



Anti - C-erbB-2 (Her-2/neu)

Rabbit clonal antibody

CAT#

CONCENTRATED READY TO USE (RTU)

DB 094-0.1 (100 μ l) DB 094-RTU-7 (7 ml) DB 094-0.2 (200 μ l) DB 094-RTU-15 (15 ml) DB 094-0.5 (500 μ l)

DB 094-1 (1 ml)

STORAGE AND APPLICATION

CONCENTRATED READY TO USE (RTU)

Storage: +4°C, Do not freeze!

Application: IHC-P, dilution 1:100 Application: IHC-P, ready to use

PRODUCT INFORMATION

Clone: A24-V

Buffer: 20 mM Tris-HCl, pH 8.0 Stabilizer: 20 mg/ml BSA Preservative: 0.05% NaN₃

Specificity: Human

Expiration: 24 months from the shipping date

Immunogen: Peptide derived from C-terminal region of human CerbB-2. Antibody recognizes the epitope between

Pro1233 - Pro1254.

Cellular localization: membrane, cytoplasm

Positive control: breast ductal carcinoma tissue

Protein accession number: P04626

IHC-P PROTOCOL - INSTRUCTION MANUAL

- 1. Deparaffinize the section in 3 changes of xylene, 10 minutes each.
- 2. Wash the section in 96%, 80% and 70% ethyl alcohol for 10 minutes each.
- 3. Rinse in distilled water, 2 x 5 minutes.
- 4. Block the endogenous peroxidase by incubating the tissue in 3% hydrogen peroxide (H₂O₂) for 10 minutes.
- Wash in distilled water, 2 x 5 minutes.
- 6. For antigen retrieval use one of the following procedures: A) Immerse the slide in Tris-EDTA buffer, pH 9.0, 0.05% Tween- 20*, and incubate at 95-97°C in water bath for 20-25 minutes, or B) Immerse the slide in citrate buffer, pH 6.0, 0.05% Tween-20**, and incubate at 95-97°C in water bath for 20-25 minutes. (Alternatively adjust to your own protocol, keeping the required pH).
- Remove the staining to room temperature and let the slide to cool down in antigen retrieval buffer for 15 minutes.
- 8. Rinse in distilled water, 2 x 5 minutes.
- Wash in PBS (phosphate buffer saline, pH 7.0-7.5) supplemented with 0.05% of Tween-20 (buffer A), 2 x 5 min
- 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):

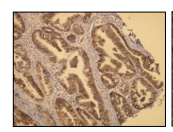
Incubate the section with primary antibody (ready to use) for 1 hour in a closed wet chamber.

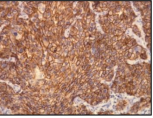
- 11. Wash 3 x 5 minutes with buffer A.
- Apply the secondary antibody (the protocol depends on the supplier), and proceed to standard immunohistochemistry protocol (HRP - Peroxide - DAB).
 Micropolymer-HRP detection kit rabbit/mouse dual of DB Biotech is suggested (http://www.dbbiotech.com/products/detection-system.html).
- 13. Wash 3 x 5 minutes with buffer A.
- 14. Apply the chromogen (DAB), 1 3 minutes.
- 15. Wash in water, 2 x 5 minutes.
- 16. Stain in hematoxylin for 5 minutes.
- 17. Wash in distilled water, 3 x 2 minutes.
- 18. 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

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.

** Citrate Buffer (10mM Citric Acid, 0.05% Tween-20, pH 6.0):





C-erbB-2 protein expression (3+) in adenocarcinoma of stomach (A) and in breast ductal carcinoma (B). Formalin fixed, paraffin embedded human tissues (4 µm sections) stained with anti - C-erbB-2 (DB 094) monospecific antibody according to related DB Biotech datasheet.

VENTANA PROTOCOL – INSTRUCTION MANUAL

SHORT APPLICATION PROTOCOL FOR VENTANA BENCHMARK SLIDE STAINING SYSTEM

PROCEDURE: U ultraView DAB

- 1. Deparafinization
- 2. Heating (72 °C) at the medium temperatures. Deparafinization.
- 3. Cell conditioning
- 4. ULTRA conditioner #1
- 5. Heating glass (95 °C), incubation 8 min. (Cell conditioner #1; buffer CC1).
- 6. ULTRA CC1 solution application 36 min.
- 7. Antibody incubation temperature
- 8. Heating glass (36 °C), incubation 4 min.
- 9. Titration
- 10. Hand apply primary antibody 100 μ l. Incubation 44 min.
- 11. ultraWash
- 12. Nuclear stain
- 13. Hematoxylin II application one drop (nuclear stain). Cover and incubate 12 min.
- 14. After nuclear stair
- 15. Bluing reagent application, one drop. After nuclear stain, cover and incubate 4 min

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 / 20 min. / 100 °C
- Incubation of primary antibody / temperature: 30 min. / 20 °C

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.
- 2. Centrifuge the vial before use.
- 3. Intended for professional In Vitro Diagnostic use in laboratories.
- Do not use after expiration date stamped on vial label.
- 5. 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 (NaN₃) which is highly toxic in higher concentrations.
 The concentration in the reagent (0.05%) is not considered as hazardous.
- 8. Disposal of waste material must be conducted in accordance with local regulations.
- 9. Wear appropriate Personal Protective Equipment to avoid contact with eyes and skin.

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