High scoring clones*	Low scoring clones*
lu-ALK	rmAb: D5F3, mAb: OTI1A4	mAb: ALK1 mAb: 5A4
PD-L1	mAb: 22C3, rmAb: 28-8 and rmAb SP263	
p63	mAb: DAK-p63 and 4A4	mAb: 7JUL
p40	mAb: BC28 and rmAb: ZR8	Many pAbs
Napsin A	mAb: IP64 and MRQ-60	pAb: 760-4446 and 352A-7x
TTF1	mAb: SPT24 and SP141	mAb: 8G7G3/1
SYP	mAb: 27G12, rmAb MRQ-40 and DAK-SYNAP	mAb: SY38
WT1	mAb: WT49 and 6F-H2, rmAb: EP122 and D8I7	
CEA	mAb: CEA31 and COL-1	mAb: TF3H8-1 and II-7
CGA	pAb: A0430§ / IR502§, mAb: LK2H10	mAb DAK-A3 and 5H7
Calretinin	rmAb: SP65, mAb DAK-Calret1, pAb 18-0211	rmAb: SP13
Podoplanin	mAb: D2-40	mAb: D2-40 #
CD56	rmAb: MRQ-42, mAb: CD564 and 123C3	mAb: 123C3 #
# Ventana plat	form § Products discontinued * on the	basis of the assessments in NordiQC



Concentrated Antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff. ¹	Suff. OPS ²	
mAb clone 12-140-10	3	Leica/Novocastra	0	0	0	3	-	-	
mAb clone CEA31	9 1	Cell Marque BioSB	6	0	3	1	67%	75%	
mAb COL-1	6 5 5 2 1 1	Thermo/Neomarkers Invitrogen/Zymed Biocare Immunologic Zytomed GeneTex	11	7	2	0	90%	94%	
mAb II-7	85	Dako/Agilent	2	19	60	4	25%	58%	
mAb CEA88	2	BioGenex	0	0	1	1	-	-	6% 5% 42%
mAb PARLAM 4	1	Monosan	0	0	1	0	-	-	6% 59% 42%
mAb BS33	1	Nordic Biosite	0	0	1	0			86% 75% 42%
Ready-To-Use Antibodies									2004 2009 2013 2016
mAb clone CEA31 760-4594	53	Ventana/Cell Marque	22	26	5	0	91%	100%	
mAb clone CEA31 236M	4	Cell Marque	1	2	1	0	-	-	
mAb clone COL-1 MAD-002095QD	2	Master Diagnostica	ο	0	1	1	-	-	mAb clone TF3H8- ⁻
mAb clone COL-1 PM058	1	Biocare	o	0	1	0	-	-	cross reacts with
mAb clone COL-1 Kit-0008	1	Maixin	1	0	0	0	-	-	BGP and NCA
mAb clone II-7 IR/IS622/GA622	47	Dako/Agilent	o	6	40	1	13%	-	mAb clone II-7 is
mAb clone II-7 PA0004	12	Leica	o	5	6	1	42%	-	difficult to optimise
mAb clone TF3H8-1 760-2507	13	Ventana/Roche	o	0	0	13	0%	-	
Total	255		43	65	122	25	-		
Proportion			17%	25%	48%	10%	42%		



300

225

150

75

NordiQC	40	CEA / RUN	54 20	018		Pa	SS:	63 %	
Table 1. Antibodies	and a	assessment marks for C	EA, run	54					
Concentrated Antibodies	1	Vendor	Optimal	Good	Borderline	Poor	Suff.1	Suff. OPS ²	
mAb BS33	1	Nordic Biosite	0	0	1	0			
mAb clone CEA31	24	Cell Marque	15	7	2	0	92%	100%	2
mAb COL-1	7 4 4	BioCare Thermo/Neomarkers Immunologic Invitrogen/Zymed Diagnostic BioSystems Genemed GeneTex Leica Zytomed	16	11	1	1	93%	100%	
mAb II-7	48	Dako/Agilent	4	10	9	25	29%	-)	
Ready-To-Use									
mAb clone CEA31 760-4594 mAp clone CEA31		Ventana/Cell Marque	57	22	4	7	88%	94%	
760-4594 ³	1	Ventana/Cell Marque	0	1	0	0	-	-	% % % %
mAb clone CEA31 236M	5	Cell Marque	3	2	o	ο	-	-	86% 75% 59% 63%
mAb clone COL-1 MAD-002095QD	1	Master Diagnostica	0	1	0	0	-	-	
mAb clone COL-1 PM058	1	Biocare	1	0	0	0	-	-	2004 2009 2013 2016 2018
mAb clone COL-1 PA0848	2	Leica	о	1	о	0	-	-	
mAb clone COL-1 Kit-0008	2	Maixin	1	0	1	о	-	-	
mAb clone COL-1	1	Sakura	1	0	o	о	-	-	mAb clone TF3H8-1
mAb clone II-7 GA622	21	Dako/Agilent	о	2	11	8	10%	-	cross reacts with
mAb clone II-7 GA622 ⁴	2	Dako/Agilent	o	0	1	1	-	-	BGP and NCA
mAb clone II-7 IR/IS622	24	Dako/Agilent	1	13	4	6	58%	90%	
mAb clone II-7 IR/IS622 ⁵	4	Dako/Agilent	1	0	1	2	-	-	mAb clone II-7 is
mAb clone II-7 PA0004	5	Leica	о	1	3	1	-	-	difficult to optimise
mAb clone TF3H8-1 760-2507	11	Ventana/Roche	о	0	9	2	0%	-	
Unknown clone	1	Leica	о	ο	1	о	-	-	
Total	272		100	71	48	53	-		
Proportion			37%	26%	18%	19%	63%		555555555555555555555555555555555555555

Lung tun	nours: Ar	itibodies,	protoco	ls and co	ontrols _{No}	
NordiQC	CEA /	RUN 54 20	18	Pass: 63	%	300
Ab	Labs 2018	Labs 2016	Change	% suff.		- 225
COL1	35	24	46 %	91 %		- 150 - 75
CEA31	120	67	79 %	89 %	%98 2004 2009 2013 2016	800 - 0 2018
1]-7	104	144	-28 %	31 %	mAb clone TF3 cross reacts BGP and N	with
TF3H8-1	11	13	-15 %	0 %	mAb clone II- difficult to opt	okokokokok iko

The significant increase in the pass rate seems to be related to an increased used of the mAbs clones CEA31 and COL-1.

CEA / RUN 54 2018



Controls / iCAPC

Positive: Appendix.

* The vast majority of the epithelial cells must show a moderate to strong cytoplasmic staining reaction.

Negative: Liver

* No cells must be positive.

iCAPC: immunohistochemistry Critical Assay Performance Control



CEA / RUN 54 2018



Controls / iCAPC

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

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FIGURE 9. mCEA iCAP. A, A moderate to strong staining reaction must be seen in the brush border of the surface epithelial cells. Virtually all epithelial cells must show a weak to moderate cytoplasmic staining reaction (LLOD). If overexpression of the CEA is desirable target for detection in adenocarcinoma, the demonstration of the staining of only of the surface of mucosa can be selected as LLOD. B, Liver: no staining reaction must be seen. C, Tonsil: scattered squamous epithelial cells show a moderate to strong cytoplasmic staining reaction (number of cells demonstrated will vary from tonsil to tonsil).



CEA / RUN 47 2016

Recommend- able clones	Retrieval	Titre	Detection	RTU	Detection
mAb COL-1	HIER, High pH	1:100 - 1:400	2- or <u>3-step</u>		
mAb CEA31	HIER, High pH	1:100 - 1:400	2- or <u>3-step</u>	Ventana	2- or <u>3-step</u>

Table 3. Optimal results for CEA for the three most commonly used concentrated antibodies on the 3 main IHC systems*

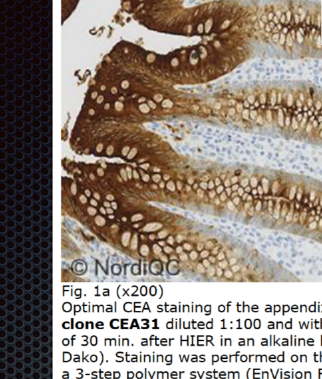
Concentrated antibodies	Da Autostainer Li OM	nk / Classic /	Vent BenchMark		Leica Bond III / Max		
	TRS pH 9.0	TRS pH 6.1	CC1 pH 8.5	CC2 pH 6.0	ER2 pH 9.0	ER1 pH 6.0	
mAb clone II-7	1/17** (6%)	0/2	0/35 (0%)	-	1/10 (10%)	0/4 (0%)	
mAb clone COL-1	1/2	-	8/13 (62%)	-	1/1	-	
mAb clone CEA31	3/3 -		3/6 (50%)	-	-	-	

* 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)

EA31





CEA / RUN 47 2016

Nord

Optimal CEA staining of the appendix using the **mAb clone CEA31** diluted 1:100 and with an incubation time of 30 min. after HIER in an alkaline buffer (TRS pH 9, Dako). Staining was performed on the Dako Omnis using a 3-step polymer system (EnVision Flex+). A weak to moderate staining reaction is seen in the vast majority of the luminal epithelial cells of the appendix, whereas the glycocalyx show an intense staining reaction. Also compare with Figs. 2a – 4a, same protocol. No background staining is seen.

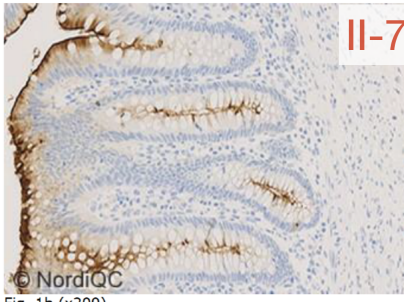


Fig. 1b (x200)

Insufficient CÉA staining of the appendix using the **mAb clone II-7** in a RTU format (Dako GA622) with an incubation time of 25 min. after HIER in an alkaline buffer (TRS pH 9, Dako). Staining was performed on the Dako Omnis using a 3-step polymer system (EnVision Flex+). In spite of very similar protocol settings the "clone II-7"-protocol only demonstrates the glycocalyx distinctively, while the cytoplasmic compartment in the vast majority of epithelial cells is unstained - same field as in Fig. 1a. Also compare with Figs. 2b - 4b, same protocol.

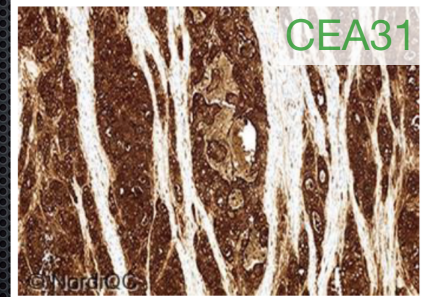


Fig. 2a (x200)

Optimal CEA staining of the colon adenocarcinoma with high level CEA expression using same protocol as in Fig. 1a. Virtually all the neoplastic cells show a strong and distinct cytoplasmic staining reaction. Weak background staining in the vicinity of the neoplastic cells, due to diffusion of antigen, is seen and accepted.

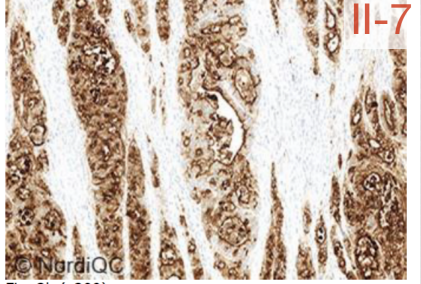


Fig. 2b (x200)

CEA staining of the colon adenocarcinoma with high level CEA expression using same insufficient protocol as in Fig. 1b – same field as in Fig. 2a. The intensity of the neoplastic cells demonstrated is reduced compared to the level expected and obtained in Fig. 2a. Less successful primary antibody: mAb clone II-7





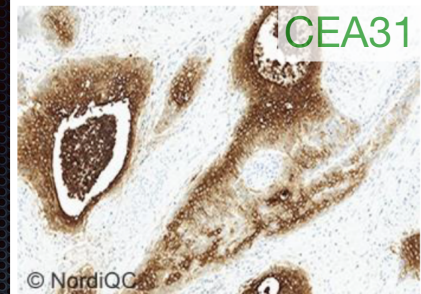
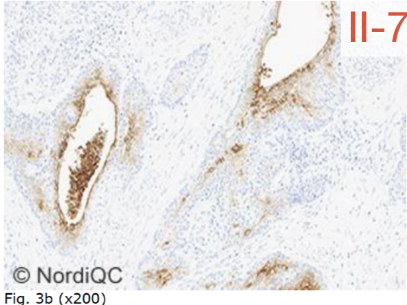


Fig. 3a (x200)

Optimal CEA staining of the urothelial carcinoma, tissue core no. 4, using same protocol as in Figs. 1a and 2a. The majority of the neoplastic cells show a strong and distinct staining reaction. No background staining is seen.



Insufficient CEA staining of the urothelial carcinoma, tissue core no. 4, using same protocol as in Figs. 1b and 2b – same field as in Fig. 3a. The proportion and intensity of the neoplastic cells demonstrated is significantly reduced compared to the level expected and obtained in Fig. 3a.

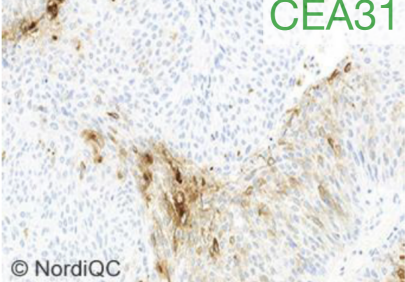


Fig. 4a (x200)

Optimal CEA staining of the urothelial carcinoma, tissue core no. 5, with low level CEA expression using same protocol as in Figs. 1a - 3a. Focally the neoplastic cells show a moderate to strong and distinct staining reaction. No background staining is seen.

© NordiQC Fig. 4b (x200)

Insufficient CEA staining of the urothelial carcinoma, tissue core no. 5, with low level CEA expression using same protocol as in Figs. 1b - 3b - same field as in Fig. 4a.

The neoplastic cells show no staining reaction and a false negative result of the tumour is seen.

