

Lung tumours

Optimization of antibodies, selection, protocols and controls

NQC Workshop 2018

Ole Nielsen, Dept. of Pathology Odense University Hospital



Lung markers in NordiQC assessments:

- lu-ALK (NQC in 2017)
- PD-L1 (NQC in 2018)
- 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 2015)
- CEA (NQC in 2016)
- Calretinin (NQC in 2018)
- CGA (NQC in 2018)
- Podoplanin (NQC in 2012)
- CD56 (NQC in 2013)



| Target | High scoring clones* | Low scoring clones* |
|------------|--|---|
| lu-ALK | rmAb: <mark>D5F3</mark> , mAb: <mark>OTI1A4</mark> | mAb: ALK1 mAb: 5A4 |
| PD-L1 | mAb: 22C3, rmAb: 28-8 and rmAb SP263 | |
| p63 | mAb: DAK-p63 and 4A4 | mAb: <mark>7JUL</mark> |
| p40 | mAb: BC28 and rmAb: ZR8 | Many pAbs |
| Napsin A | mAb: IP64 and MRQ-60 | pAb: <mark>760-4446</mark> and <mark>352A-7x</mark> |
| TTF1 | mAb: SPT24 and SP141 | mAb: <mark>8G7G3/</mark> 1 |
| SYP | mAb: 27G12, rmAb MRQ-40 and DAK-SYNAP | mAb: <mark>SY38</mark> |
| WT1 | mAb: WT49 and 6F-H2 | |
| 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: <mark>D2-40</mark> | mAb: D2-40 # |
| CD56 | rmAb: MRQ-42, mAb: CD564 and 123C3 | mAb: 123C3 # |

[#] Ventana platform § 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: DAK-p63

Producer: Dako

Product no. / lot no.: M7317 / 20032413

Diluent: Da Vinci Green

Dilution factor: 1:100

Incubation time / temperature: 32 min. / 36°C

Epitope retrieval, HIER

Device: On Board / On Machine

Buffer: Ventana Ultra CC1

Heating time at max. temp.: 56 min.

Maximum heating temp.: 100°C

Visualization system

Producer: Ventana

Product / no: OptiView DAB IHC Detection Kit / 760-700

Incubation time linker: 8 min.
Incubation time polymer: 8 min.
Incubation temperature: 36°C

Chromogen

Producer: Ventana

Product / no: OptiView DAB IHC Detection Kit / 760-700

Incubation time / temperature: 8 min. / 22°C

Enhancement: 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 Salate And Andrew 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: <mark>D5F3</mark> , mAb: <mark>OTI1A4</mark> | mAb: ALK1 mAb: 5A4 |
| PD-L1 | mAb: 22C3, rmAb: 28-8 and rmAb SP263 | |
| p63 | mAb: DAK-p63 and 4A4 | mAb: <mark>7JUL</mark> |
| p40 | mAb: BC28 and rmAb: ZR8 | Many pAbs |
| Napsin A | mAb: IP64 and MRQ-60 | pAb: <mark>760-4446</mark> and <mark>352A-7x</mark> |
| TTF1 | mAb: SPT24 and SP141 | mAb: <mark>8G7G3/</mark> 1 |
| SYP | mAb: 27G12, rmAb MRQ-40 and DAK-SYNAP | mAb: <mark>SY38</mark> |
| WT1 | mAb: WT49 and 6F-H2 | |
| 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: <mark>D2-40</mark> | mAb: D2-40 # |
| CD56 | rmAb: MRQ-42, mAb: CD564 and 123C3 | mAb: 123C3 # |

[#] Ventana platform § Products discontinued

^{*} on the basis of the assessments in NordiQC

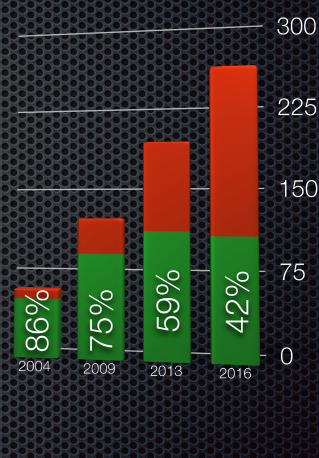




CEA / RUN 47 2016

Pass: 42 %

| Table 1 Antibodies | and a | ssessment marks for CE | Λ rup 4 | 00000000000000000000000000000000000000 | 0,000,000,000 | | 00000000 | 20202020 |
|---|-----------------------|---|---------|--|---------------|------|--------------------|---------------------------|
| 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 | 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 | - | - |
| mAb PARLAM 4 | 1 | Monosan | 0 | 0 | 1 | 0 | - | - |
| mAb BS33 | 1 | Nordic Biosite | 0 | 0 | 1 | 0 | | |
| Ready-To-Use Antibodies | | | | | | | | |
| 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 | 0 | 0 | 1 | 1 | - | - |
| mAb clone COL-1 PM058 | 1 | Biocare | 0 | 0 | 1 | 0 | - | - |
| mAb clone COL-1 Kit-0008 | 1 | Maixin | 1 | 0 | 0 | 0 | - | - |
| mAb clone II-7 IR/IS622/GA622 | 47 | Dako/Agilent | 0 | 6 | 40 | 1 | 13% | - |
| mAb clone II-7 PA0004 | 12 | Leica | 0 | 5 | 6 | 1 | 42% | - |
| mAb clone TF3H8-1 760-2507 | 13 | Ventana/Roche | 0 | 0 | 0 | 13 | 0% | _ |
| Total | 255 | | 43 | 65 | 122 | 25 | - | |
| Proportion | | | 17% | 25% | 48% | 10% | 42% | |



mAb clone TF3H8-1 cross reacts with BGP and NCA

mAb clone II-7 is difficult to optimise





CEA / RUN 47 2016

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 47 2016

Controls / iCAPC

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

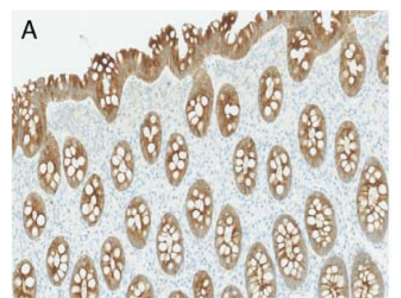
Emina E. Torlakovic, MD, PhD,*† Søren Nielsen, HT, CT,‡§ Glenn Francis, MBBS, FRCPA, MBA, FFSc (RCPA), ||¶ John Garratt, RT,†** Blake Gilks, MD, FRCPC,†††

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‡§





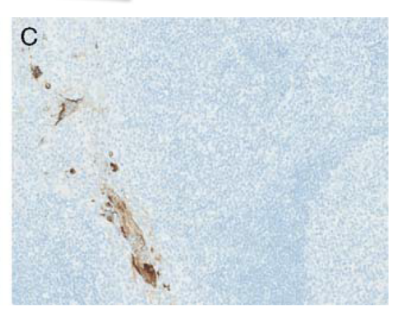


FIGURE 9. mCEA iCAP. A, A moderate to strong staining reaction must be seen in the brush border of the surface epithelial cells. Virtually all epithelial cells must show a weak to moderate cytoplasmic staining reaction (LLOD). If overexpression of the CEA is desirable target for detection in adenocarcinoma, 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 | 1 | 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)



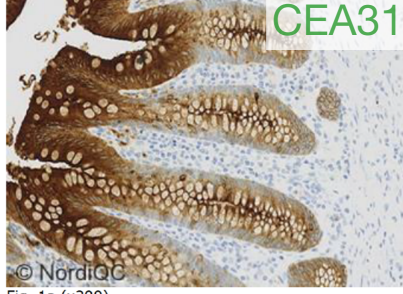


Fig. 1a (x200)

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.

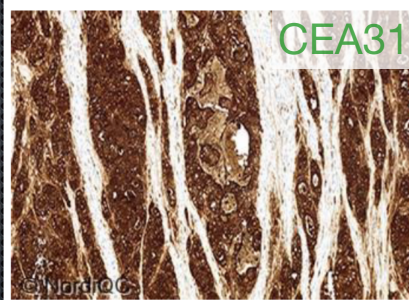


Fig. 2a (x200)

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.

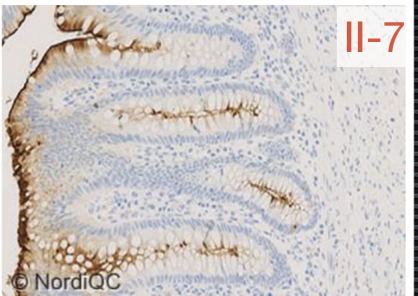


Fig. 1b (x200)

Insufficient CEA 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.

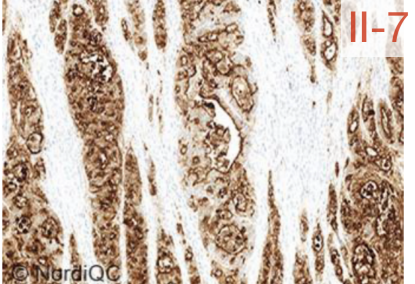


Fig. 2b (x200)

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.

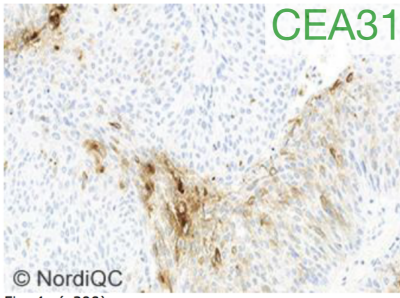


Fig. 4a (x200)

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.

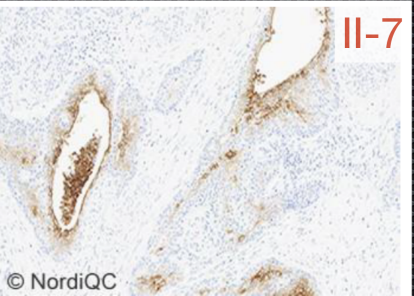


Fig. 3b (x200)
Insufficient CEA staining of the urothelial carcinoma, tissue core no. 4, using same protocol as in Figs. 1b and 2b – same field as in Fig. 3a. The proportion and intensity of the neoplastic cells demonstrated is significantly reduced compared to the level expected and obtained in Fig. 3a.

