



# **Anti - Cytokeratin 14**

# Rabbit clonal antibody

#### CAT#

CONCENTRATED **READY TO USE (RTU)** 

DB 099-0.1 DB 099-RTU-7  $(100 \mu l)$ (7 ml) DB 099-0.2  $(200 \mu I)$ DB 099-RTU-15 (15 ml) DB 099-0.5 (500 µl)

DB 099-1 (1 ml)

#### STORAGE AND APPLICATION

CONCENTRATED

+4°C +4°C, Do not freeze! Storage: Storage:

Application: IHC-P, Application: IHC-P,

dilution 1:100 ready to use

READY TO USE (RTU)

# PRODUCT INFORMATION

Clone:

20 mM Tris-HCl, pH 8.0 Buffer: Stabilizer: 20 mg/ml BSA Preservative: 0.05% NaN<sub>3</sub>

Specificity: Human

24 months from the shipping date Expiration:

Immunogen: Peptide derived from C-terminal region of human cytokeratin 14. Antibody recognizes the epitope

between Val455 - Lys471.

Cellular localization: cytoplasm

skin squamous cell carcinoma tissue, tonsil tissue Positive control:

Protein accession number: P02533

#### IHC-P PROTOCOL - INSTRUCTION MANUAL

- Deparaffinize the section in 3 changes of xylene, 5 minutes each.
- Wash the section in 96%, 80% and 70% ethyl alcohol for 10 minutes each.
- 3. Rinse in distilled water.
- Block the endogenous peroxidase by incubating the tissue in 3% hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>) for 10 minutes
- Wash in distilled water for 5 minutes.
- For antigen retrieval: immerse the slide in Tris-EDTA buffer, pH 9.0, 0.05% Tween-20\*, and incubate at 95°C in water bath for 30 minutes.(Alternatively adjust to your own protocol, keeping the required pH)
- Remove the staining to room temperature and let the slide to cool (in Tris-EDTA buffer, pH 9.0) for 15 minutes.
- Rinse in distilled water
- Wash in 0.05 M Tris-HCl, pH 7.6 buffer supplemented with 0.2% of Tween-20 (buffer A) for 5 minutes
- 10. CONCENTRATED:

Incubate the section with primary antibody at the dilution 1:100 for 1 hour in the closed wet chamber.

# **READY TO USE (RTU)**

Incubate the section with primary antibody (ready to use) for 1 hour in a closed wet chamber.

- Wash twice 5 minutes with buffer A.
- Apply the secondary antibody (the protocol depends on the supplier), and proceed to standard immunohistochemistry protocol (HRP Peroxide DAB). Micropolymer-HRP detection kit rabbit/mouse dual of DB Biotech is suggested (http://www.dbbiotech.com/products/detection-system.html)
- Wash twice 5 minutes with buffer A.
- Apply the chromogen (DAB), 1 3 minutes.
- Wash in water for 10 minutes.
- Stain in hematoxylin for 5 minutes
- Wash in water for 10 minutes.
- Dehydrate the section in 2 changes of 96% ethyl alcohol for 5 minutes each.
- Wash the section in 2 changes of xylene for 2 minutes each.
- Mount the slide for observation.

#### \*Tris-EDTA Buffer (10mM Tris Base, 1mM EDTA solution, 0.05% Tween-20, pH 9.0):

Tris ------- 1.21 g; EDTA ------ 0.37 g; Distilled water ------ 1000 ml Mix to dissolve in 700 ml of distilled water. Adjust pH to 9.0 with 1M HCl and then add 0.5 ml of Tween-20 and mix well. Adjust the final volume to 1 liter with distilled water. Store this solution at room temperature for 3 months or at 4°C for longer storage.

В

#### VENTANA PROTOCOL - INSTRUCTION MANUAL

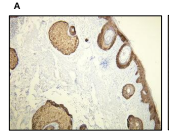
SHORT APPLICATION PROTOCOL FOR VENTANA BENCHMARK SLIDE STAINING SYSTEM

#### PROCEDURE: U ultraView DAB

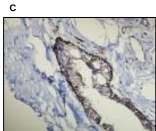
- Deparafinization
- Heating (72 °C) at the medium temperatures. Deparafinization.
- Cell conditioning 3.
- 4. ULTRA conditioner #1
- Heating glass (95 °C), incubation 8 min. (Cell conditioner #1; buffer CC1).
- 6 ULTRA CC1 solution application - 64 min.
- Antibody incubation temperature
- 8. Heating glass (36 °C), incubation 4 min.
- 9. Titration
- 10. Hand apply - primary antibody 100 µl. Incubation 36 min.
- 11. ultraWash
- 12. Nuclear stain
- 13. Hematoxylin II application - one drop (nuclear stain). Cover and incubate 12 min.
- After nuclear stain
- Bluing reagent application, one drop. After nuclear stain, cover and incubate 4 min

#### **PRECAUTIONS**

- We strongly recommend to use DB Primary Antibody Diluent (catalog number DB D-125, or DB D-250), eventually alternative diluent (containing protease free BSA at the concentrations ≥ 1mg/ml) for dilution of concentrated antibodies, otherwise the warranty might be voided.
- Centrifuge the vial before use.
- Intended for professional In Vitro Diagnostic use in laboratories.
- Do not use after expiration date stamped on vial label.
- Avoid contamination of the reagent.
- Any discrepancies in the recommended procedures stated in the working protocol may affect the final results.
- The reagent contains sodium azide  $(NaN_3)$  which is highly toxic in higher concentrations. The concentration in the reagent (0.05%) is not considered as hazardous.
- Disposal of waste material must be conducted in accordance with local regulations
- Wear appropriate Personal Protective Equipment to avoid contact with eyes and skin.







CK14 expression in the epidermal and skin-adnexal epithelial cells (A), myoepithelial cells of DCIS of the breast (B) and ductal myoepithelial cells of the breast (C). All, formalin fixed, paraffin embedded human tissues (4 µm sections) stained with anti - Cytokeratin 14 (DB 099) monospecific clonal antibody according to related DB Biotech datasheet.

## LEICA BOND MAX PROTOCOL -**INSTRUCTION MANUAL**

SHORT APPLICATION PROTOCOL FOR LEICA BOND MAX SLIDE STAINING SYSTEM

### Protocol F:

- Visualization system: BOND Refine DS9800
- Epitope retrieval / heating time / temperature: ER2 / 30 min. / 100 °C
- Incubation of primary antibody / temperature: 30 min. / 20 °C

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