Less successful primary antibody: mAb clone II-7





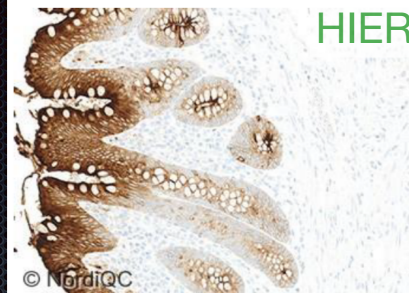


Fig. 5a (x200)

Optimal CEA staining of the appendix using the mAb clone CEA31 diluted 1:400 and with an incubation time of 30 min. after **HIER** in an alkaline buffer (CC1, Ventana). Staining was performed on the Ventana BenchMark using a 3-step multimer system (OptiView)

A weak to moderate staining reaction is seen in the vast majority of the luminal epithelial cells of the appendix, whereas the glycocalyx show an intense staining reaction. Compare also to Fig. 6a, same protocol. © NordiQC

Fig. 5b (x200)

Insufficient CEA staining of the appendix using the mAb clone CEA31 with similar protocol settings as used in Fig. 5a. Only difference was the use of proteolytic pretreatment (Protease 1, Ventana for 8 min.) instead of HIER. Proteolytic pre-treatment results in a drastic reduction in staining intensity. Only the glycocalyx is distinctively demonstrated, while the cytoplasmic compartment of the epithelial cells is unstained - same field as in Fig. 5a. Compare also to Fig. 6b, same protocol.

Protease

© NordiQC

Fig. 6b (x200)

Insufficient CEA staining of the urothelial carcinoma, tissue core no. 5, with low level CEA expression using same protocol as in Fig. 5b – same field as in Fig. 6a. The neoplastic cells show no staining reaction and a false negative result in this tumour is seen. Inappropriate retrieval - use of proteolysis

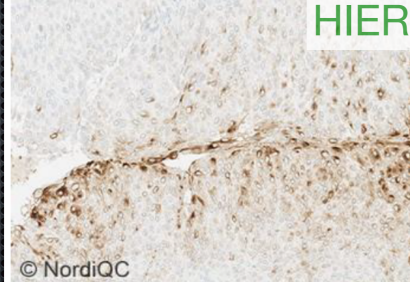


Fig. 6a (x200)

Optimal CEA staining of the urothelial carcinoma, tissue core no. 5, with low level CEA expression using same protocol as in Fig 5a. Focally the neoplastic cells show a moderate to strong and distinct staining reaction.



Inappropriate antibody -NCA and BGP cross reaction

CEA31

© NordiQC

Fig. 7a (x200)

CEA / RUN 47 2016

Optimal CEA staining of the liver using same protocol as in Figs. 5a and 6a based on the **mAb clone CEA31**. No staining reaction is seen in the Kupffer cells, leucocytes and the bile canaliculi. No background staining is seen.

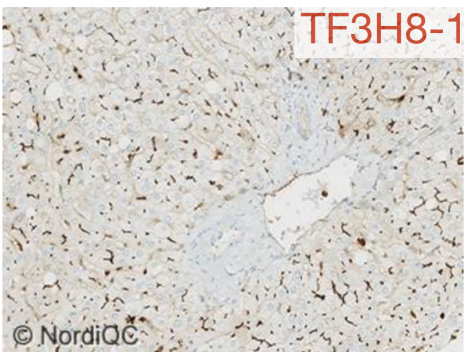


Fig. 7b (x200)

Insufficient CEA staining of the liver using the **mAb clone TF3H8-1.** Both the Kupffer cells, leucocytes and bile canaliculi are stained due to a cross reaction of the Ab to NCA (CEACAM6) and BGP (CEACAM1) – same field as in Fig. 7a.

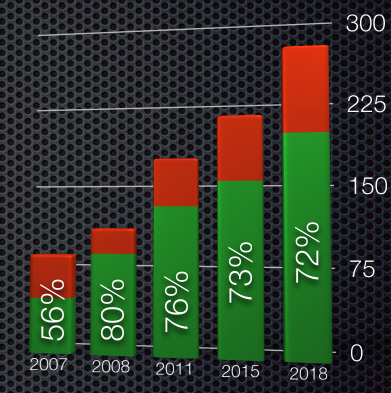


 Calretinin / RUN 52 2018
 Pass: 72 %

 odies and assessment marks for CR, run 52

 oties and assessment marks for

Concentrated antibodies vendor Optimal Good Borderline Poor Suff. ¹ OPS ² mAb clone 2E7 1 Immunologic 1 0 0 - - mAb clone CAL6 7 Leica/Novocastra 1 1 2 0 - - mAb clone CAL6 7 Leica/Novocastra 1 3 0 3 57% - mAb clone BAK-Cairet 1 Dako/Agilent 9 8 8 9 50% 81% rmAb clone SP13 3 Cell Marque 0 0 0 - - pAb 18-0211 12 Invitrogen/Thermo 3 3 4 2 50% 100% pAb, 232A 2 Cell Marque 0 0 1 0 - - pAb 61-0006 1 Genemed 0 0 1 0 0 1 - pAb ReK03 14 Leica/Novocastra 1 11<	Table 1. Antibodies and assessment marks for CR, run 52									
mAb clone 5A5 3 Leica/Novocastra 1 1 2 0 - mAb clone CAL6 7 Leica/Novocastra 1 3 0 3 57% - mAb clone DAK-Calret 34 Dako/Agilent 9 88 88 9 50% 81% rmAb clone BSR235 1 Nordic Biosite 1 0 0 0 - - rmAb clone SP13 2 Cell Marque 1 0 0 0 - - pAb 18-0211 12 Invitrogen/Thermo 3 3 44 2 50% 100% pAb 18-0211 12 Invitrogen/Thermo 3 3 44 2 50% 100% pAb, 232A 2 Cell Marque 0 0 1 0 - - pAb, CP092C 1 Biocare 0 0 1 0 - - mAb clone CAL6 14 Leica/Novocastra 1 11 11 2 0 86% 92% mAb clone DAK	Concentrated antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff.1		
mAb clone SAS 1 Monosan 1 1 1 2 0 - - mAb clone CAL6 7 Leica/Novocastra 1 3 0 3 57% - mAb clone DAK-Cairet 3 Dako/Agilent 9 8 8 9 50% 81% rmAb clone BSR235 1 Nordic Biosite 1 0 0 0 - - rmAb clone SP13 2 Cell Marque 1 3 4 1 44% - pAb 18-0211 12 Invitrogen/Thermo 3 3 4 2 50% 100% pAb 61-0006 1 Genemed 0 0 1 0 - - pAb 61-0006 1 Biocare 0 0 1 0 - - pAb 70-052 1 Zytomed Systems 0 0 0 1 - - mAb clone CAL6 14 Leica/Novocastra 1 11 11 2 0 86% 92% mA	mAb clone 2E7	1	Immunologic	1	0	0	0	-	-	
mAb clone DAK-Cairet 34 Dako/Agilent 9 8 8 9 50% 81% rmAb clone BSR235 1 Nordic Biosite 1 0 0 0 - - rmAb clone SP13 2 Cell Marque 1 0 0 0 - - pAb 18-0211 12 Invitrogen/Thermo 3 3 4 2 50% 100% pAb, 322A 2 Cell Marque 0 0 2 0 - - pAb 61-0006 1 Genemed 0 0 1 0 - - pAb 61-0006 1 Enemed 0 0 1 0 - - pAb 61-0006 1 Enemed 0 0 1 - - pAb CP092C 1 Biocare 0 0 1 - - mAb clone CAL6 14 Leica/Novocastra 1 11 1 2	mAb clone 5A5			1	1	2	0	-	-	
1 34 Dako/Agilent 9 8 8 9 50% 81% rmAb clone BSR235 1 Nordic Biosite 1 0 0 0 - - rmAb clone SP13 2 Cell Marque 1 3 4 1 44% - pAb 18-0211 12 Invitrogen/Thermo 3 3 4 2 50% 100% pAb 18-0211 12 Invitrogen/Thermo 3 3 4 2 50% 100% pAb 18-0211 12 Invitrogen/Thermo 3 3 4 2 50% 100% pAb 50-006 1 Genemed 0 0 1 0 - - pAb 61-0006 1 Biocare 0 0 1 0 - - pAb RBK003 1 Zytomed Systems 0 0 0 - - mAb clone CAL6 14 Leica/Novocastra 1 11 1 2 0 94% 97% mAb clone DAK-Cairet 1 <td>mAb clone CAL6</td> <td>7</td> <td>Leica/Novocastra</td> <td>1</td> <td>3</td> <td>0</td> <td>3</td> <td>57%</td> <td>-</td>	mAb clone CAL6	7	Leica/Novocastra	1	3	0	3	57%	-	
3 Cell Marque Immunologic 2 1 3 4 1 44% - pAb 18-0211 12 Invitrogen/Thermo 3 3 4 2 50% 100% pAb, 232A 2 Cell Marque 0 0 2 0 - - pAb 61-0006 1 Genemed 0 0 1 0 - - pAb 61-0006 1 Genemed 0 0 1 0 - - pAb 61-0006 1 Genemed 0 0 1 0 - - pAb 62006 14 Sytomed Systems 0 0 0 1 - - mAb clone CAL6 14 Leica/Novocastra 1 11 2 0 86% 92% mAb clone DAK-Cairet 1 Leica/Novocastra 1 11 0 0 - - mAb clone DAK-Cairet 1 Leica/Novocastra 1 0 0 - - mAb clone DAK-Cairet 1 Maixin		34	Dako/Agilent	9	8	8	9	50%	81%	
rmAb clone SP13 2 Immunologic Spring Bioscience 2 1 3 4 1 44% - pAb 18-0211 12 Invitrogen/Thermo 3 3 4 2 50% 100% pAb, 232A 2 Cell Marque 0 0 2 0 - - pAb, 232A 2 Cell Marque 0 0 1 0 - - pAb, 232A 2 Cell Marque 0 0 1 0 - - pAb, 2006 1 Biocare 0 0 1 0 - - pAb RBK03 1 Zytomed Systems 0 0 0 1 - - mAb clone CAL6 PA0346 14 Leica/Novocastra 1 11 1 2 0 86% 92% mAb clone DAK-Calret 1 S/IR627 ⁴ 35 Dako/Agilent 14 19 2 0 94% 97% mAb clone MX027 1 Maixin 1 0 0 1 - -	rmAb clone BSR235	1	Nordic Biosite	1	0	0	0	-	-	
pAb, 232A 2 Cell Marque 0 0 2 0 - - pAb 61-0006 1 Genemed 0 0 1 0 - - pAb, CP092C 1 Biocare 0 0 1 0 - - pAb, REK003 1 Zytomed Systems 0 0 0 1 - - Ready-To-Use antibodies 1 Leica/Novocastra 1 11 2 0 86% 92% mAb clone CAL6 14 Leica/Novocastra 1 11 2 0 86% 92% mAb clone DAK-Cairet 1 Leica/Novocastra 1 0 0 - - mAb clone DAK-Cairet 15 Dako/Agilent 14 19 2 0 94% 97% mAb clone MX-Cairet 1 Maixin 1 0 0 - - mAb clone MX027 1 Maixin 1 0 0 - - mAb SP13 232R 1 Cell Marque 0	rmAb clone SP13	2 2	Immunologic Spring Bioscience	1	3	4	1	44%	-	
pAb 61-0006 1 Genemed 0 0 1 0 - - pAb, CP092C 1 Biocare 0 0 1 0 - - pAb RBK003 1 Zytomed Systems 0 0 0 1 - - Ready-To-Use antibodies - - - - - - - MAb Clone CAL6 PA0346 14 Leica/Novocastra 1 11 2 0 86% 92% mAb clone DAK-Calret 1 IS/IR627 1 Leica/Novocastra 1 0 0 - - mAb clone DAK-Calret 1 IS/IR627 20 Dako/Agilent 14 19 2 0 94% 97% mAb clone DAK-Calret 1 IS/IR627 20 Dako/Agilent 0 4 11 5 20% - mAb clone MX027 MAB-0716 1 Maixin 1 0 0 0 - - mAb SP13 XMAD- 000315QD 1 Master Diagnostica 0 0 1 0 - - pAb S2	pAb 18-0211	12	Invitrogen/Thermo	3	3	4	2	50%	100%	
pAb, CP092C 1 Biocare 0 0 1 0 - - pAb RBK003 1 Zytomed Systems 0 0 0 1 - - Ready-To-Use antibodies I Leica/Novocastra 1 11 2 0 86% 92% mAb clone CAL6 PA0346 14 Leica/Novocastra 1 0 0 0 - - mAb clone CAL6 PA03463 14 Leica/Novocastra 1 0 0 0 - - mAb clone DAK-Cairet 1 IS/IR627 35 Dako/Agilent 14 19 2 0 94% 97% mAb clone MX027 mAb clone MX027 35 Dako/Agilent 0 4 11 5 20% - rmAb SP13 MAD- 000315QD 1 Maixin 1 0 0 1 - - rmAb SP13 RMPD010 1 Diagnostic Biosystems 0 1 0 - - rmAb SP13 RMPD010 1 Diagnostic Biosystems 0 1 0 - -		2			0	2	0	-	-	
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Ready-To-Use antibodiesII <td></td> <td>1</td> <td></td> <td></td> <td></td> <td></td> <td></td> <td>-</td> <td>-</td>		1						-	-	
antibodies Image: constraint of the second sec		1	Zytomed Systems	0	0	0	1	-	-	
PA0346 14 Leica/Novocastra 1 11 2 0 86% 92% mAb clone CAL6 1 Leica/Novocastra 1 0 0 0 - - mAb clone DAK-Cairet 35 Dako/Agilent 14 19 2 0 94% 97% mAb clone DAK-Cairet 35 Dako/Agilent 14 19 2 0 94% 97% mAb clone DAK-Cairet 20 Dako/Agilent 0 4 11 5 20% - mAb clone MX027 Maixin 1 0 0 0 - - mAb SP13 232R 1 Cell Marque 0 0 1 0 - - rmAb SP13 3 MAD- 0 0 1 0 0 - - - rmAb clone SP65 790- 1 Master Diagnostica 0 1 0 0 - - rmAb sp13 RMPD010 1 Diagnostic Biosystems 0 1 0 2 0 - - pAb 2										
PA03463 1 Leica/Novocastra 1 0 0 0 - <td></td> <td>14</td> <td>Leica/Novocastra</td> <td>1</td> <td>11</td> <td>2</td> <td>0</td> <td>86%</td> <td>92%</td>		14	Leica/Novocastra	1	11	2	0	86%	92%	
1 IS/IR627 35 Dako/Agilent 14 19 2 0 94% 97% mAb clone DAK-Calret 20 Dako/Agilent 0 4 11 5 20% - mAb clone MX027 1 Maixin 1 0 0 0 - - mAb clone MX027 1 Maixin 1 0 0 0 - - rmAb SP13 232R 1 Cell Marque 0 0 1 0 - - rmAb SP13 MAD- 000315QD 1 Master Diagnostica 0 0 1 0 - - rmAb SP13 RMPD010 1 Diagnostic Biosystems 0 1 0 - - rmAb clone SP65 790- 4467 118 Ventana/Roche 86 20 10 2 90% 96% pAb 8223-C010 1 Sakura Finetek 0 1 0 - - - Unknown RTU Ab 1 0 0 1 0 - - -		1	Leica/Novocastra	1	0	0	0	-	-	
1 IS/IR6274 20 Dako/Aglient 0 4 11 5 20% - mAb clone MX027 1 Maixin 1 0 0 0 - - rmAb SP13 232R 1 Cell Marque 0 0 1 0 - - rmAb SP13 MAD- 000315QD 1 Cell Marque 0 0 1 0 - - rmAb SP13 MAD- 000315QD 1 Diagnostic Biosystems 0 1 0 0 - - rmAb SP13 RMPD010 1 Diagnostic Biosystems 0 1 0 0 - - rmAb clone SP65 790- 4467 118 Ventana/Roche 86 20 10 2 90% 96% pAb 232A-78 2 Cell Marque 0 0 2 0 - - pAb 8223-C010 1 Sakura Finetek 0 1 0 0 - - Unknown RTU Ab 1 0 0 1 0 - -		35	Dako/Agilent	14	19	2	0	94%	97%	
MAB-0716 1 Maixin 1 0 0 0 - - rmAb SP13 232R 1 Cell Marque 0 0 1 0 - - rmAb SP13 MAD- 000315QD 1 Master Diagnostica 0 0 1 0 - - rmAb SP13 RMPD010 1 Diagnostic Biosystems 0 1 0 0 - - rmAb clone SP65 790- 4467 118 Ventana/Roche 86 20 100 2 90% 96% pAb 232A-78 2 Cell Marque 0 0 1 0 - - pAb 8223-C010 1 Sakura Finetek 0 1 0 0 - - Unknown RTU Ab 1 - 0 0 1 0 - -		20	Dako/Agilent	0	4	11	5	20%	-	
rmAb SP13 MAD- 000315QD1Master Diagnostica0010rmAb SP13 RMPD010 rmAb clone SP65 790- 44671Diagnostic Biosystems0100rmAb clone SP65 790- 4467118Ventana/Roche8620100290%96%pAb 232A-782Cell Marque0020pAb 8223-C0101Sakura Finetek0100Unknown RTU Ab1		1	Maixin	1	0	0	0	-	-	
000315QD 1 Master Diagnostica 0 0 1 0 - - - rmAb SP13 RMPD010 1 Diagnostic Biosystems 0 1 0 0 - - rmAb clone SP65 790- 4467 118 Ventana/Roche 86 20 10 2 90% 96% pAb 232A-78 2 Cell Marque 0 0 2 0 - - pAb 8223-C010 1 Sakura Finetek 0 1 0 0 - - Unknown RTU Ab 1 Image: Colore of the second of	rmAb SP13 232R	1	Cell Marque	0	0	1	0	-	-	
rmAb clone SP65 790- 4467 118 Ventana/Roche 86 20 10 2 90% 96% pAb 232A-78 2 Cell Marque 0 0 2 0 - - pAb 8223-C010 1 Sakura Finetek 0 1 0 0 - - Unknown RTU Ab 1 Image: Constraint of the second		1	Master Diagnostica	0	0	1	0	-	-	
4467 118 Ventana/Roche 86 20 10 2 90% 96% pAb 232A-78 2 Cell Marque 0 0 2 0 - - pAb 8223-C010 1 Sakura Finetek 0 1 0 0 - - Unknown RTU Ab 1 Image: Constraint of the second	rmAb SP13 RMPD010	1	Diagnostic Biosystems	0	1	0	0	-	-	
pAb 8223-C010 1 Sakura Finetek 0 1 0 0 - - Unknown RTU Ab 1 0 0 1 0 - - -		118	Ventana/Roche	86	20	10	2	90%	96%	
Unknown RTU Ab 1 0 0 1 0	pAb 232A-78	2	Cell Marque	0	0	2	0	-	-	
	pAb 8223-C010	1	Sakura Finetek	0	1	0	0	-	-	
Total 269 120 74 52 23 -	Unknown RTU Ab	1		0	0	1	0	-	-	
	Total	269		120	74	52	23	-		
Proportion 45% 27% 19% 9% 72%	Proportion			45%	27%	19%	9%	72%		