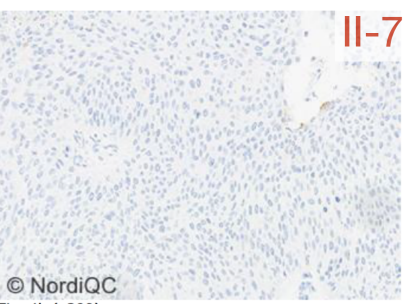


Fig. 4b (x200)

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



CEA / RUN 47 2016

C Madioc

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.

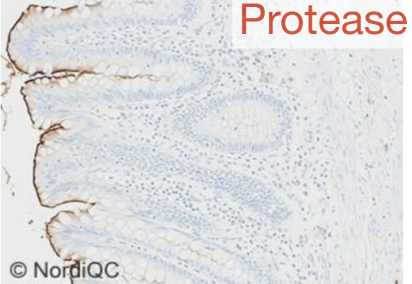


Fig. 5b (x200)

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.

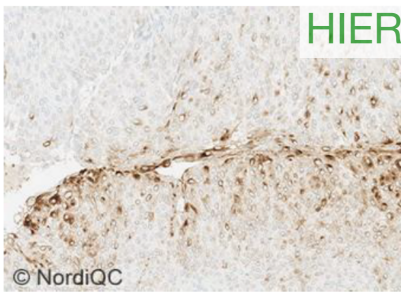


Fig. 6a (x200)

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.

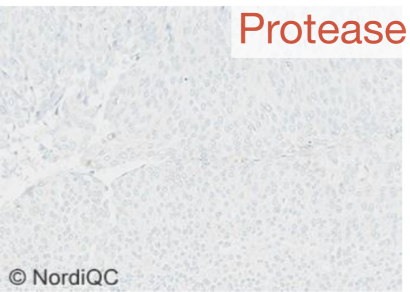


Fig. 6b (x200)

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



Inappropriate antibody - NCA and BGP cross reaction



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.

© NordiQČ

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.

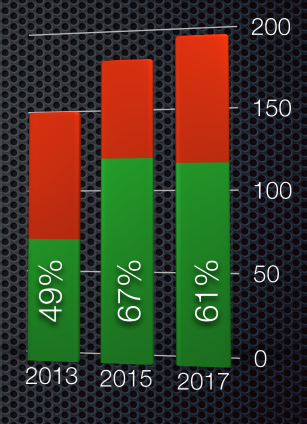




lu-ALK / RUN 51 2017

Pass: 61 %

| Table 1. Antibodies and assessment marks for lu-ALK, run 51 | | | | | | | | |
|---|------------------------|---|---------|------|------------|------|--------|---------------------------|
| Concentrated antibodies | n | Vendor | Optimal | Good | Borderline | Poor | Suff.1 | Suff. OPS ² |
| mAb clone 5A4 | 43 1 1 1 1 | Leica/Novocastra Abcam Biocare Monosan ThermoFisher | 1 | 15 | 24 | 7 | 34% | 22% |
| mAb clone ALK1 | 2 1 | Dako Cell Marque | 0 | 0 | 0 | 3 | - | - |
| rmAb clone D5F3 | 23 | Cell Signaling | 6 | 12 | 3 | 2 | 78% | 94% |
| mAb clone OTI1A4 | 13 | ORIGENE | 10 | 3 | 0 | 0 | 100% | 100% |
| Ready-To-Use antibodies | | | | | | | | |
| mAb clone 5A4 PA0306 | 6 | Leica/Novocatra | 0 | 0 | 6 | 0 | - | - |
| mAb clone 5A4 MAB-0281 | 1 | Maixin | 0 | 0 | 1 | 0 | - | - |
| mAb 5A4 MAD-001720QD | 1 | Master Diagnostica | 0 | 0 | 1 | 0 | - | - |
| mAb clone 5A4 MS-1104-R7 | 1 | ThermoFisher | 0 | 1 | 0 | 0 | - | - |
| mAb ALK1 IR641 | 9 | Dako | 0 | 0 | 1 | 8 | - | - |
| mAb clone ALK1 GA641 | 4 | Dako | 0 | 0 | 0 | 4 | - | - |
| mAb clone ALK1 790/800-2918 | 7 | Ventana | 0 | 0 | 2 | 5 | - | - |
| rmAb clone SP8 AN770 | 1 | BioGenex | 0 | 0 | 0 | 1 | - | - |
| rmAb clone D5F3 790-4796 | 70 | Ventana | 53 | 12 | 4 | 1 | 93% | 100% |
| rmAb clone D5F3 790-4796 ³ | 2 | Ventana | 1 | 0 | 1 | 0 | - | - |
| mAb clone OTI1A4 8344-C010 | 1 | Sakura Finetek | 1 | 0 | 0 | 0 | - | - |
| Total | 189 | | 72 | 43 | 43 | 31 | - | |
| Proportion | | | 38% | 23% | 23% | 16% | 61% | |



mAb clone ALK1 is not "Fit for purpose"

mAb clone 5A5 is difficult to optimise





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*

| tile 4 illalli Int | ne 4 main the systems. | | | | | | | | |
|---------------------------|---------------------------------|---------------|---------------|---------------|---------------------------------|---------------|-------------------------|---------------|--|
| Concentrated antibodies | Dako Autostainer Link / Classic | | Dako Omnis | | Ventana BenchMark XT / Ultra | | 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)





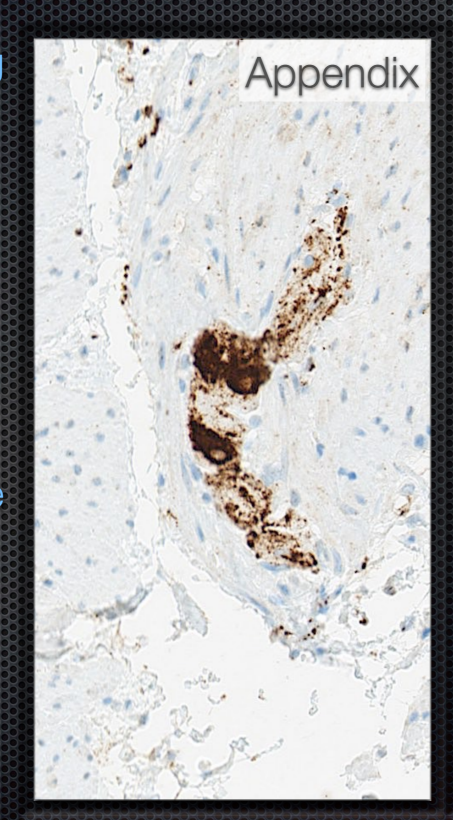
lu-ALK / RUN 51 2017

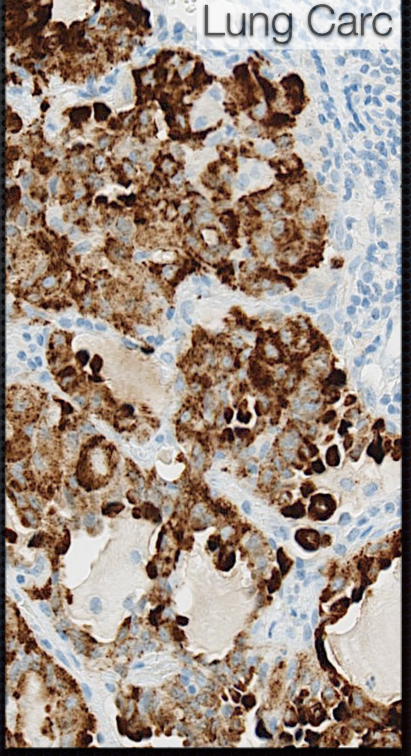
Controls / iCAPC

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

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

Negative: Lung cancer without ALK rearrangements





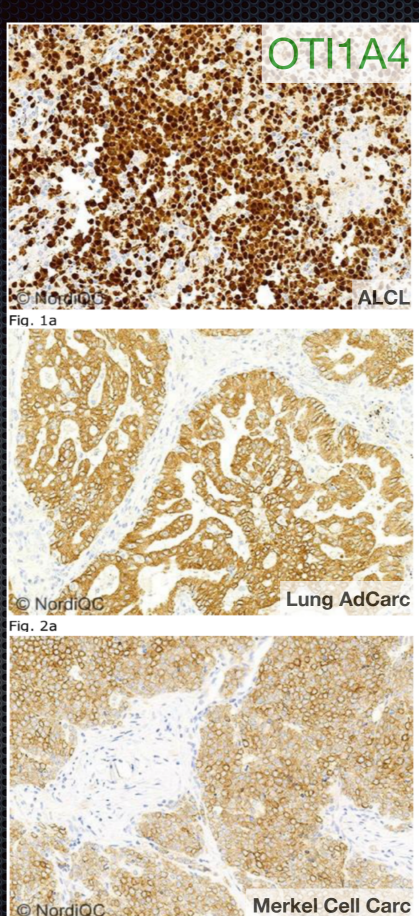
lu-ALK / RUN 51 2017

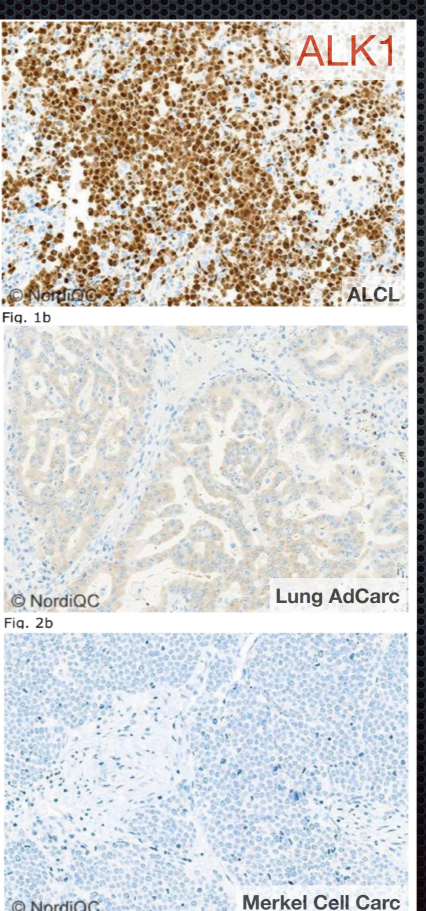
Lung tumours: Antibodies, protocols and controls

