

Anti - PLAP (Placental Alkaline Phosphatase)

Rabbit clonal antibody

CAT#

DB 047-1

1.

 CONCENTRATED

 DB 047-0.1
 (100 µl)

 DB 047-0.2
 (200 µl)

 DB 047-0.5
 (500 µl)

READY TO USE (RTU) DB 047-RTU-7 (7 ml) DB 047-RTU-15 (15 ml)

STORAGE AND APPLICATION CONCENTRATED

(1 ml)

Storage: +4°C Application: IHC-P, dilution 1:100 READY TO USE (RTU)

Storage: +4°C, Do not freeze! Application: IHC-P, ready to use

PRODUCT INFORMATION	
Clone:	P16-D
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 the middle portion of human PLAP sequence. Antibody recognizes the epitope between Glu238 - Leu250.
Cellular localization: membrane	
Positive control:	human testicular tumor tissue
Protein accession number: P05187	

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).
- ULTRA CC1 solution application 64 min.
- Antibody incubation temperature
- 8. Heating glass (36 °C), incubation 4 min.
- 9. Titration
- Hand apply primary antibody 100 µl. Incubation 32 min.
 ultraWash
- 11. ultraWash
 12. Nuclear stain
- 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

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

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.
- 4. 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.
 Disoosal of waste material must be conducted in accordance with local regulations.
- Wear appropriate Personal Protective Equipment to avoid contact with eyes and skin.

Wash the section in 96%, 80% and 70% ethyl alcohol for 10 minutes each.
 Rinse in distilled water.

IHC-P PROTOCOL – INSTRUCTION MANUAL

 Block the endogenous peroxidase by incubating the tissue in 3% hydrogen peroxide (H₂O₂) for 10 minutes.

Deparaffinize the section in 3 changes of xylene, 5 minutes each.

- 5. Wash in distilled water for 5 minutes.
- For antigen retrieval: immerse the slide in the citrate buffer, pH 6.0, 0.05% Tween-20*, and incubate in water bath at 96-98°C for 30-40 minutes (Alternatively adjust to your own protocol, keeping the required pH)
- Transfer the slide to room temperature and let it cool down (in citrate buffer, pH 6.0) for 15 minutes.
- 8. Rinse in distilled water.
- 9. 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.

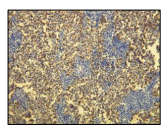
READY TO USE (RTU):

Incubate the section with primary antibody (ready to use) for 1 hour in a closed wet chamber. Wash twice 5 minutes with buffer A.

- 11. Wash twice 5 minutes with buffer /
- 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 (<u>http://www.dbbiotech.com/products/detection-system.html</u>).
- 13. Wash twice 5 minutes with buffer A.
- 14. Apply the chromogen (DAB), 1 3 minutes.
- 15. Wash in water for 10 minutes.
- 16. Stain in hematoxylin for 5 minutes.
- 17. Wash in water for 10 minutes.
- 18. Dehydrate the section in 2 changes of 96% ethyl alcohol for 5 minutes each.
- 19. Wash the section in 2 changes of xylene for 2 minutes each.
- 20. Mount the slide for observation.

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

Citric acid (anhydrous) ------ 1.92 g; Distilled water ------ 1000 ml Mix to dissolve in 700 ml of distilled water. Adjust pH to 6.0 with 1M NaOH 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.



Diffuse cytoplasmic PLAP positivity in the typical seminoma. Formalin fixed, paraffin embedded human tissue (4 µm section) stained with anti - PLAP (DB 047) monospecific clonal antibody according to related DB Biotech datasheet.