The mAb clone DAK-Calret 1 based RTU system developed for the Autostainer - but used on the Dako Omnis platform

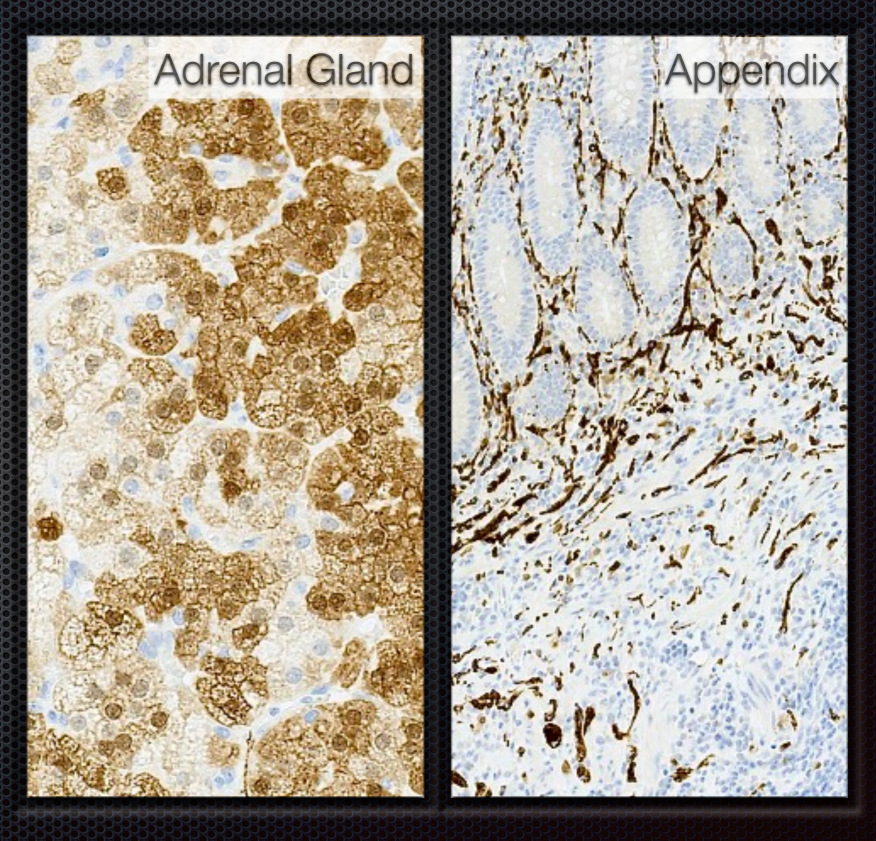
Calretinin / RUN 52 2018



Controls / iCAPC

Adrenal gland will serve as a "low-level expressor" (LE) positive tissue control, in which an at least weak to moderate, distinct cytoplasmic and nuclear staining of the majority of the cortical epithelial cells must be seen.

Appendix serves both as negative tissue and "high-level expressor" (HE) positive tissue control. Columnar epithelial cells and smooth muscle cells should be negative, while strong, distinct cytoplasmic and nuclear staining of the peripheral nerves (ganglion cells and axons) and macrophages should be seen. Furthermore, fat cells in the submucosa of the appendix could serve as an additional LE positive tissue control.



Calretinin / RUN 52 2018



Recommend- able clones	Retrieval	Titre	Detection	RTU	Detection
mAb DAK- Calret1	HIER, High pH	1:20 - 1:100	3-step	Dako	2 or 3-step
mAb 5A5	HIER, High pH	1:100	3-step		
mAb CAL6	HIER, High pH	1:15	3-step	Leica	3-step
pAb 18-0211	HIER, High pH	1:50 - 1:150	3-step		

rmAb SP65 HIER, High pH

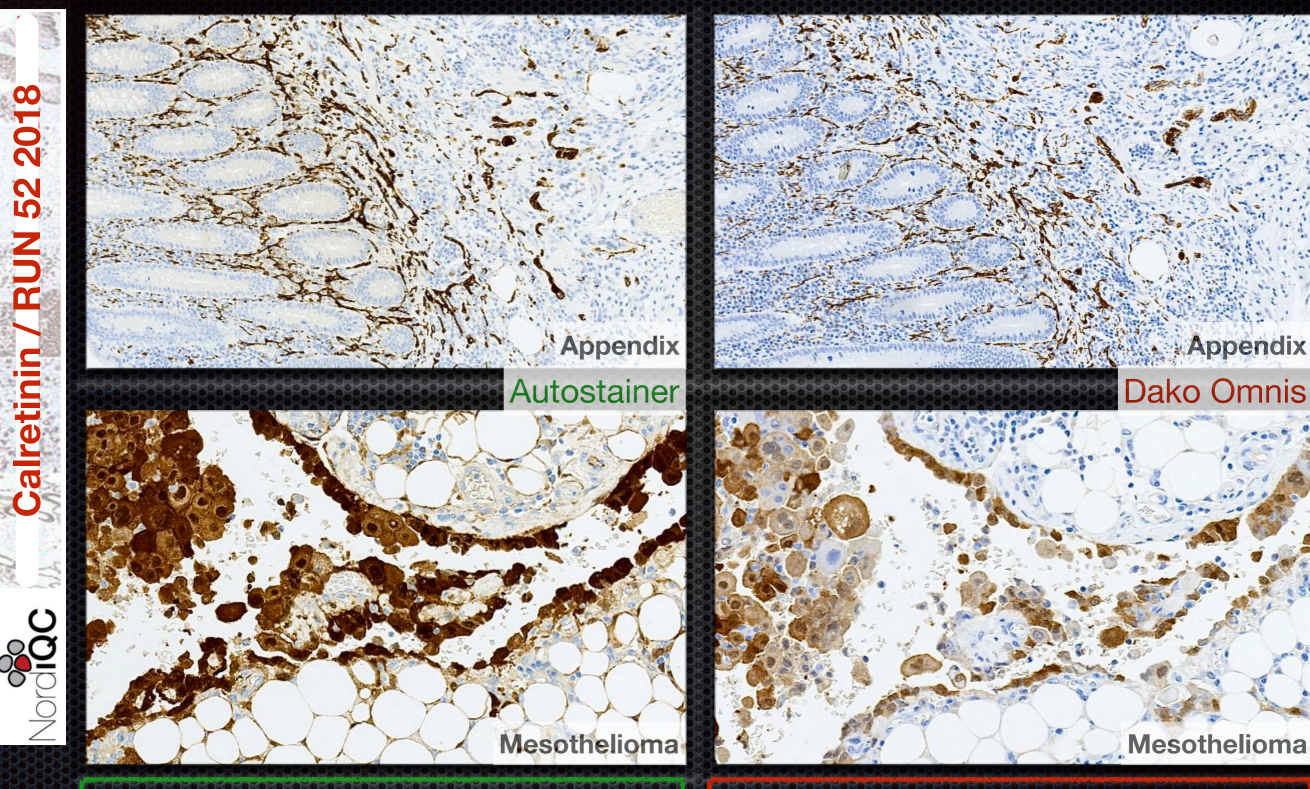
NordiQC

Ventana

2 or 3-step

Table 3. Proportion of optimal results for CR for the most commonly used antibodies as concentrates on the 4 main IHC systems*									
Concentrated		Dako Autostainer Link /		Dako		ana	Leica		
antibodies			Omnis		BenchMark		Bond II	Г/мах	
	Clas	SIC			GX / XT	/ Ultra			
	TRS pH	TRS pH	TRS pH	TRS pH		CC2 pH	ER2 pH	ER1 pH	
	9.0	6.1	9.0	6.1	CC1 pH 8.5	6.0	9.0	6.0	
mAb clone CAL6	-	-	1/2 **	-	0/1	-	0/2	0/1	
mAb clone DAK-Calret 1	3/10 (30%)	-	0/6	-	0/6	-	5/7 (71%)	0/1	
rmAb clone SP13	-	-	-	-	0/4	-	-	-	
pAb 18-0211	1/2	1/1	-	-	0/6	-	0/1	1/1	