Fig. 3b









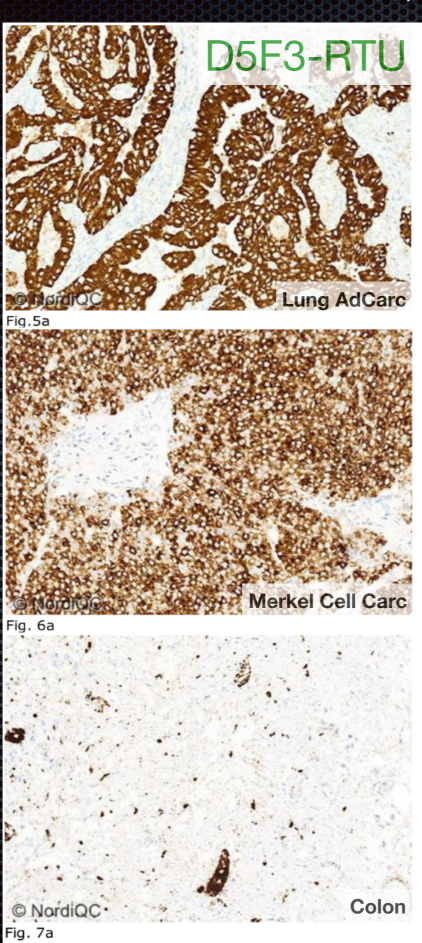
Less
successful
primary
antibody:
mAb clone
ALK1

u-ALK / RUN 51 2017

Lung tumours: Antibodies, protocols and controls







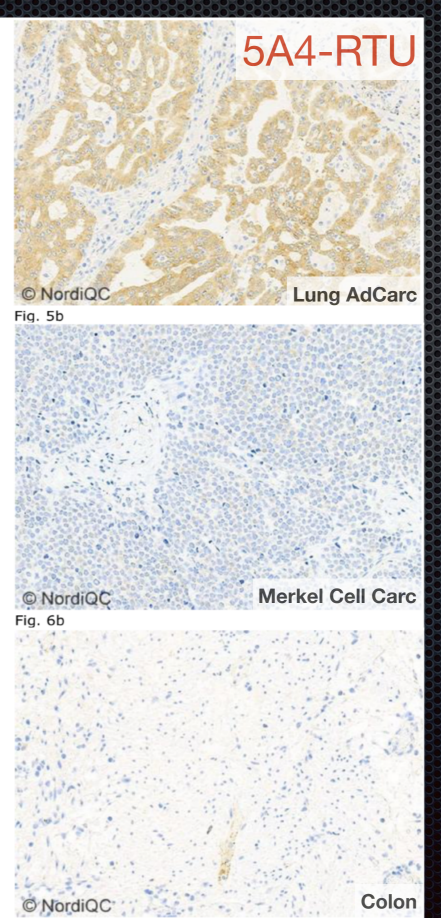


Fig. 7b

Less
successful
RTU system
based on
mAb clone
5A4





p63 / RUN 48 2016

Pass: 82 %

| 1 VOI GIGO | 20 | A IN HER CALL CALLS TO BRIDGE WAS | 200000000000000000000000000000000000000 | | 6 J w | (1000) | | |
|--|--|---|---|------|------------|--------|--------------------|---------------------------|
| Table 1. Antibodies ar | nd a | ssessment marks for p63 | , run 48 | | | | | |
| Concentrated antibodies | n | Vendor | Optimal | Good | Borderline | Poor | Suff. ¹ | Suff. OPS ² |
| mAb clone 4A4 | 26 4 3 2 2 1 1 1 1 | BioCare Medical ImmunoLogic Dako Zeta Corporation Thermo Scientific Zytomed Systems BioGenex Diagnostic BioSystems Klinipath Minarini Nordic Biosite Santa Cruz | 13 | 20 | 11 | 2 | 72% | 76% |
| 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 SFI-6 | 2 | DCS Immunoline | 0 | 0 | 2 | 0 | - | - |
| rmAb clone BSR6 | 1 | Nordic Biosite | 0 | 0 | 1 | 0 | - | - |
| rmAb clone DBR16.1 | 1 | Diagnostic Biosystems | 1 | 0 | 0 | 0 | | |
| rmAb clone EPR5701 | 1 | Epitomics | 0 | 0 | 1 | 0 | - | - |
| Unknown 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 PM163 | 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 | 0 | 0 | - | - |
| Total | 274 | | 120 | 104 | 35 | 15 | - | |
| Proportion | | | 44 % | 38 % | 13 % | 5 % | 82 % | |







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 | (0 | Venta | na | Leica | | |
|----------------------|--------------|------------|----------------------|------------|------------|------------|--|
| antibodies | Autostaine | r / Omnis | BenchMark XT / Ultra | | Bond II | I / 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 applied as listed above, HIER buffers and detection kits used as recommended by the vendors of the respective platforms.

^{** (}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* | Laboratory modified protocol 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.





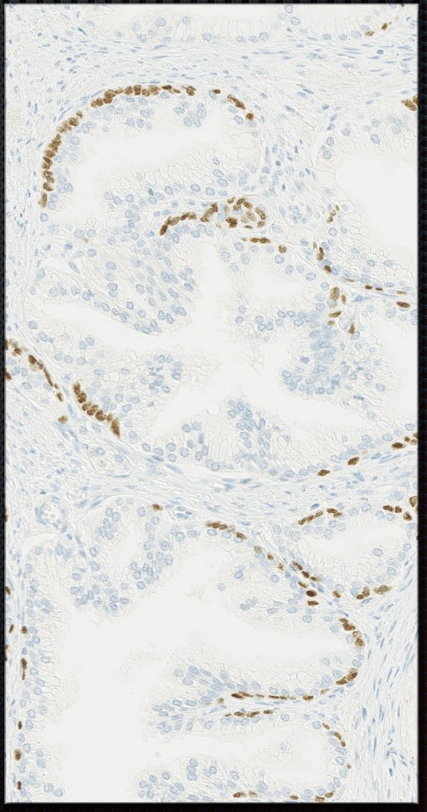
p63 / RUN 41 2014

Controls / iCAPC

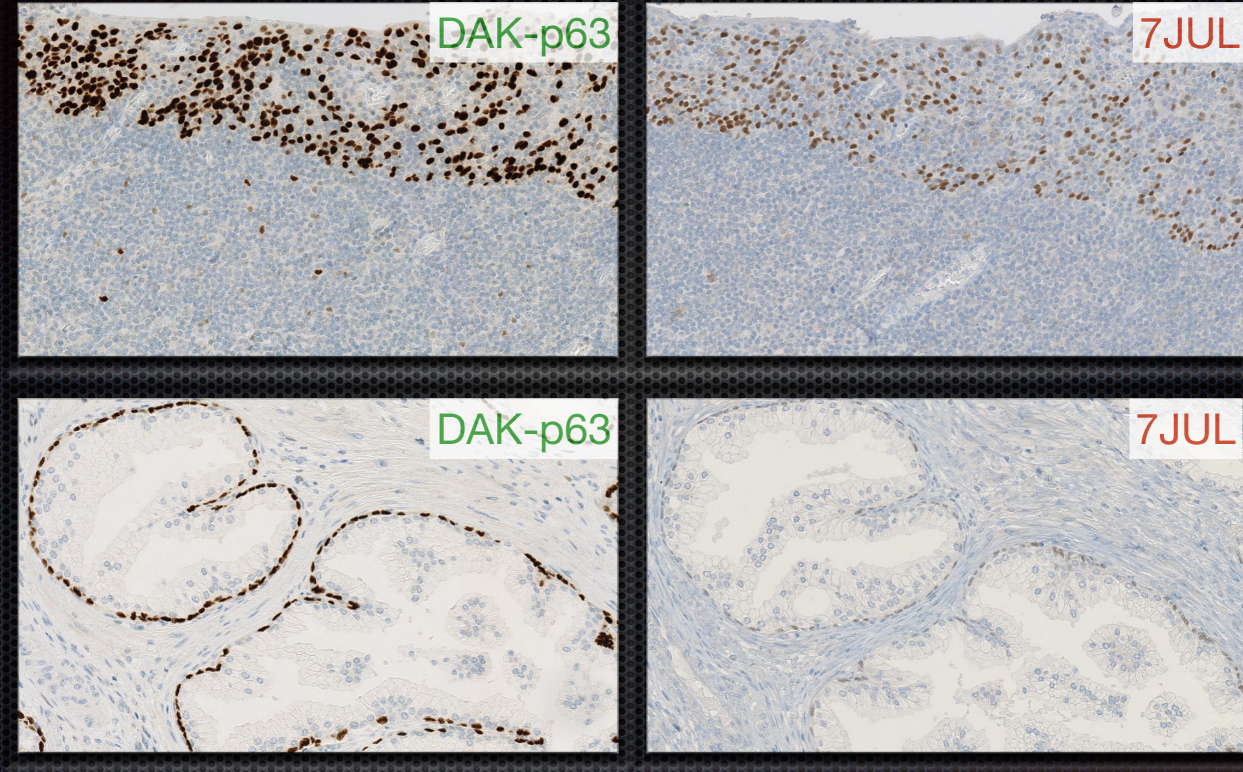
Positive: Tonsil or prostate.

- * 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.