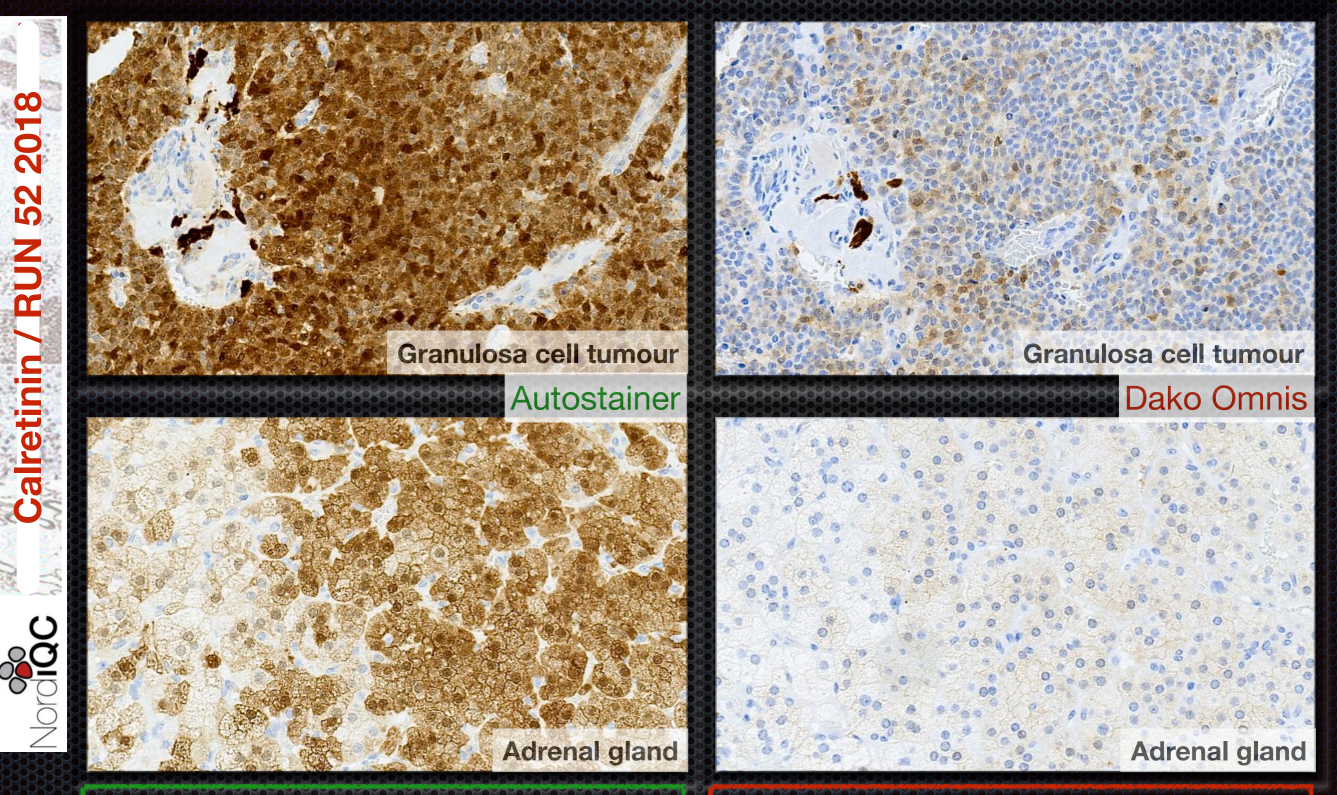


The mAb clone DAK-Calret 1 based RTU system developed for the Autostainer.

The mAb clone DAK-Calret 1 based RTU system developed for the Autostainer - but used on the Dako Omnis platform.

NordiQC

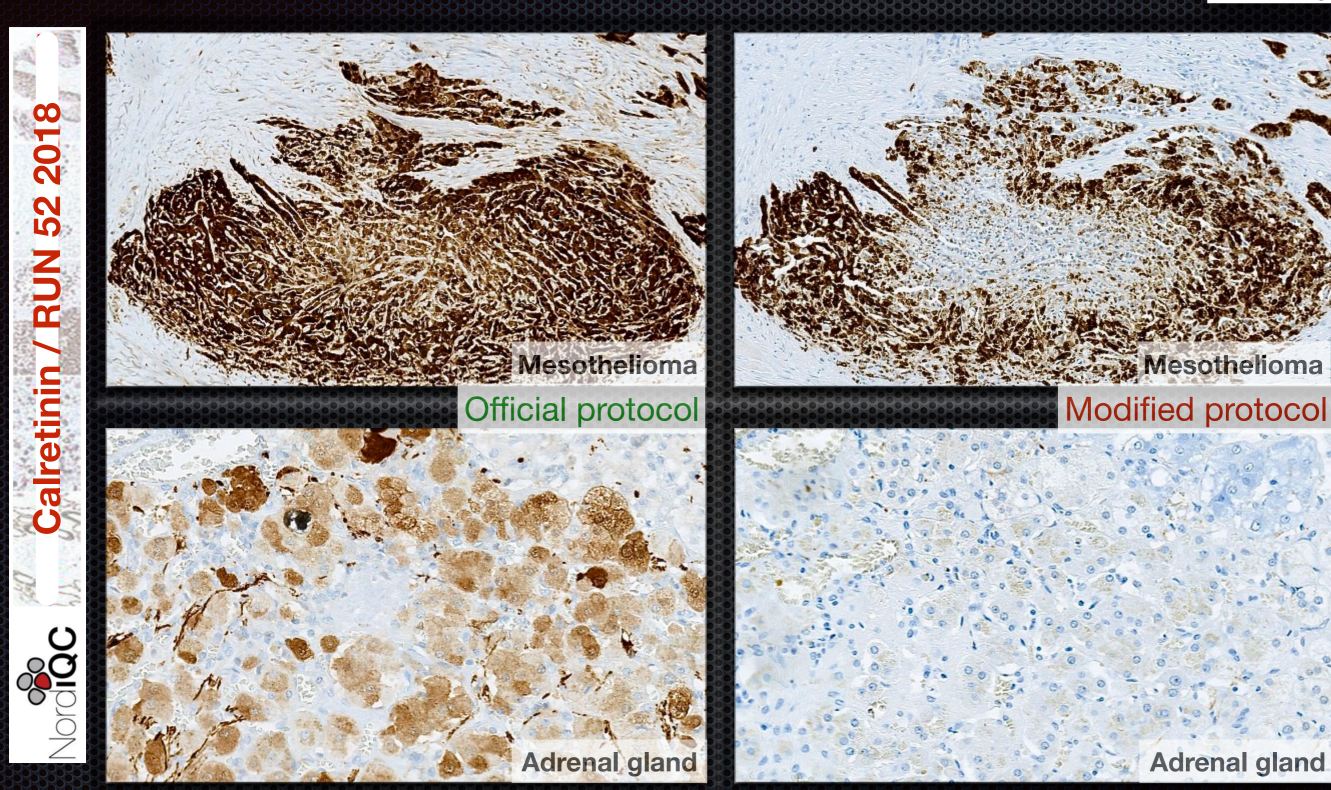
Lung tumours: Antibodies, protocols and controls



The mAb clone DAK-Calret 1 based RTU system developed for the Autostainer.

The mAb clone DAK-Calret 1 based RTU system developed for the Autostainer - but used on the Dako Omnis platform





The rmAb clone SP65 based RTU system for the Ventana BenchMark platform. Official protocol: CC1 32 min / Ab 16 / Optiview The rmAb clone SP65 based RTU system for the Ventana BenchMark platform. Modified protocol: CC1 4 min / Ab 16 / Optiview+Amp



150

100

lu-ALK / RUN 51 2017 Pass: 61 % Table 1. Antibodies and assessment marks for lu-ALK, run 51 200 Suff.1 Suff. Concentrated antibodies n Vendor Optimal Good Borderline Poor OPS^2 43 Leica/Novocastra 1 Abcam 7 mAb clone **5A4** 1 15 24 34% 22% 1 Biocare Monosan 1 1 ThermoFisher 2 Dako 3 mAb clone ALK1 0 0 0 --1 Cell Margue 3 2 rmAb clone **D5F3** 23 Cell Signaling 6 12 78% 94% 3 0 0 mAb clone OTI1A4 13 ORIGENE 10 100% 100% 67% 49% % 50 Ready-To-Use 61 antibodies mAb clone 5A4 Leica/Novocatra 0 0 6 0 6 --PA0306 0 mAb clone 5A4 2013 Maixin 0 0 1 0 2015 1 --2017 MAB-0281 mAb 5A4 Master Diagnostica 0 0 0 1 1 -MAD-001720QD mAb clone 5A4 0 0 0 ThermoFisher 1 1 -_ MS-1104-R7 mAb ALK1 9 Dako 0 0 1 8 --IR641 mAb clone ALK1 is mAb clone ALK1 4 Dako 0 0 0 4 -GA641 not "Fit for purpose" mAb clone ALK1 7 5 Ventana 0 0 2 -790/800-2918 rmAb clone SP8 1 BioGenex 0 0 0 1 --AN770 mAb clone 5A5 is rmAb clone D5F3 70 100% Ventana 53 12 4 1 93% 790-4796 difficult to optimise rmAb clone D5F3 2 0 0 1 1 Ventana --790-4796³ mAb clone OTI1A4 1 Sakura Finetek 1 0 0 0 _ _ 8344-C010 189 72 43 31 Total 43 -

38%

23%

23%

16%

61%

Proportion

lu-ALK / RUN 51 2017



Recommend- able clones	Retrieval	Titre	Detection	RTU	Detection
mAb OTI1A4	HIER, High pH	1:100 - 1:1500	3-step		
rmAb D5F3	HIER, High pH	1:50 - 1:200	3-step +/- amp	Ventana	3-step + amp
mAb 5A4	HIER, High pH	1:20	3-step + amp	No optimal	

Table 3. Proportion of optimal results for lu-ALK for the most commonly used antibodies as concentrate on the 4 main IHC systems*

Concentrated antibodies	Dako Autostainer Link / Classic		Dako Omnis		Vent BenchMark		Leica Bond III / Max			
	TRS pH 9.0	TRS pH 6.1	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 5A4	0/9** (0%)		0/3	-	1/22 (5%)	-	0/9 (0%)	0/1		
mAb clone OTI1A4	2/2	-	5/5 (100%)	-	1/2	-	1/1	-		
rmAb clone D5F3	2/3	0/1	0/3	- (2/6 (33%)	-	2/7 (29%)	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)

lu-ALK / RUN 51 2017



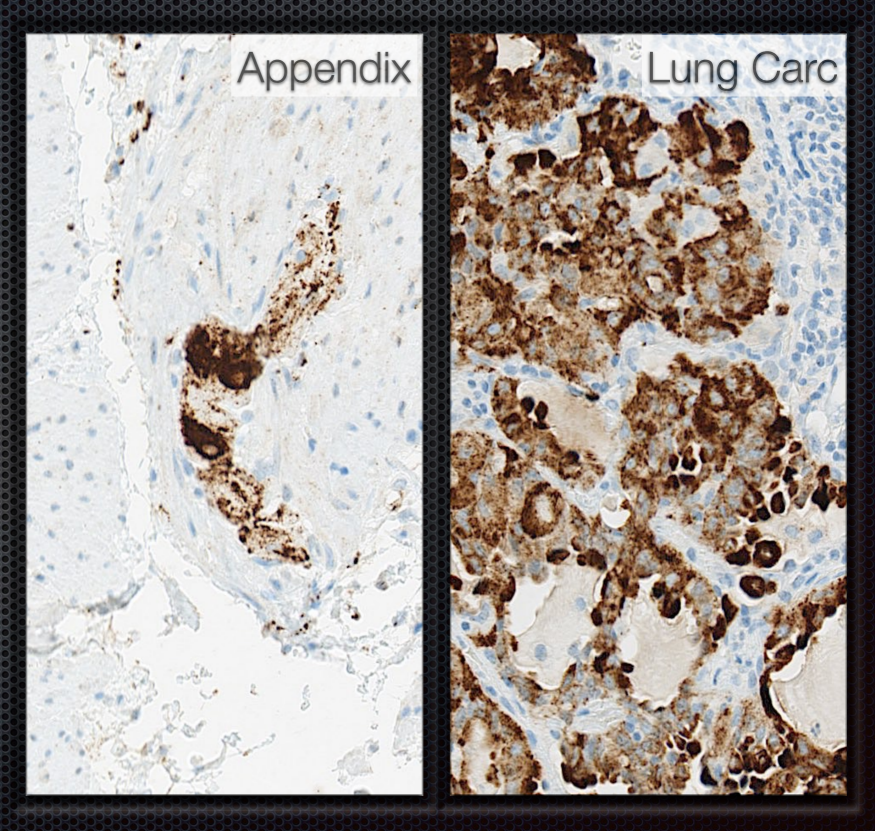
Controls / iCAPC

Positive: ALCL and lung adenocarcinoma with FISH verified ALK rearrangements and normal appendix.

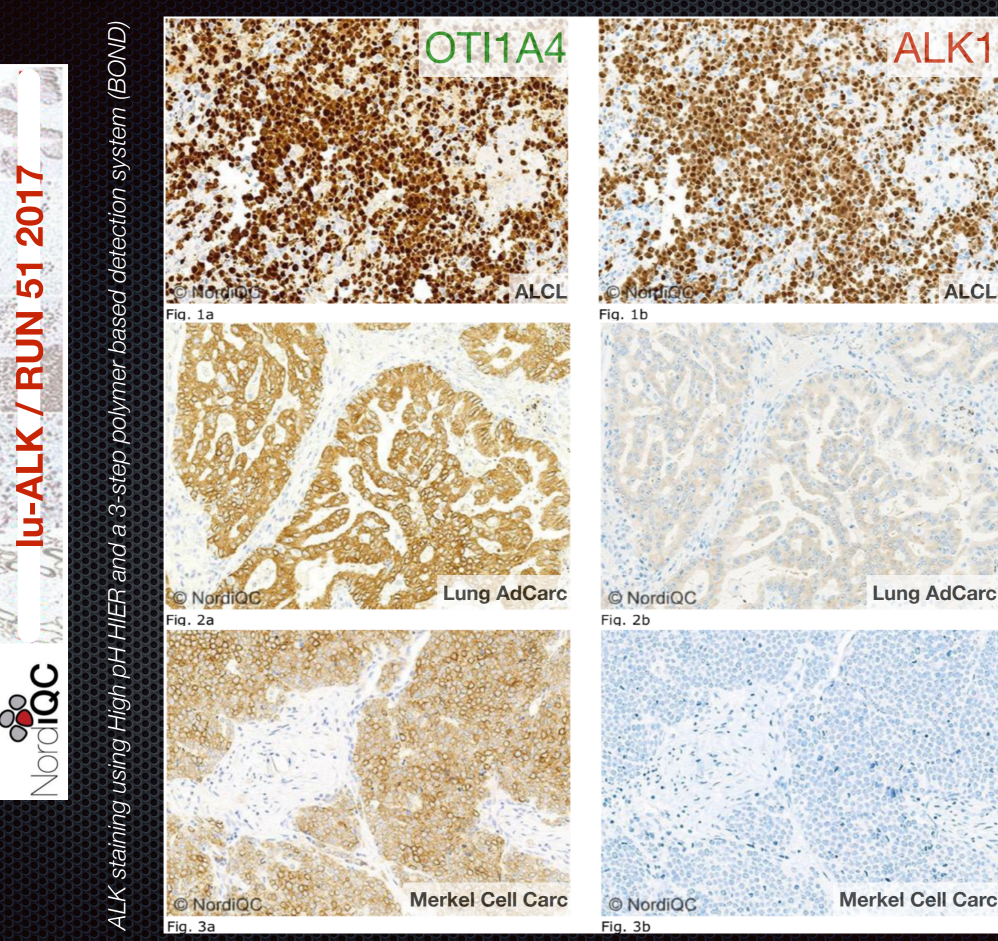
NordiQC

* A weak to strong granular cytoplasmic staining reaction should be seen in the ganglion cells in appendix.