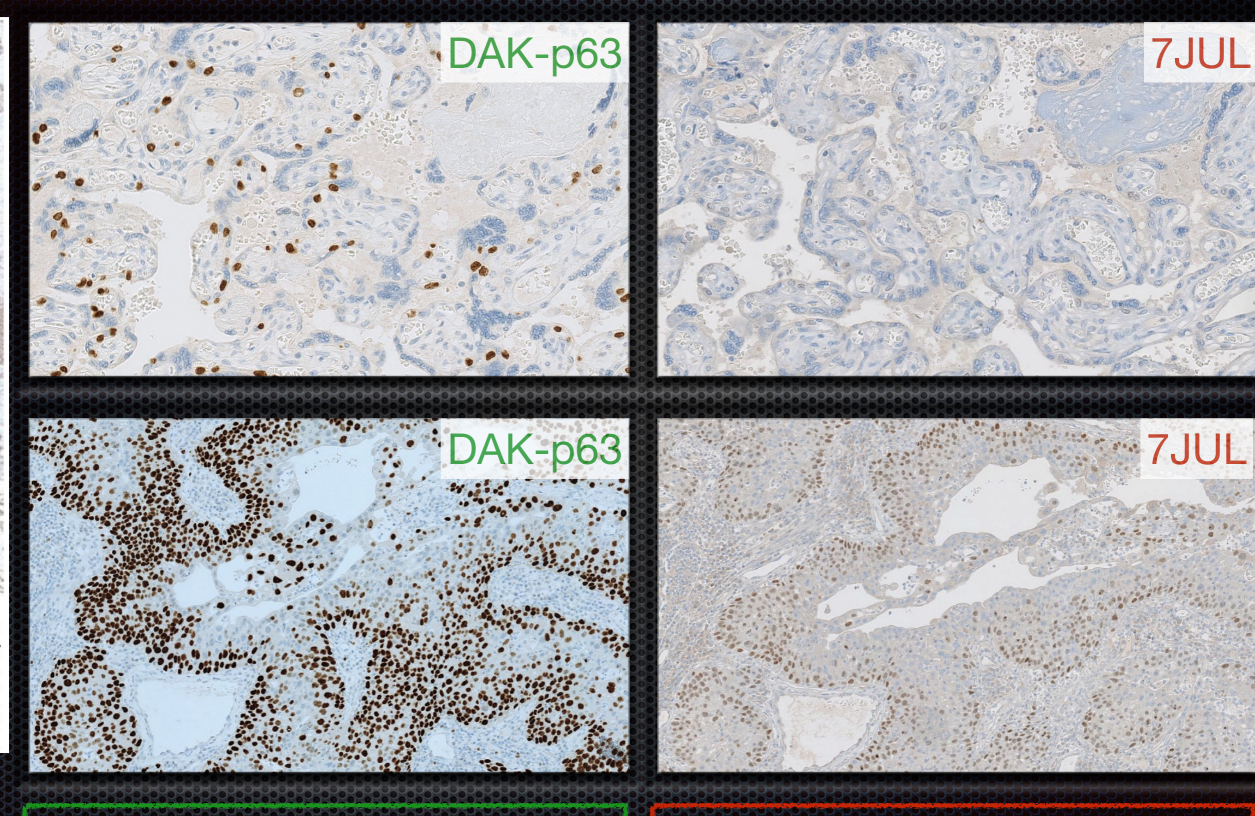


mAb clone DAK-p63 optimally calibrated

The use of a less successful primary Ab: 7JUL

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

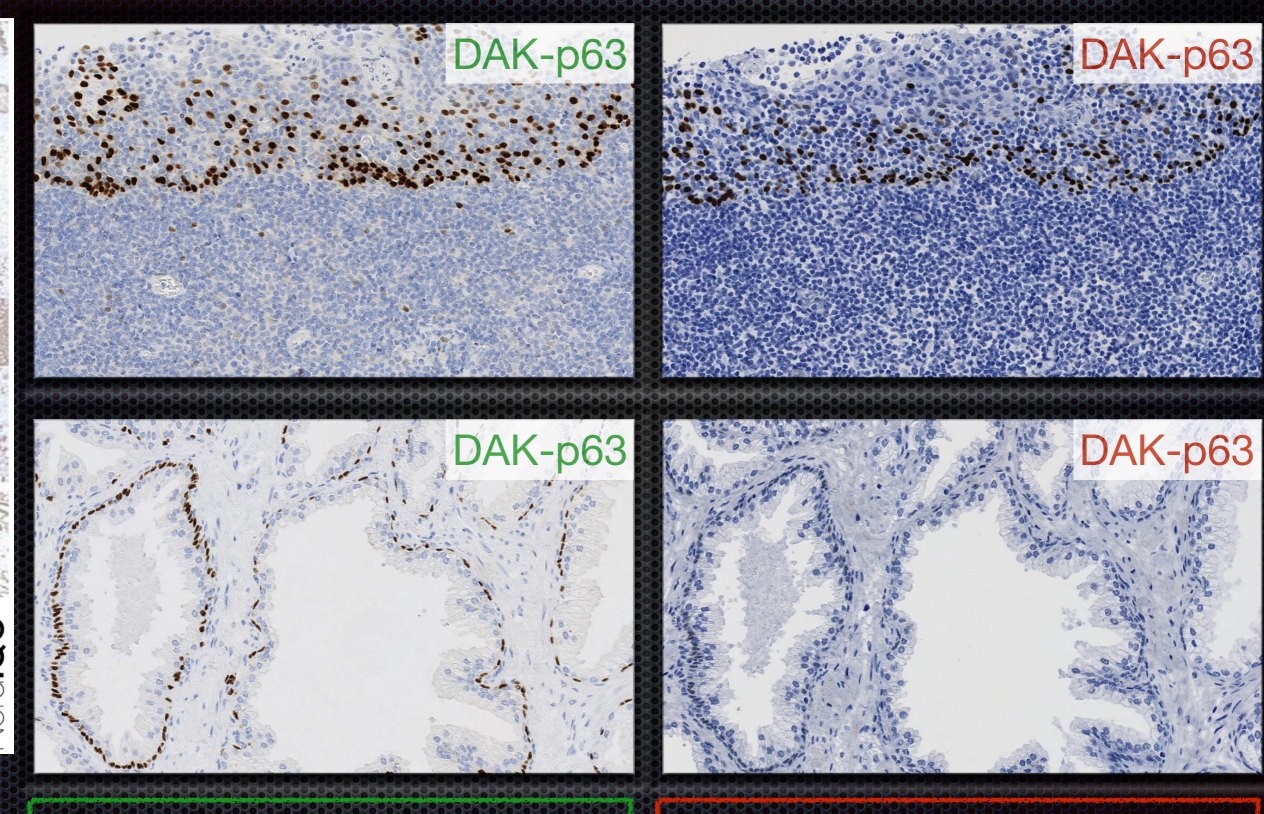




mAb clone DAK-p63 optimally calibrated

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

Sufficient HIER (64')