<u>Negative</u>: Lung cancer without ALK rearrangements

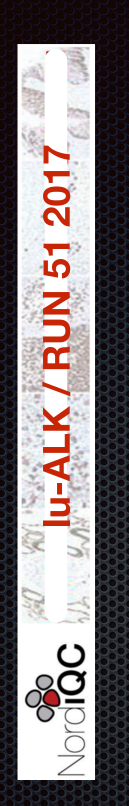




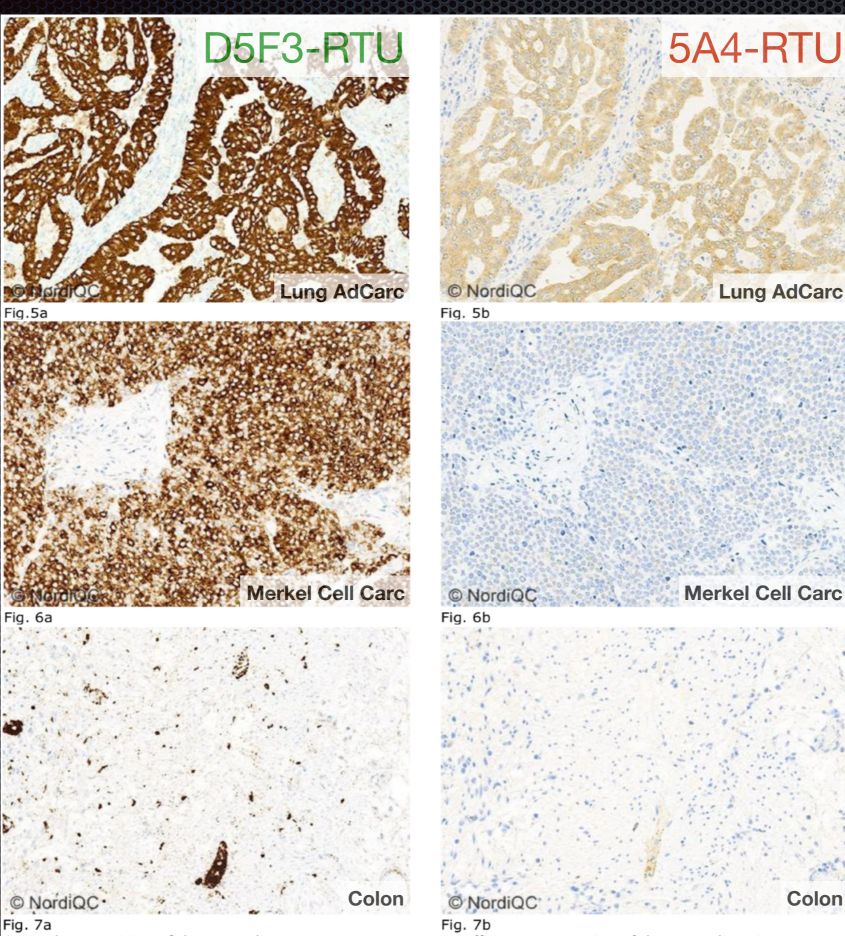


Less successful primary antibody: mAb clone ALK1









Less successful **RTU** system based on mAb clone 5A4

Colon



Table 1. Antibodies a	nd a	ssessment marks for p6	3, run 48										300
Concentrated antibodies	n	Vendor	Optimal	Good	Borderline	Poor	Suff.1	Suff. OPS ²					
	26 4 3 2 2	BioCare Medical ImmunoLogic Dako Zeta Corporation Thermo Scientific Zytomed Systems											225
mAb clone 4A4	2 1 1 1 1 1 1	BioGenex Diagnostic BioSystems Klinipath Minarini Nordic Biosite Santa Cruz	13	20	11	2	72%	76%			0%	82%	- 150 - 75
mAb clone DAK-p63	47	Dako	20	21	6	0	87%	91%		95%			
mAb clone 7JUL	12	Leica/Novocastra	0	1	3	8	8%	-		S			
mAb clone SFI-6	2	DCS Immunoline	0	0	2	0	-	-		O			
rmAb clone BSR6	1	Nordic Biosite	0	0	1	0	-	-	83%				0
rmAb clone DBR16.1	1	Diagnostic Biosystems	1	0	0	0			2006	2000	0014	0010	- 0
rmAb clone EPR5701	1	Epitomics	0	0	1	0	-	-	2000	2009	2014	2016	
Jnknown Ab	1	Unknown	1	0	0	0	-	-					
Ready-To-Use antibodies													
mAb clone 4A4 790-4509	102	Ventana	59	36	5	2	93%	95%					
mAb clone DAK-p63 IR662	46	Dako	21	23	2	0	96%	94%					
mAb clone 4A4 P M163	3	BioCare	1	1	1	0	-	-					
mAb clone 7JUL PA0103	5	Leica/Novocastra	0	0	3	2	-	-					
mAb clone 4A4 AM418	2	BioGenex	0	1	0	1	-	-					
mAb clone 4A4 ARB- 56695	1	Nordic Biosite	1	0	0	0	-	-					
mAb clone MX013 MAB-0694	1	Maixin	0	1	0	0	-	-					
mAb clone 4A4 MAD- 000479QD	3	Master Diagnostica SL	3	0	o	0	-	-					
Total	274		120	104	35	15	-						
Proportion	<u> </u>		44 %	38 %	13 %	5 %	82 %		559595959595				



p63 / RUN 48 2016

Recommend- able clones	Retrieval	Titre	Detection	RTU	Detection
mAb 4A4	HIER, High pH	1:50 - 1:600	3-step	Ventana	3-step
mAb DAK- p63	HIER, High pH	1:50 - 1:300	2- or <u>3-step</u>	Dako	2- or <u>3-step</u>

Table 3. Proportion of optimal results for p63 using concentrated antibodies on the 3 main IHC systems*									
Concentrated	Dak		Ventar	na	Leica				
antibodies	Autostainer	r / Omnis	BenchMark X	(T / 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 DAK-p63	3/15 (20%)**	0/1	14/20 (70%)	-	1/4 (25%)	-			
mAb clone 4A4	0/6 (0%)	-	9/22 (41%)	- /	2/8 (25%)	0/1			
* Antibody concentration platforms.	tion applied as listed at	oove, HIER buffers	and detection kits use	ed as recommend	ded by the vendors	of the respective			

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

p63 RTU antibodies / Plug-and-play ??



RTU: Ready To Use or Ready To Optimise??

Table 4 summarises the proportion of sufficient and optimal marks for the most commonly used RTU systems. The performance is evaluated both as a true plug-and-play system performed according to the recommendations provided by the vendor and by a laboratory modified system changing basal protocol settings. Only protocols performed on the specific IHC stainer device were included, whereas e.g. Dako RTU Ab formats applied on a Ventana stainer were excluded.

Table 4. Proportion of sufficient and optimal results for p63 for the most commonly used RTU IHC systems

RTU systems		mended settings*		ry modified settings**
	Sufficient	Optimal	Sufficient	Optimal
Dako AS48 mAb DAK-p63 IR662	93% (14/15)	60% (9/15)	94% (16/17)	24% (4/17)
VMS Ultra/XT mAb 4A4 790-4509	60% (3/5)	20% (1/5)	95% (89/94)	60% (56/94)

* 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 >25%, detection kit – only protocols performed on the specified vendor IHC stainer included.

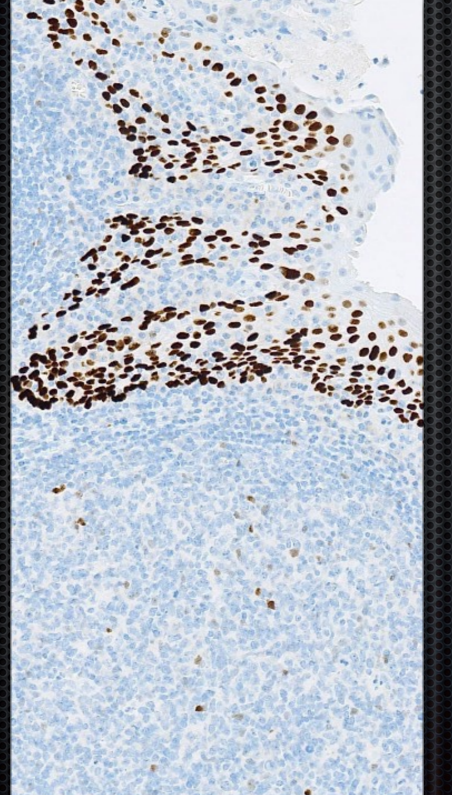
p63 / RUN 41 2014

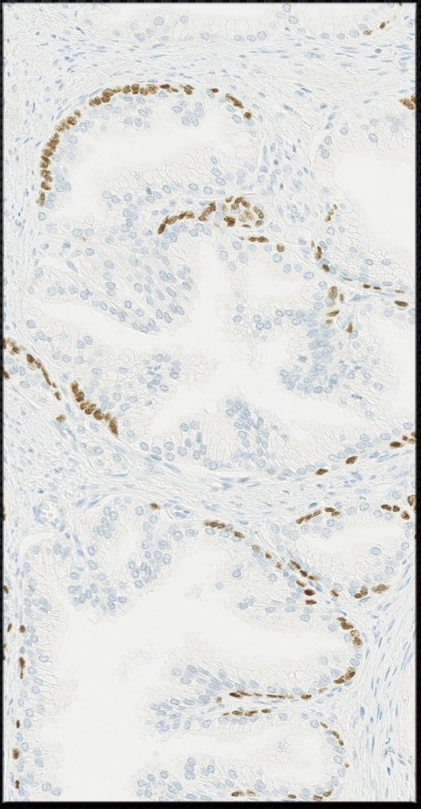


Controls / iCAPC

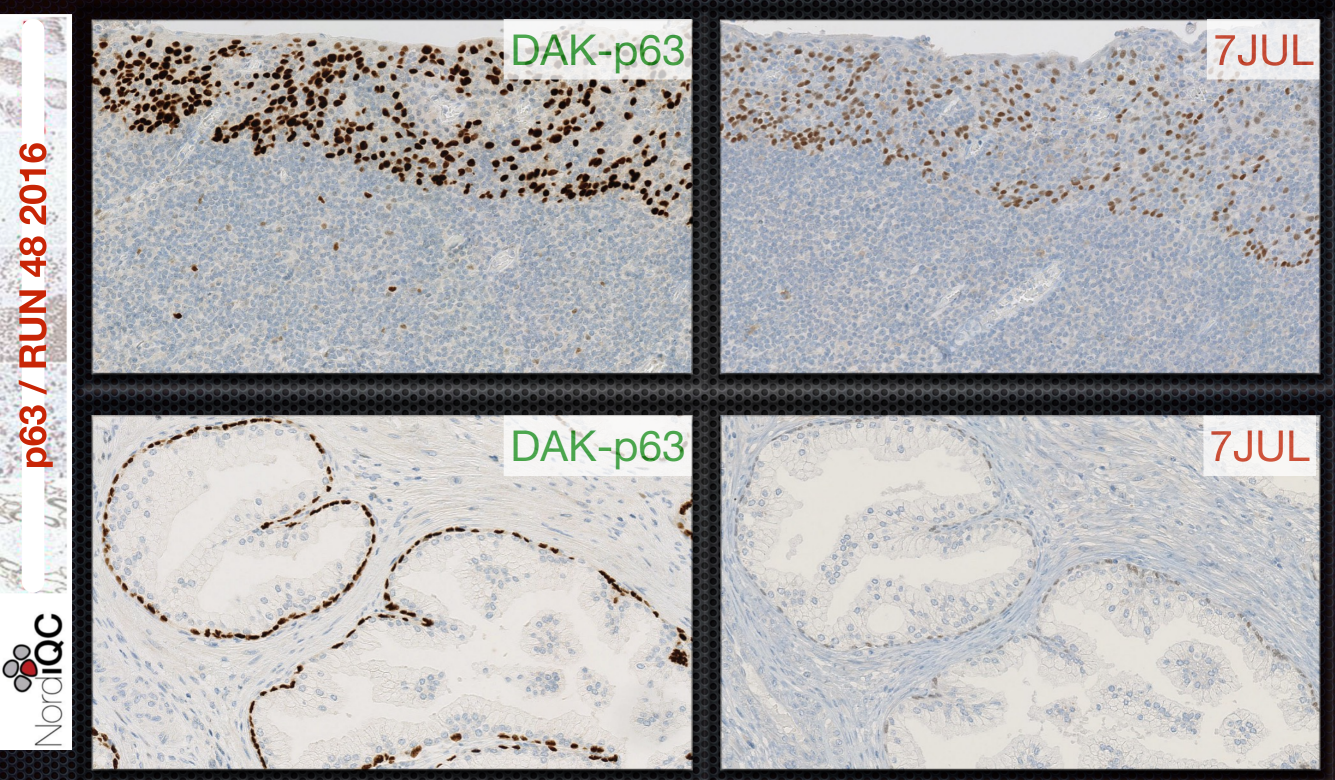
<u>Positive</u>: Tonsil or prostate.

- * Basal cells of prostate glands and squamous epithelial cells of tonsil must show a moderate to strong nuclear staining reaction.
- In the tonsil scattered lymphocytes must show a weak to moderate nuclear staining reaction.