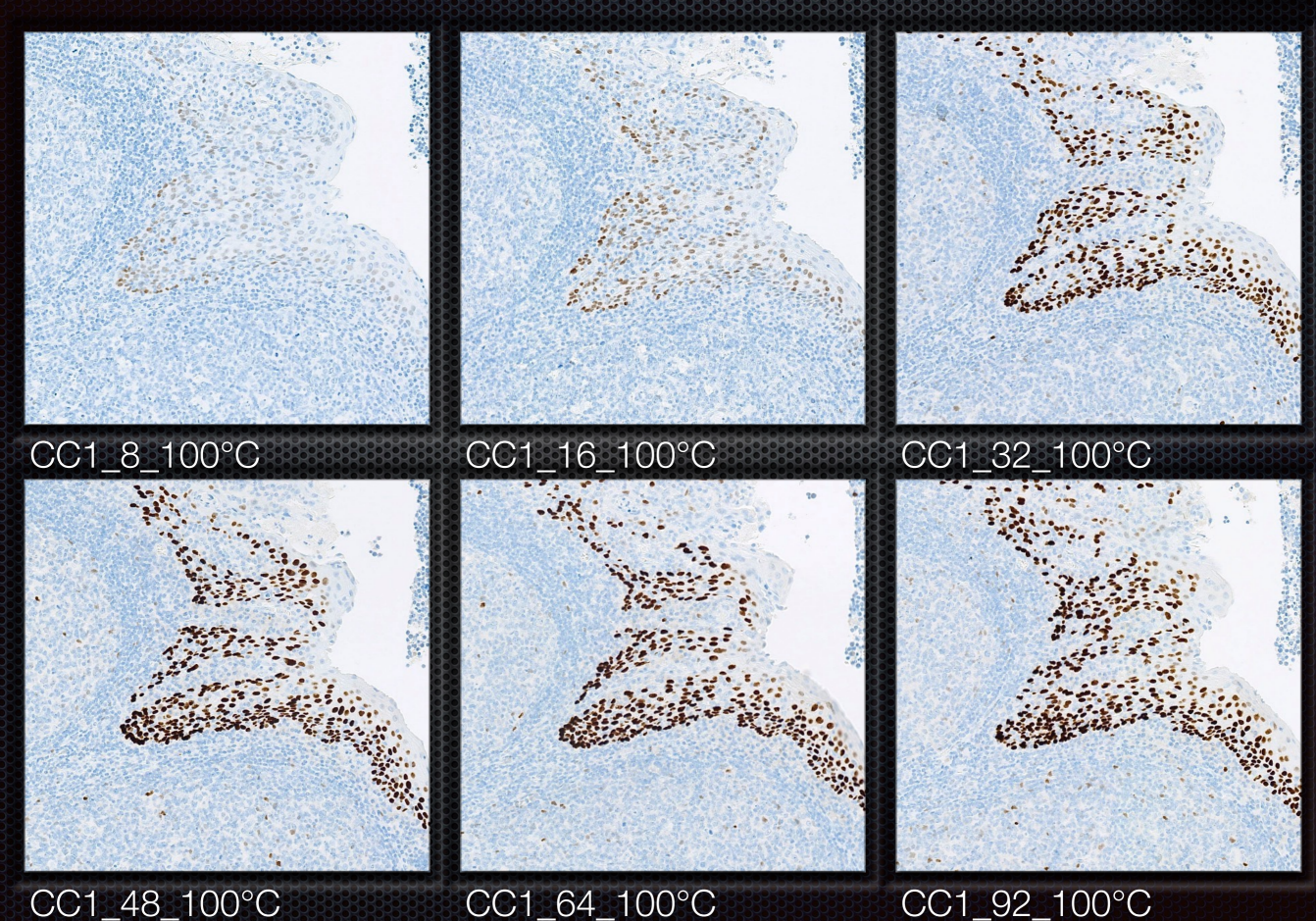


mAb clone 4A4 / CC1 64 min

mAb clone 4A4 / CC1 24 min

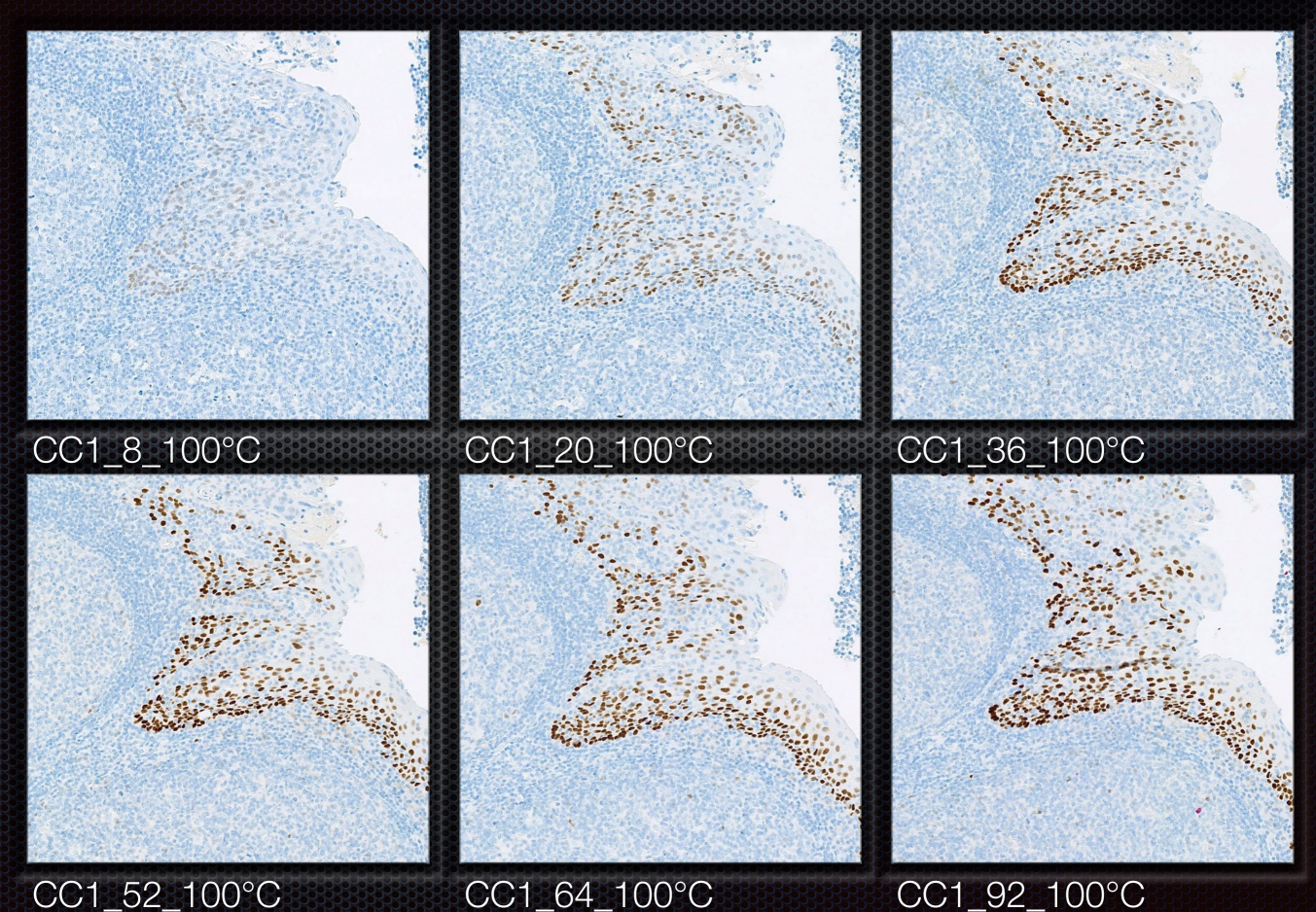
p63, 4A4 - OptiView (3-step) - Various HIER time Nordicc





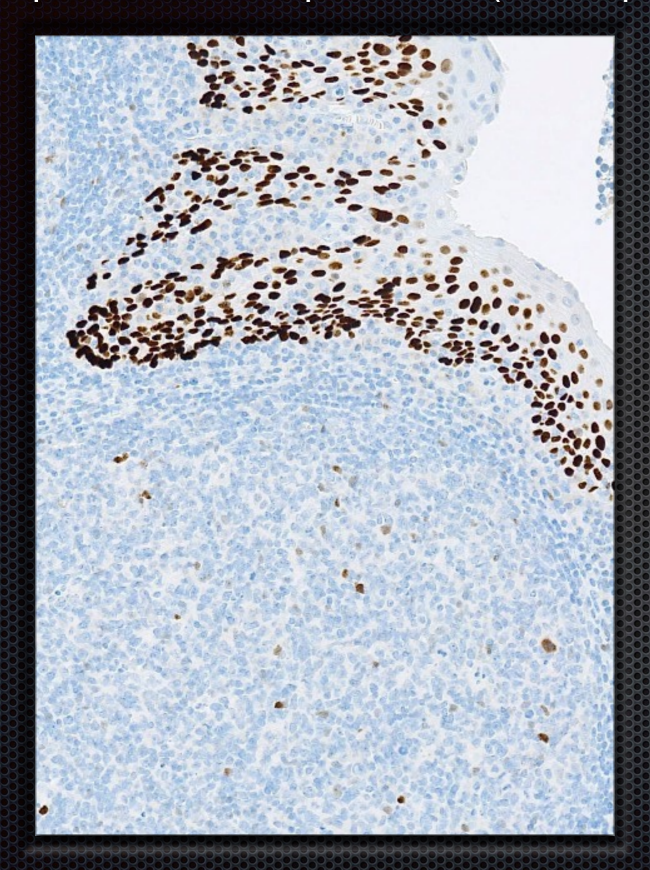
p63, 4A4 - UltraView (2-step) - Various HIER time Nordicc

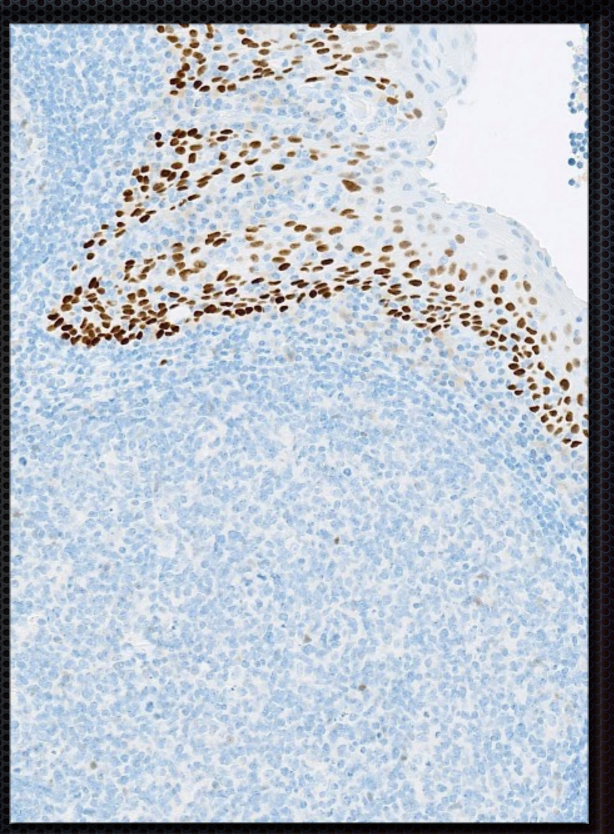




p63, 4A4 OptiView (3-step) vs UltraView (2-step) Nordicc







OptiView - HIER CC1_48_100

UltraView - HIER CC1_52_100

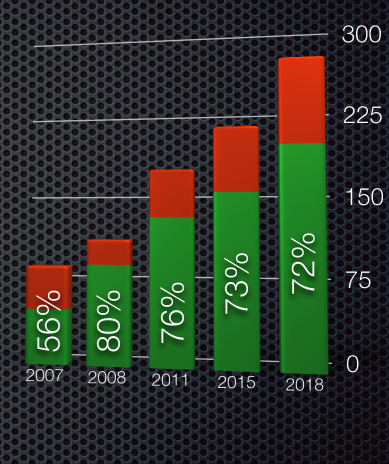




Calretinin / RUN 52 2018

Pass: 72 %

| Table 1. Antibodies a | nd a | ssessment marks fo | or CR, ru | ın 52 | | | | | |
|---|------------------|--|-----------|-------|------------|------|--------------------|---------------------------|--|
| Concentrated antibodies | n | Vendor | Optimal | Good | Borderline | Poor | Suff. ¹ | Suff. OPS ² | |
| mAb clone 2E7 | 1 | Immunologic | 1 | 0 | 0 | 0 | - | - | |
| mAb clone 5A5 | 3 1 | Leica/Novocastra Monosan | 1 | 1 | 2 | 0 | - | - | |
| mAb clone CAL6 | 7 | Leica/Novocastra | 1 | 3 | 0 | 3 | 57% | - | |
| mAb clone DAK-Calret 1 | 34 | Dako/Agilent | 9 | 8 | 8 | 9 | 50% | 81% | |
| rmAb clone BSR235 | 1 | Nordic Biosite | 1 | 0 | 0 | 0 | - | - | |
| rmAb clone SP13 | 3 2 2 2 | Cell Marque Immunologic Spring Bioscience Thermo Scientific | 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, 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 CAL6 | 1 | Leica/Novocastra | 1 | 0 | 0 | 0 | | - | |
| mAb clone DAK-Calret 1 IS/IR627 | 35 | Dako/Agilent | 14 | 19 | 2 | 0 | 94% | 97% | |
| mAb clone DAK-Calret 1 IS/IR627 ⁴ | 20 | Dako/Agilent | 0 | 4 | 11 | 5 | 20% | - | |
| mAb clone MX027 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 | - | - | |
| rmAh 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 | - | - | |
| Total | 269 | | 120 | 74 | 52 | 23 | - | | |
| 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