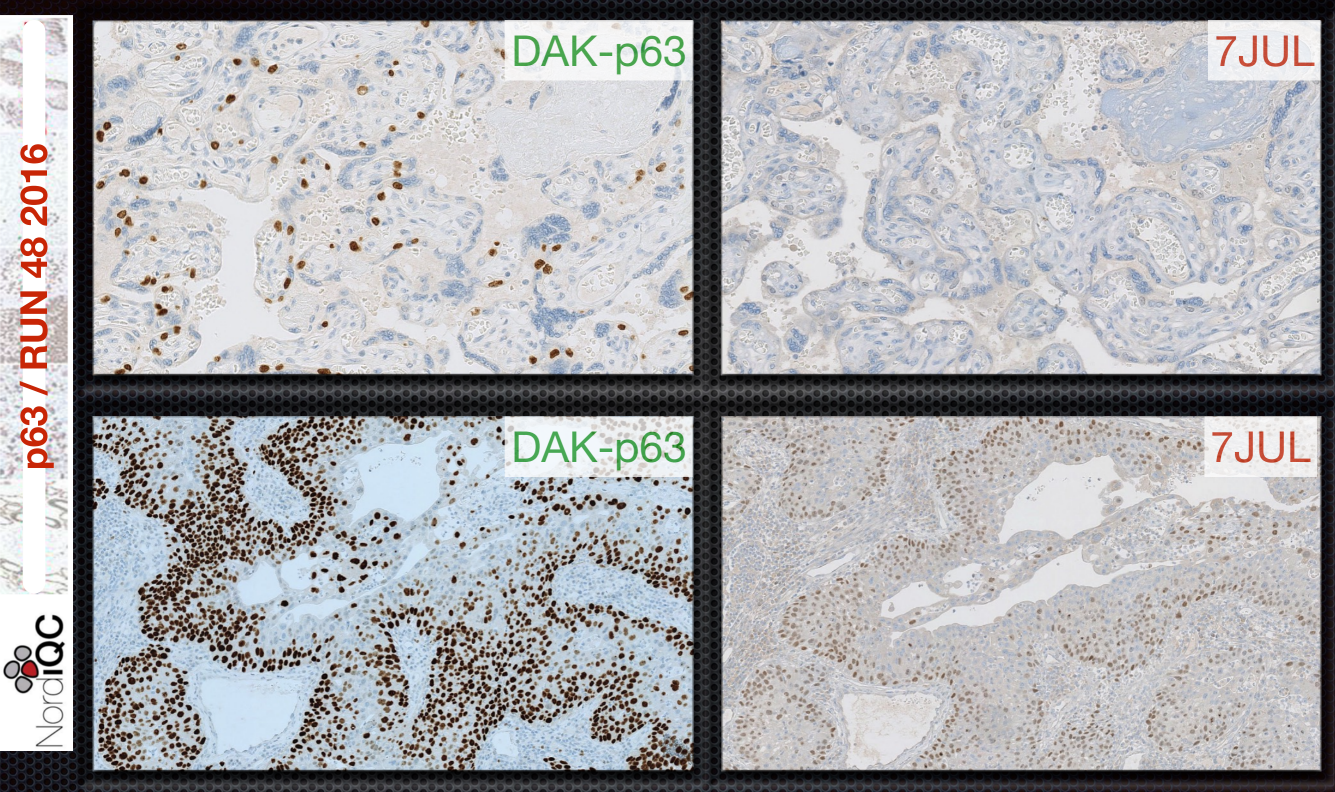


mAb clone DAK-p63 optimally calibrated

The use of a less successful primary Ab: 7JUL

				KOKOKOKOK	OKOKOKOKOK	KOKOKOKO	KOKOKOKOK	OKOKOKOKO
mAb clone DAK-p63	47	Dako	20	21	6	0	87%	91%
mAb clone 7JUL	12	Leica/Novocastra	0	1	3	8	8%	-

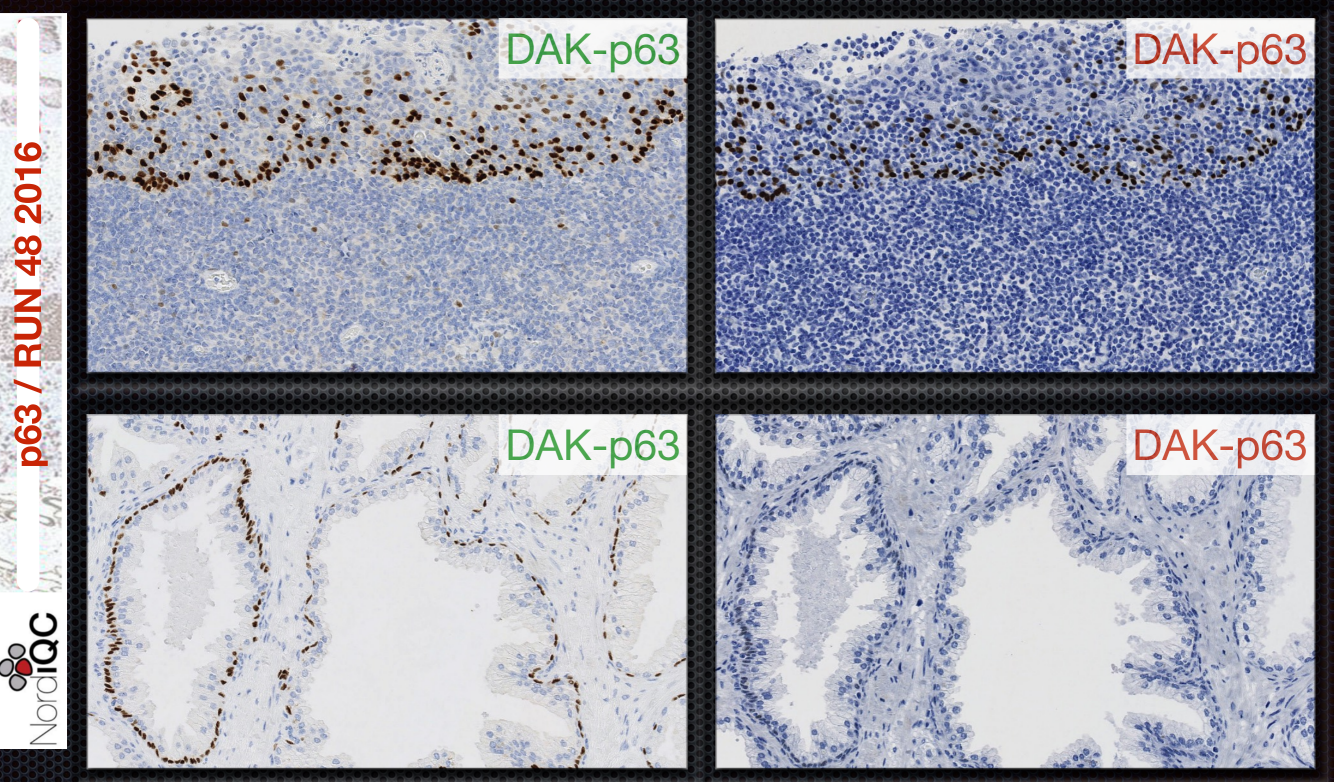




mAb clone DAK-p63 optimally calibrated

The use of a less successful primary Ab: 7JUL

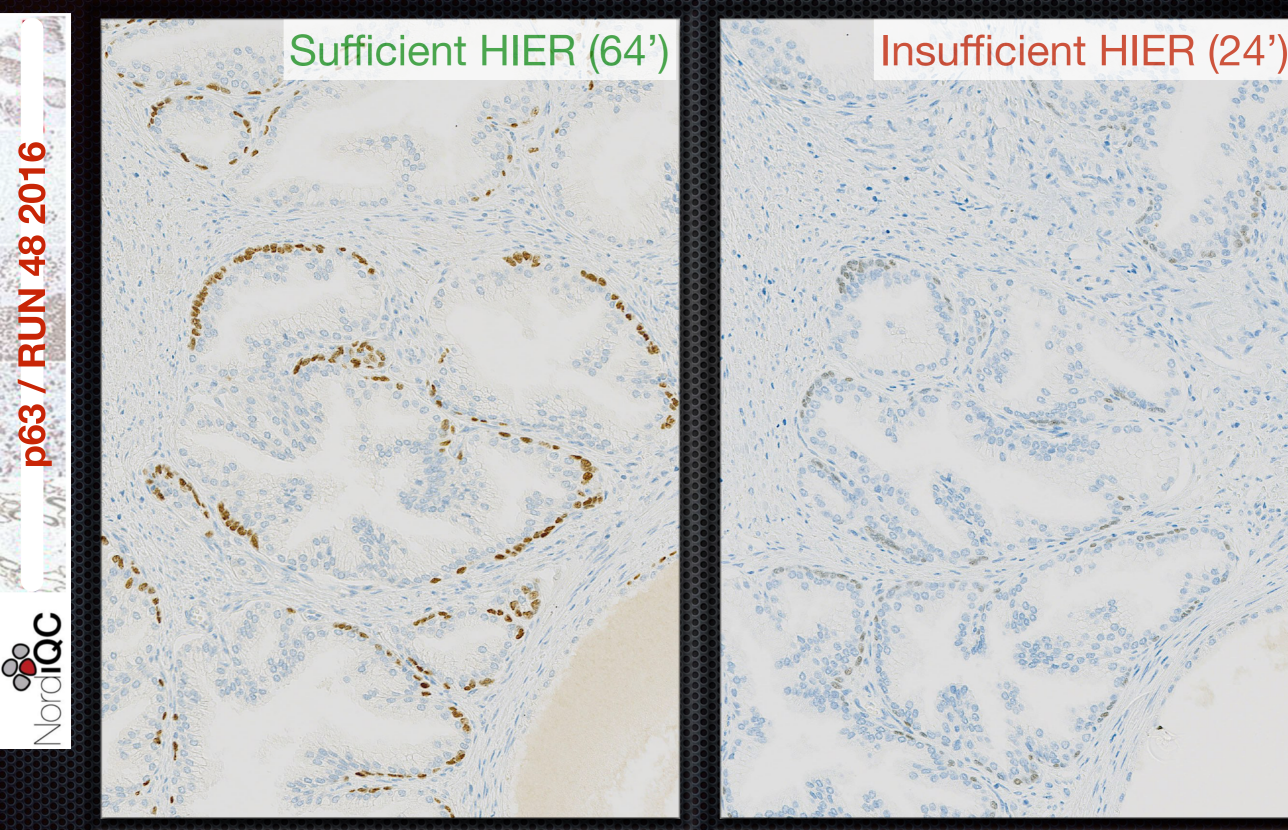




mAb clone DAK-p63 optimally calibrated in a sensitive 3-step polymer system.

Combination of the use of a less sensitive 2step polymer based detection system and strong Hematoxylin counter stain