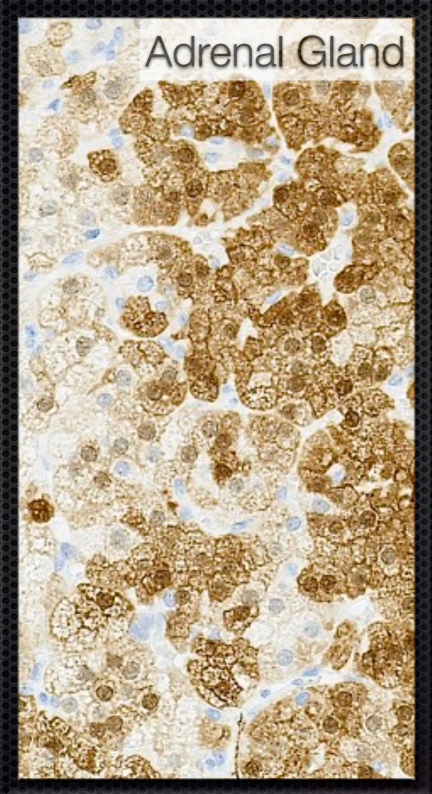


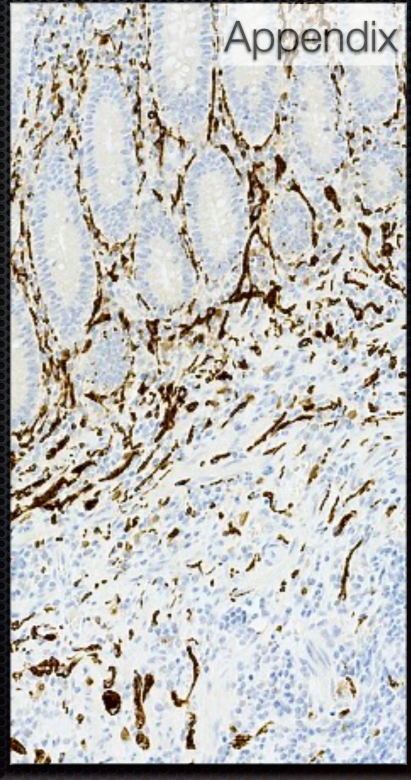
Calretinin / RUN 52 2018

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.









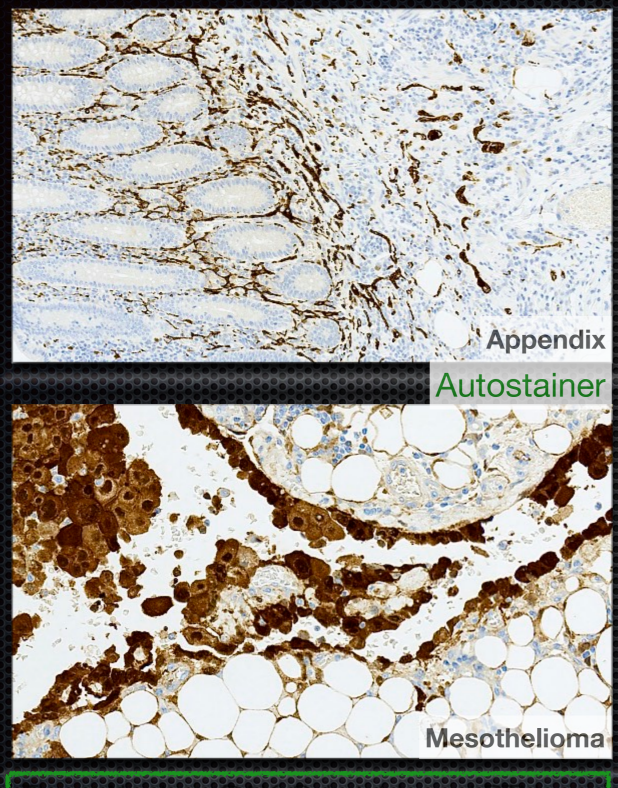
Calretinin / RUN 52 2018

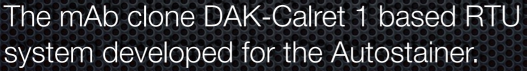
| Recommend- able clones | Retrieval | Titre | Detection | RTU | Detection |
|---------------------------|---------------|--------------|-----------|---------|-------------|
| mAb DAK- Calret1 | HIER, High pH | 1:20 - 1:100 | 3-step | Dako | 2 or 3-step |
| mAb 5A5 | HIER, High pH | 1:100 | 3-step | | |
| 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 | | | Ventana | 2 or 3-step |

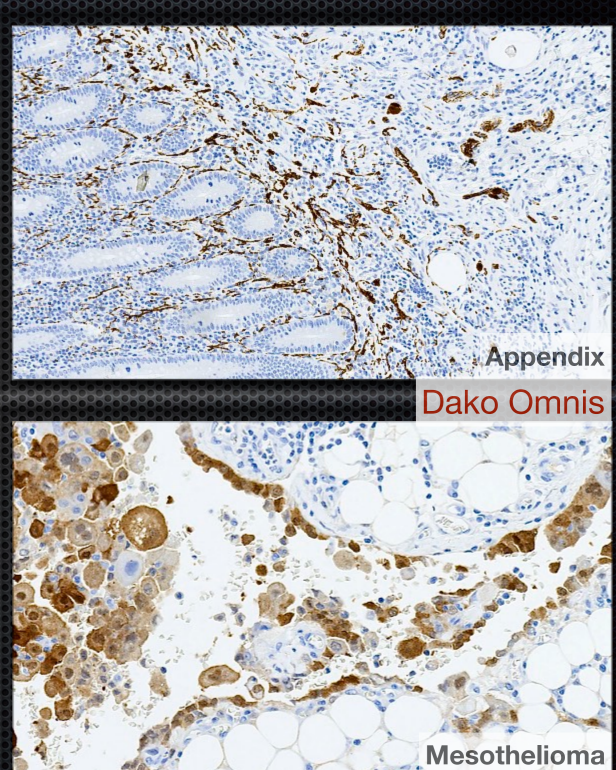
Table 3. Proportion of optimal results for CR for the most commonly used antibodies as concentrates on the 4 main IHC systems*

| Concentrated antibodies | Dako Autostainer Link / Classic | | Dako Omnis | | Ventana BenchMark GX / XT / Ultra | | 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 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 |





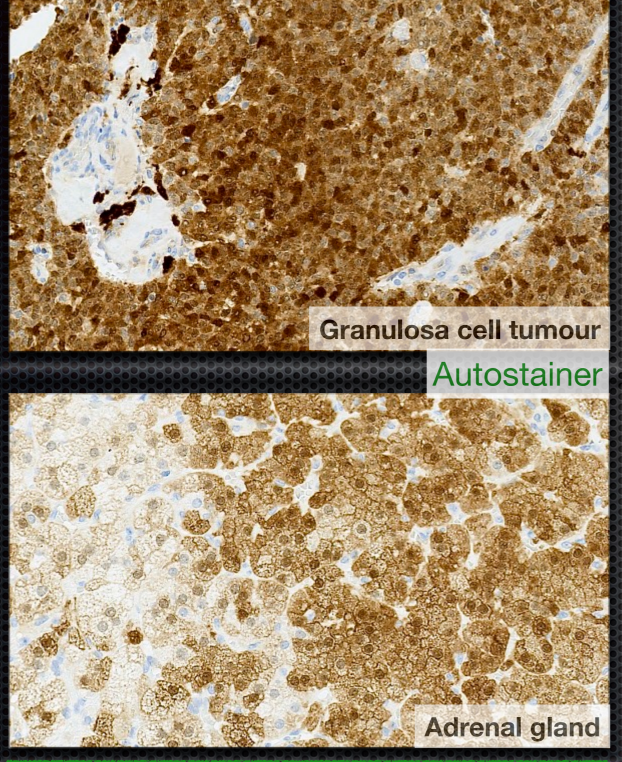


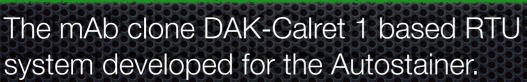


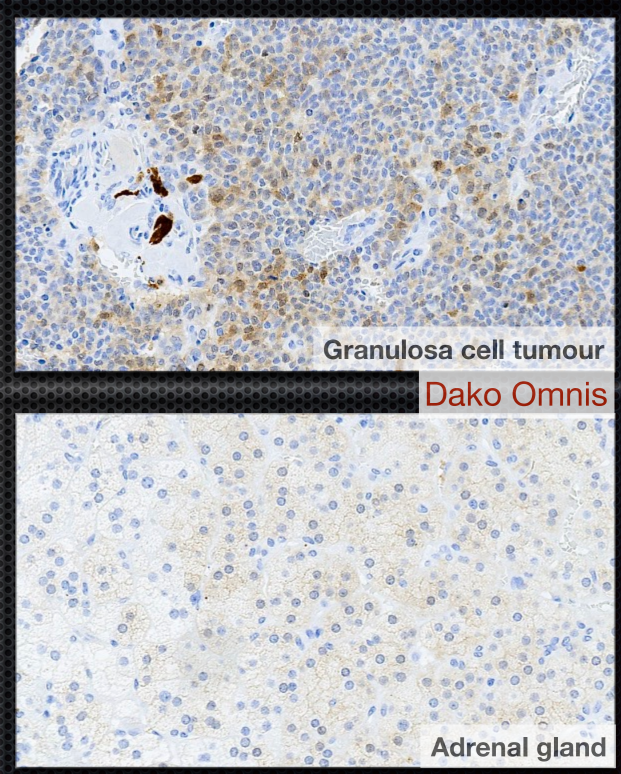
The mAb clone DAK-Calret 1 based RTU system developed for the Autostainer - but used on the Dako Omnis platform.





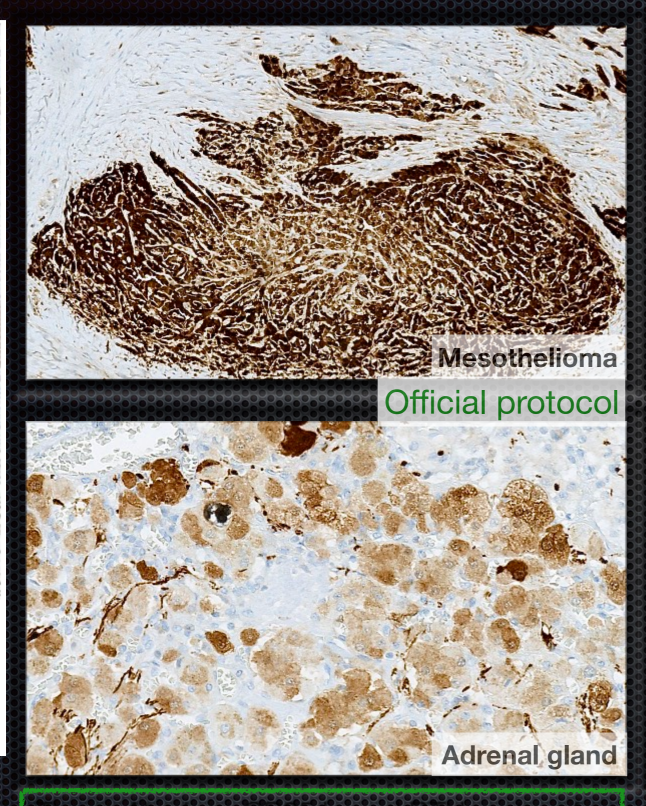


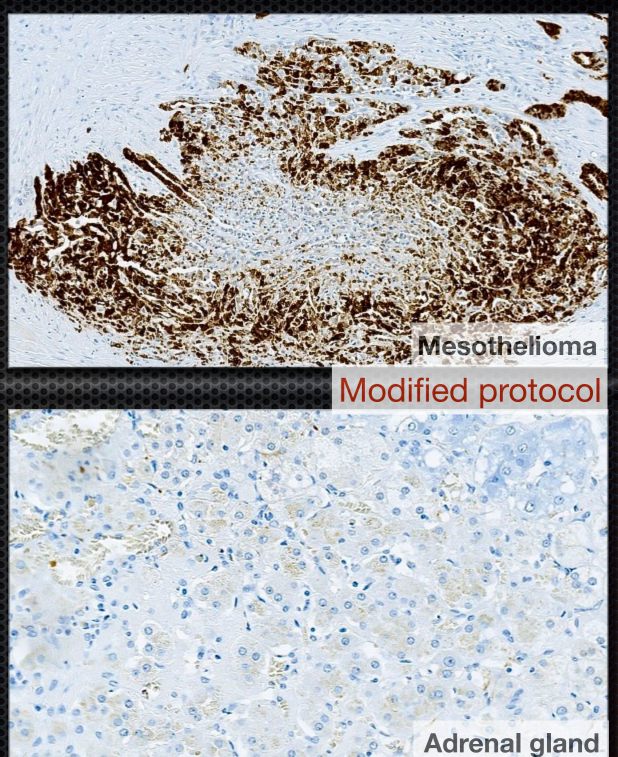




The mAb clone DAK-Calret 1 based RTU system developed for the Autostainer - but used on the Dako Omnis platform







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





p40 / RUN 48 2016

Pass: 74 %

| Table 1. Antibodies and assessment marks for p40, run 48 | | | | | | | | | |
|--|------------------------|---|---------|------|------------|------|--------------------|---------------------------|--|
| Concentrated antibodies | n | Vendor | Optimal | Good | Borderline | Poor | Suff. ¹ | Suff. OPS ² | |
| mAb clone BC28 | 77 6 2 2 1 | Biocare Zytomed Menarini abcam Nordic Biosite | 52 | 24 | 10 | 2 | 86% | 89% | |
| rmAb clone ZR8 | 12 1 1 | Immunologic Zeta Corporation BioSB | 1 | 6 | 2 | 5 | 50% | 67% | |
| pAb AC13030 | 8 | Biocare | 0 | 2 | 6 | 0 | - | - | |
| pAb RP163 | 5 | Diagnostic Biosystems | 0 | 1 | 1 | 3 | - | - | |
| pAb PC373 | 4 | Calbiochem, Merck | 0 | 1 | 0 | 3 | - | - | |
| pAb RBK054 | 3 | Zytomed | 0 | 0 | 1 | 2 | - | - | |
| pAb PI049 | 1 | DCS | 0 | 1 | 0 | 0 | - | - | |
| pAb PP123 | 1 | Pathnsitu | 0 | 0 | 1 | 0 | - | - | |
| Ready-To-Use antibodies | | | | | | | | | |
| mAb clone BC28 API/IPI/AVI 3066 | 13 | Biocare | 5 | 8 | 0 | 0 | 100% | 100% | |
| mAb clone BC28 790-4950 | 39 | Ventana | 19 | 15 | 5 | 0 | 87% | 94% | |
| mAb clone BC28 MSG097 | 1 | Zytomed | 1 | 0 | 0 | 0 | - | - | |
| mAb clone ZR8 MAD-000686QD | 3 | Master Diagnostica | 0 | 2 | 1 | 0 | - | - | |
| pAb API 3030 | 6 | Biocare | 0 | 0 | 4 | 2 | - | - | |
| pAb RAB-066 | 1 | Maixin | 0 | 1 | 0 | 0 | - | - | |
| pAb A00112 | 1 | Loxo GmbH | 0 | 0 | 1 | 0 | - | - | |
| Total | 188 | | 78 | 61 | 32 | 17 | - | | |
| Proportion | | | 42% | 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*

| Concentrated antibodies | Dako Autostainer Link / Classic | | • | | 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 |
| mAb clone BC28 | 7/20** (35%) | 1/1 | 6/8 (75%) | 1/1 | 30/42 (71%) | - | 3/6 (50%) | 0/1 |

^{*} 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)

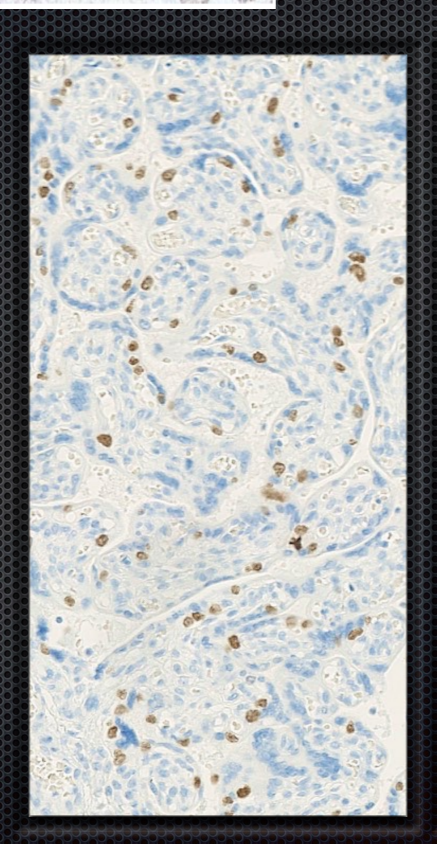
* Cytotrophoblasts must show an at least weak to moderate, distinct nuclear staining reaction.

Positive: Tonsil

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

Negative: Tonsil

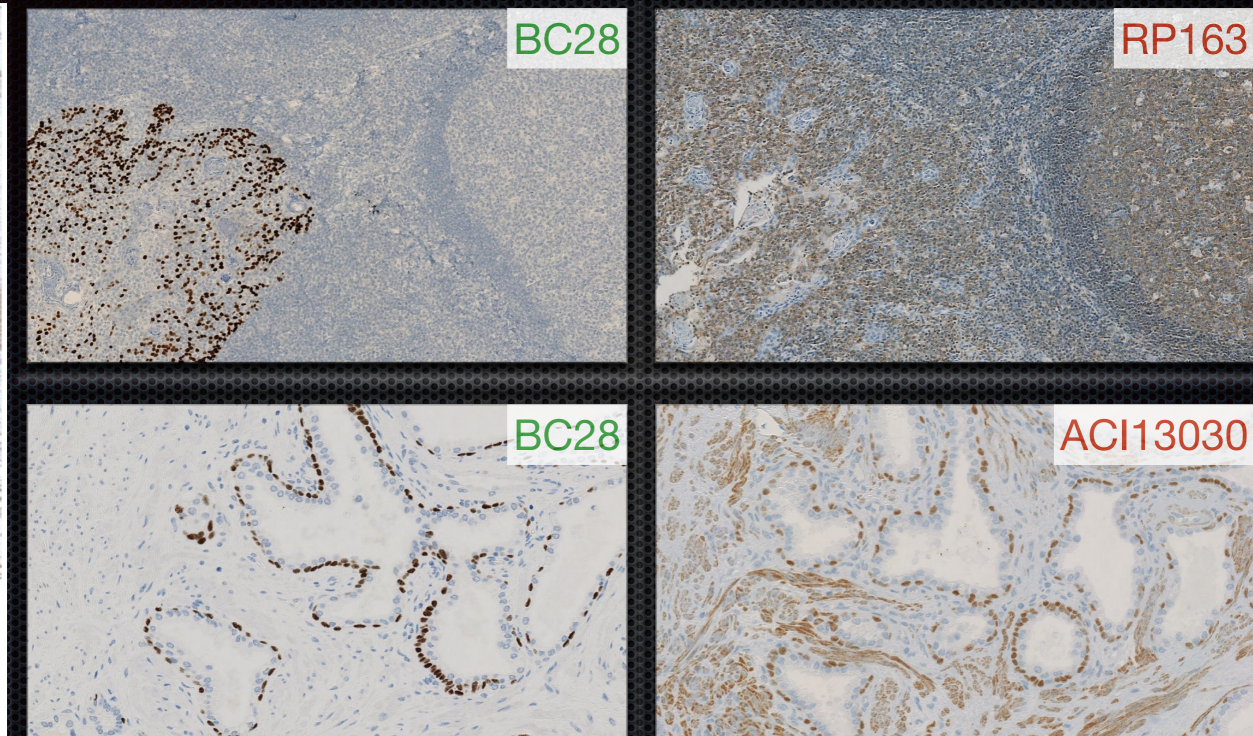
* Lymphocyttes must be negative.