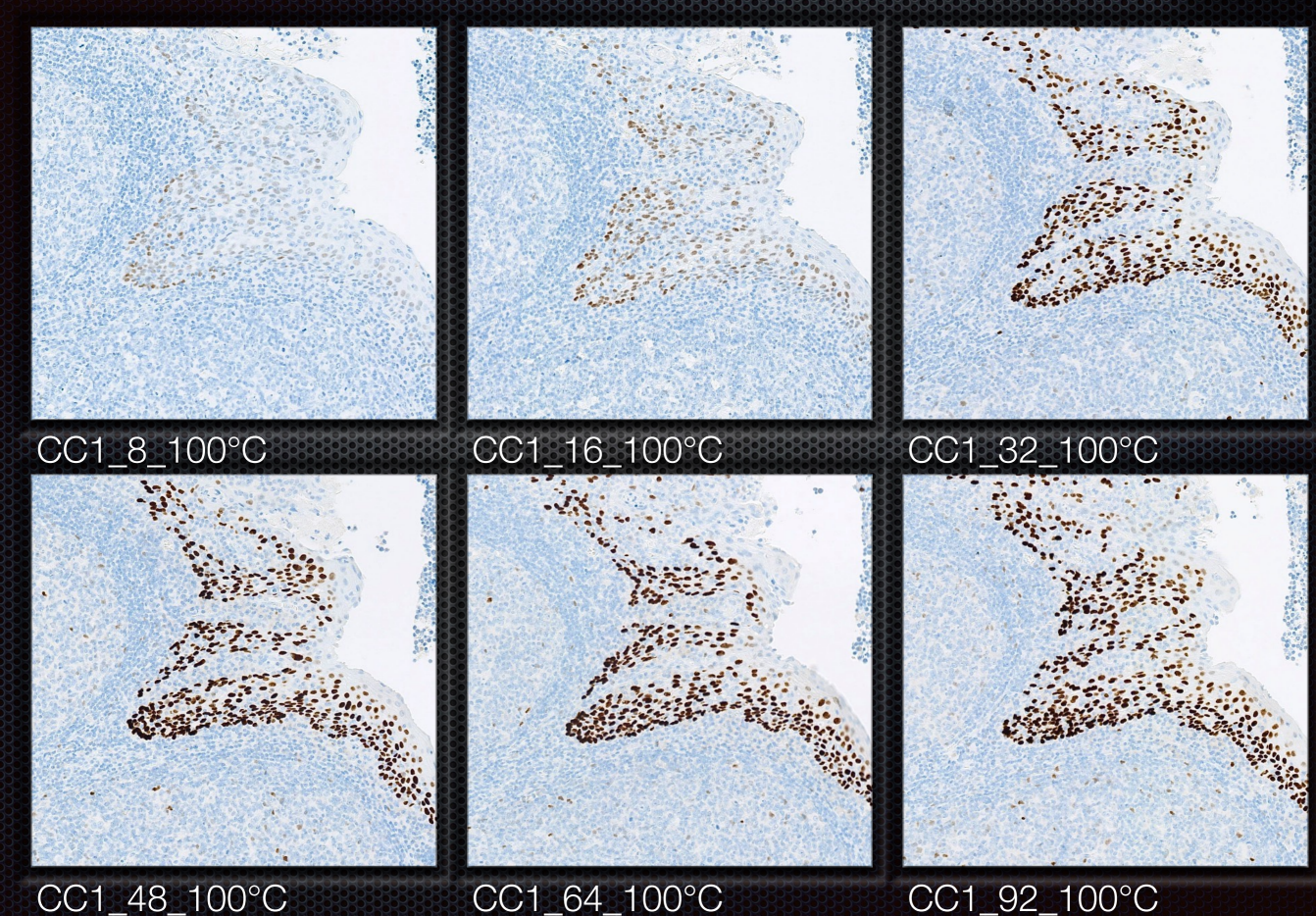


mAb clone 4A4 / CC1 64 min

mAb clone 4A4 / CC1 24 min

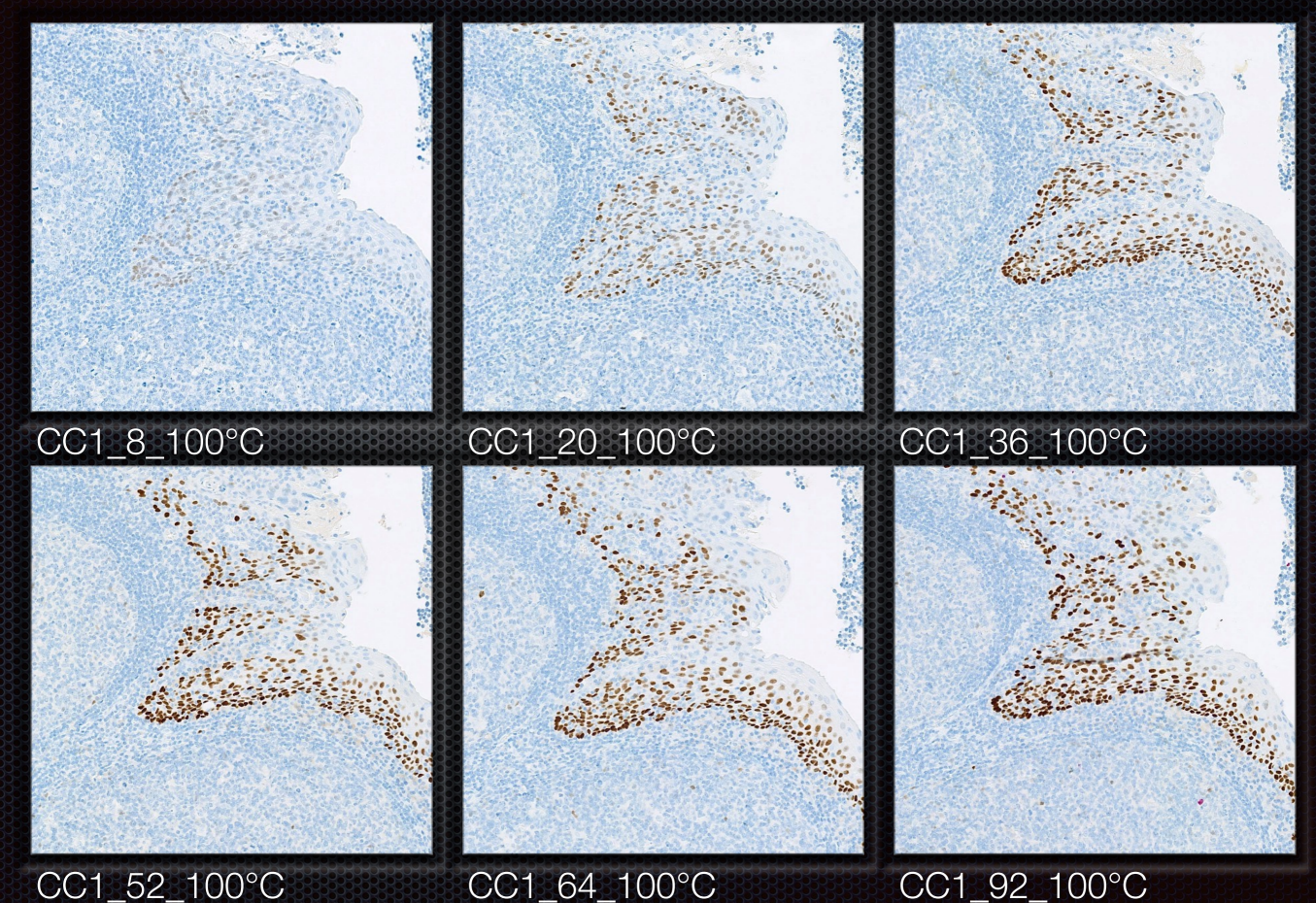
p63, 4A4 - OptiView (3-step) - Various HIER time Nordicc



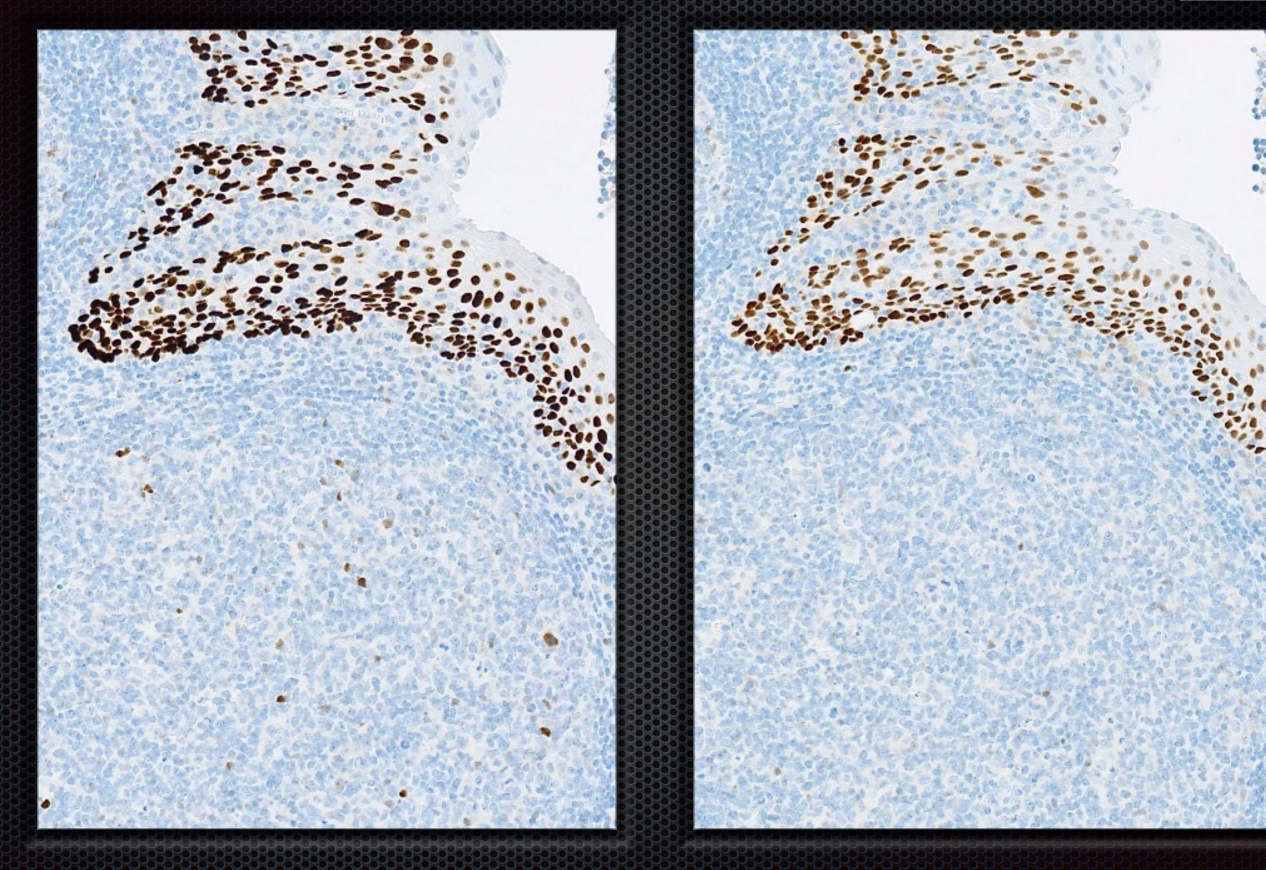


p63, 4A4 - UltraView (2-step) - Various HIER time Nordicc





p63, 4A4 OptiView (3-step) vs UltraView (2-step) Nordicc



OptiView - HIER CC1_48_100

UltraView - HIER CC1_52_100

Pass: 74 %

p40 / RUN 48 2016



200

150

100

50

0

Table 1. Antibodies and assessment marks for p40, run 48 Suff.¹ Suff. Concentrated antibodies Vendor Good Borderline Poor n Optimal OPS² 77 Biocare Zytomed 6 mAb clone BC28 2 2 Menarini 52 24 10 86% 89% 2 abcam 1 Nordic Biosite Immunologic 12 74% rmAb clone ZR8 Zeta Corporation 1 6 2 5 50% 67% 1 BioSB 1 pAb AC13030 8 Biocare 0 2 6 0 _ pAb **RP163 Diagnostic Biosystems** 5 0 1 3 56% 1 _ pAb **PC373** Calbiochem, Merck 0 0 3 1 4 _ pAb RBK054 3 Zytomed 0 0 1 2 _ pAb **PI049** DCS 0 1 0 1 0 _ 2015 2016 pAb **PP123** 0 0 0 1 1 Pathnsitu Ready-To-Use antibodies mAb clone BC28 100% 5 13 Biocare 8 0 0 100% API/IPI/AVI 3066 mAb clone BC28 5 39 19 15 0 87% 94% Ventana 790-4950 mAb clone BC28 0 0 1 0 1 Zytomed --**MSG097** mAb clone ZR8 Master Diagnostica 3 0 2 1 0 _ MAD-000686QD pAb API 3030 0 0 4 2 6 Biocare _ pAb RAB-066 0 1 Maixin 0 1 0 _ pAb **A00112** 0 0 1 0 Loxo GmbH 1 _ 78 17 Total 188 61 32 _

42%

Proportion

32%

17%

9%

74%



P40 / RUN 48 2016

Recommend- able clones	Retrieval	Titre	Detection	RTU	Detection
mAb BC28	HIER, High pH	1:20 - 1:100	3-step	Ventana	3-step
mAb ZR8	HIER, High pH	1:200	3-step + amp		

Table 3. Proportion of optimal results for p40 for the BC28 antibody as concentrate on the 4 main IHC systems*

Dako Autostainer Link / Classic		Dako OMNIS		Ventana BenchMark GX / XT / Ultra		Leica Bond III / Max	
TRS pH	TRS pH	TRS pH	TRS pH	CC1 pH	CC2 pH	ER2 pH	ER1 pH
9.0	6.1	9.0	6.1	8.5	6.0	9.0	6.0
7/20** (35%)	1/1	6/8 (75%)	1/1	30/42 (71%)	-	3/6 (50%)	0/1
	Autostain Clas TRS pH 9.0	Autostainer Link / ClassicTRS pHTRS pH9.06.17/20**1/1	Autostainer Link / Classic OM TRS pH TRS pH TRS pH 9.0 6.1 9.0 7/20** 1/1 6/8	Autostainer Link / Classic OMNIS TRS pH TRS pH TRS pH 9.0 6.1 9.0 6.1 7/20** 1/1 6/8 1/1	Autostainer Link / Classic OMNIS BenchMar TRS pH TRS pH TRS pH CC1 pH 9.0 6.1 9.0 6.1 8.5 7/20** 1/1 6/8 1/1 30/42	Autostainer Link / Classic OMNIS BenchMark GX / XT TRS pH TRS pH TRS pH CC1 pH CC2 pH 9.0 6.1 9.0 6.1 8.5 6.0 7/20** 1/1 6/8 1/1 30/42 -	Autostainer Link / ClassicOMNISBenchMark GX / XT / UltraBond II Bond IITRS pHTRS pHTRS pHTRS pHCC1 pHCC2 pHER2 pH9.06.19.06.18.56.09.07/20**1/16/81/130/423/6

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

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

p40 / RUN 44 2015



Controls / iCAPC

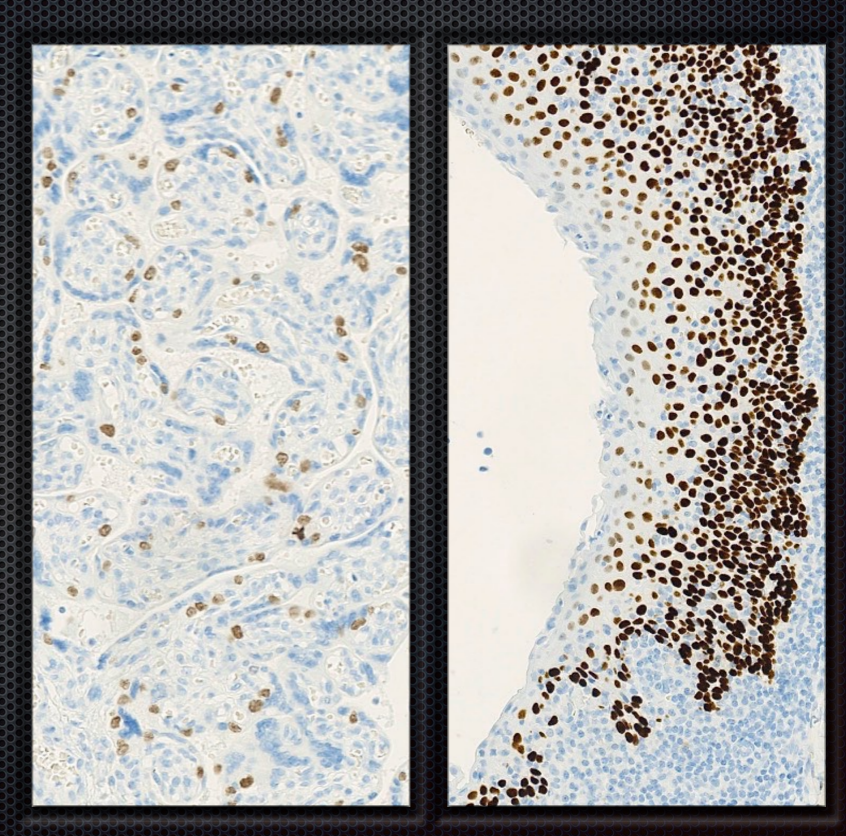
 Positive: Placenta (LLOD)
 * Cytotrophoblasts must show an at least weak to moderate, distinct nuclear staining reaction.