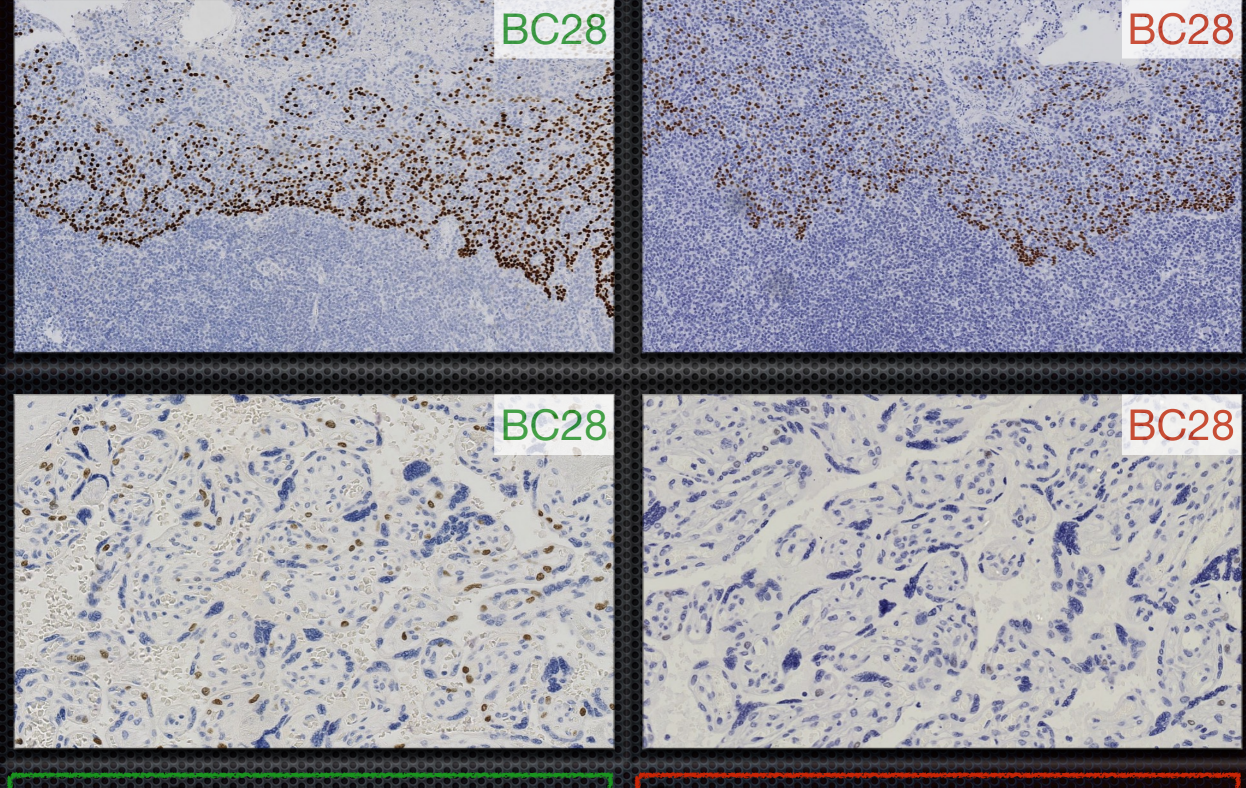




mAb clone BC28 optimally calibrated in a sensitive 3-step polymer system.

Poor signal to noise ratio using various pAb

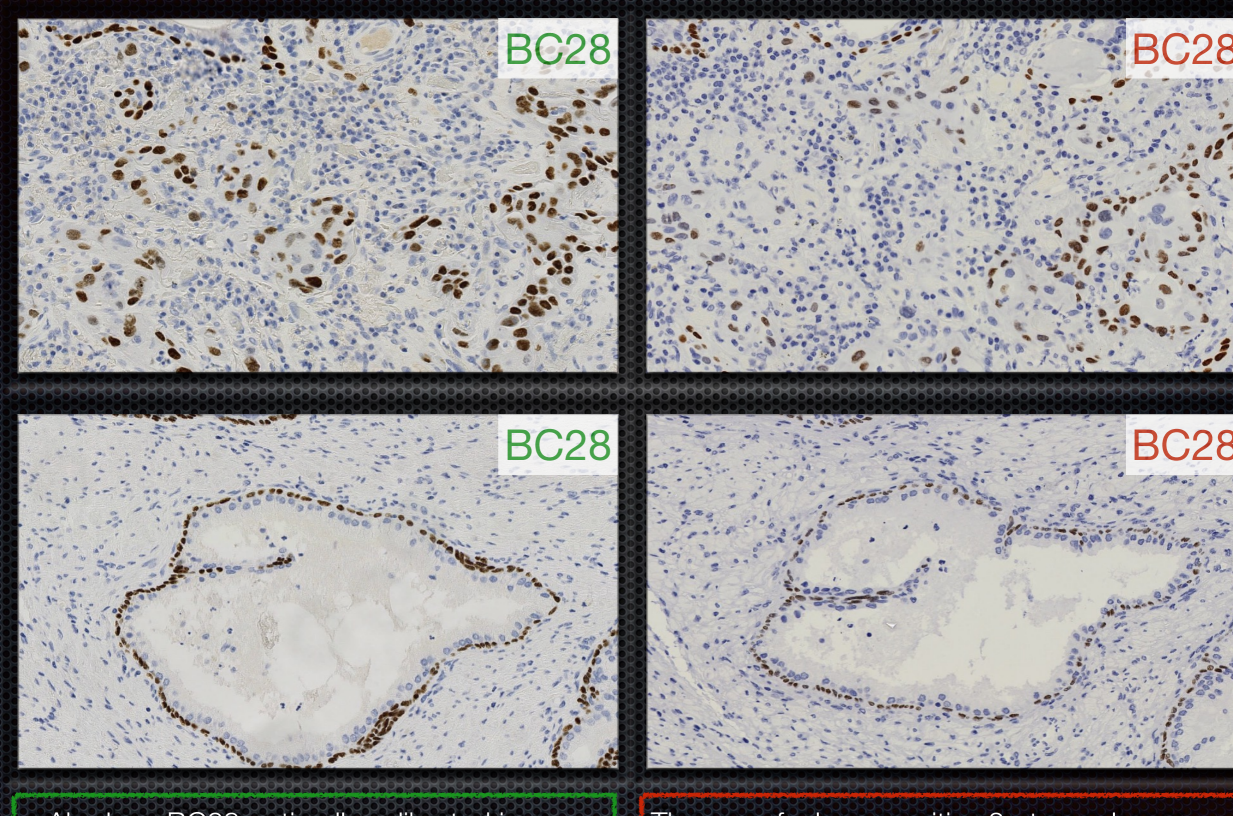




mAb clone BC28 optimally calibrated in a sensitive 3-step polymer system.

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



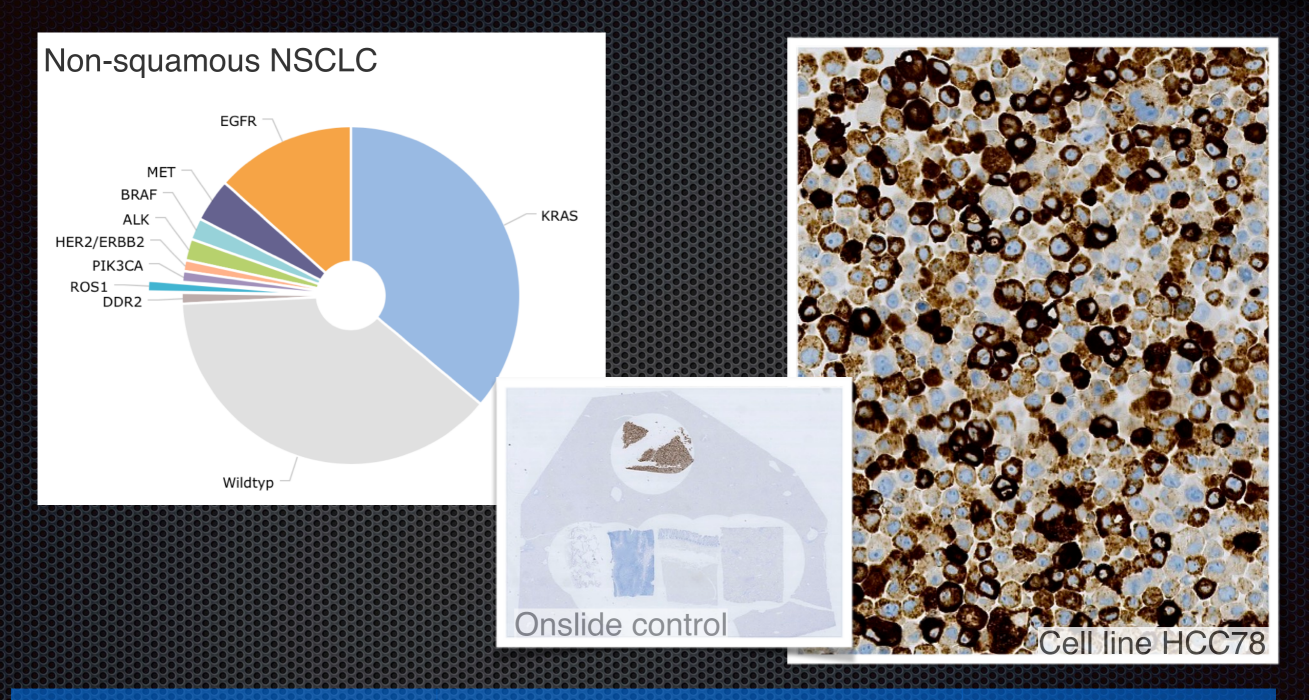


mAb clone BC28 optimally calibrated in a sensitive 3-step polymer system.

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

Driver mutations in lungcancer / ROS1





| Clone | Company | Retrieval | Titre | Detection |
|-----------|----------------|-----------|---------------|---|
| rmAb D4D6 | Cell Signaling | High pH* | 1:50 - 1:300* | 3-step polymer / multimer detection* |