Positive: Tonsil

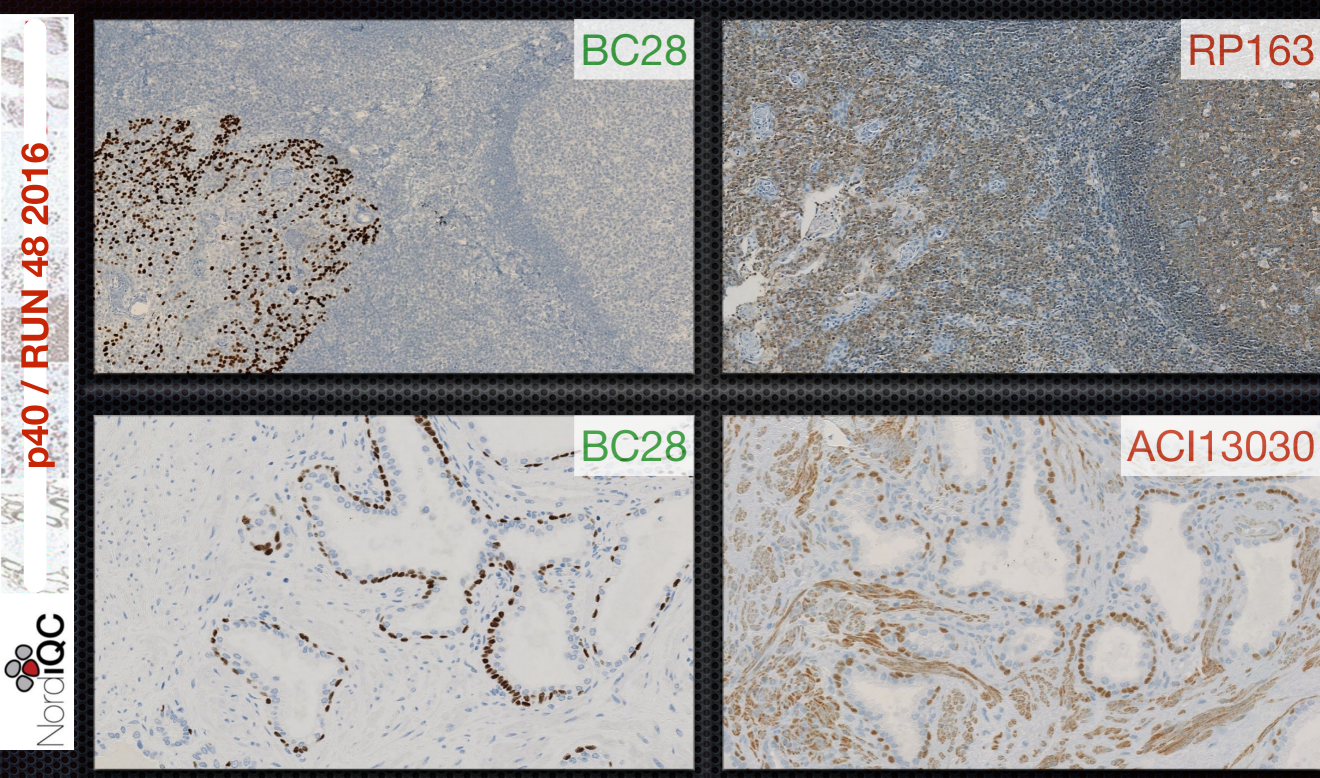
 Virtually all squamous epithelial cells must show a moderate to strong, distinct nuclear staining reaction.

Negative: Tonsil

 Lymphocyttes must be negative.



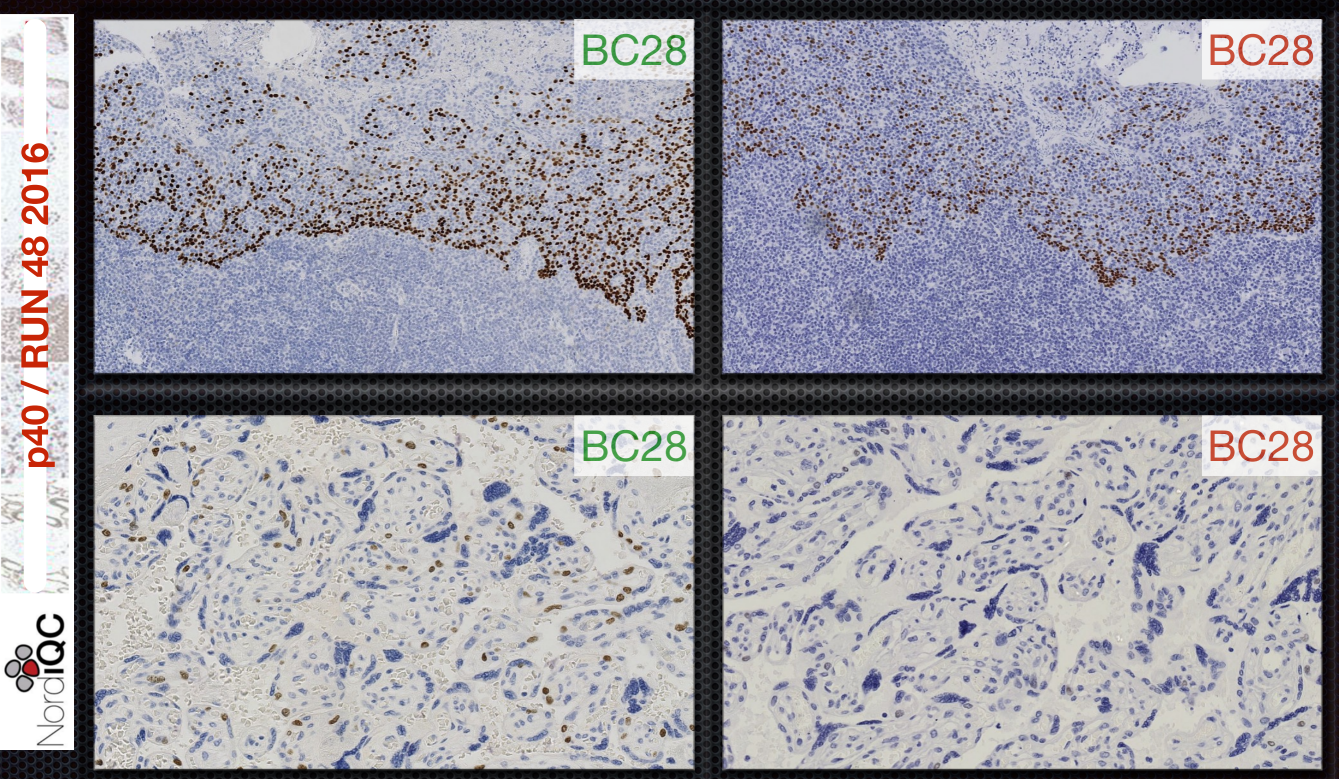




mAb clone BC28 optimally calibrated in a sensitive 3-step polymer system.

Poor signal to noise ratio using various pAb

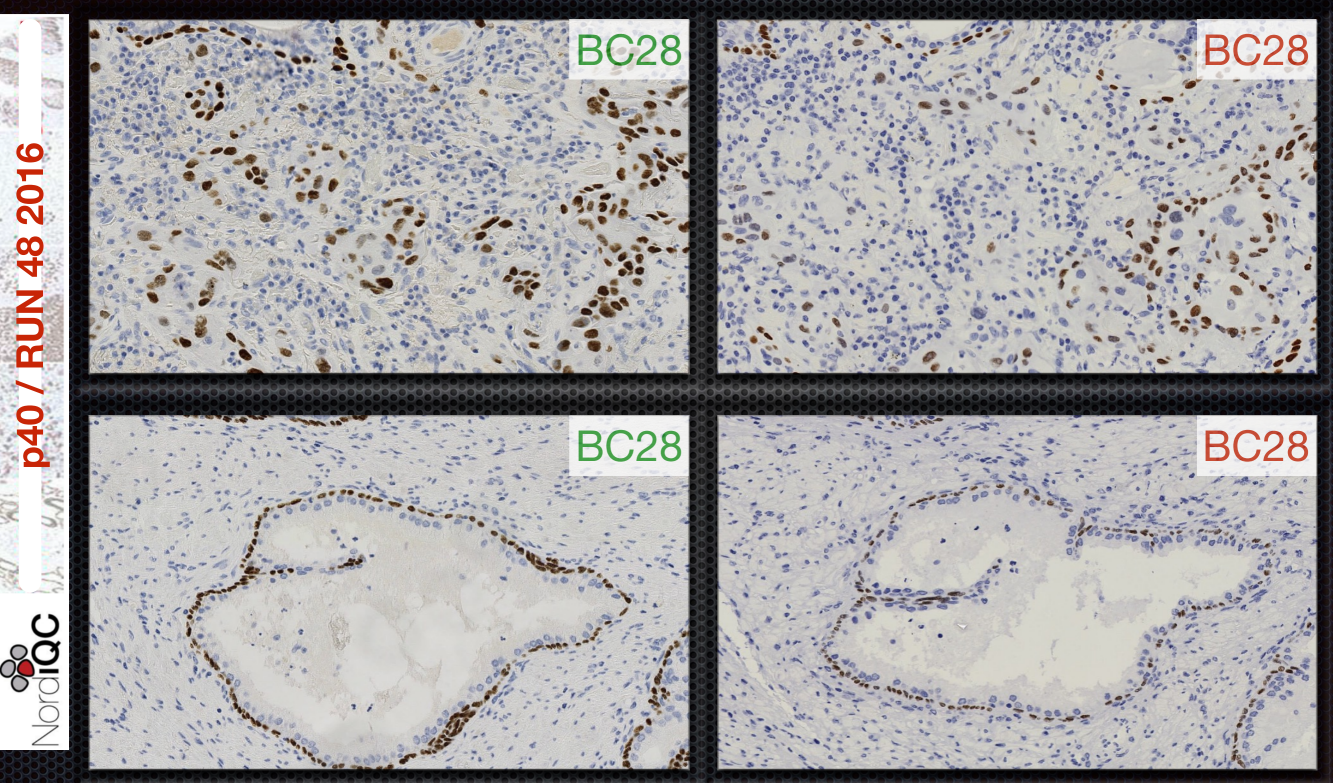




mAb clone BC28 optimally calibrated in a sensitive 3-step polymer system.

The use of a less sensitive 2-step polymer based detection system.



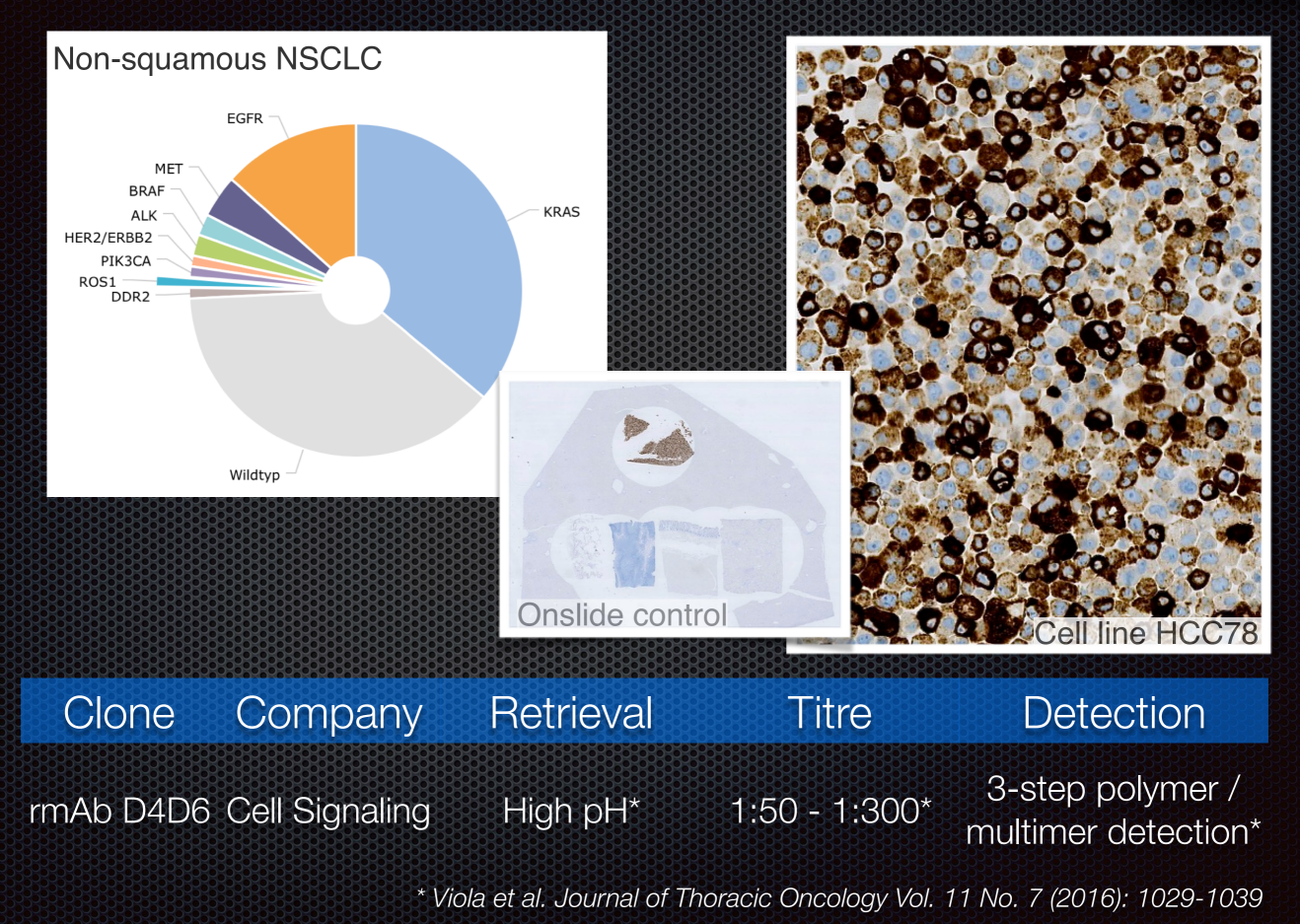


mAb clone BC28 optimally calibrated in a sensitive 3-step polymer system.

The use of a less sensitive 2-step polymer based detection system.

Driver mutations in lungcancer / ROS1









Thank you for your attention!