

# For Research Use Only (RUO)

# p504s/AMACR

Mouse Monoclonal Antibody

# [Catalog Number]

#### REF 0504

# [Package Size]

Ready to use:1mL2mL3mL5mL6mLConcentrated:0.1mL0.2 mL0.5mL1.0mL

# [Intended Use]

Mouse Monoclonal anti-p504s/AMACR antibody is intended for use to qualitatively identify p504s/AMACR antigen by light microscopy in sections of formalin-fixed, paraffin-embedded tissue using IHC detection methodology.

# This antibody is for research use only (RUO).

# [Specimen Collection and Preparation for Analysis]

Formalin-fixed, paraffin-embedded tissues.

Each section should be cut to the appropriate thickness  $(2-5 \ \mu m)$  for the primary antibody being used and placed on a positively charged glass microscope slide.

# [Storage and Handling]

Store at 2-8°C. Do not freeze.

Do not use product beyond the expiration date for the storage method.

# [Reagents Provided]

#### Clone: 13H4

Buffer: 10mM pH 7.4 Phosphate Puffer Saline (PBS).

Stabilizer: 0.05% bovine serum (BSA).

Preservative: 0.05% sodium azide (NaN<sub>3</sub>).

Ready-to-use antibody concentration: 2-5µg/mL.

Concentrated antibody concentration: 50-200µg/mL.

# [Staining Procedure]

 Deparaffinized slides in 3 changes of xylene (or Dewax solution), 10 minutes each. and hydrate through a graded series alcohols.

- 2. Wash the section in 90%, 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  $(H_2O_2)$  for 10 minutes.
- 5. Wash in distilled water, 2 x 5 minutes.
- Antigen retrieval: Place slides in a pressure cooker filled with Epitope Retrieval Solution (EDTA, pH 9.0) buffer.
- 7. Wash in PBS 2 x 5 minutes.
- Concentrated Antibody Dilution
  Suggested Dilution: 1:100-1:200
  The optimal dilution for a specific application under a given set of experimental conditions should be determined by the investigator.
- 9. Add 100 $\mu$ L primary antibody, Incubate for 30 minutes . Wash in PBS 2 x 5 minutes.
- Add 100µL secondary antibody (the protocol depends on the supplier), and proceed to standard immunohistochemistry protocol. Wash in PBS 2 x 5 minutes.
- 11. Add 100µL DAB solution (the protocol depends on the supplier), Incubate for 2-10 minutes. Wash in PBS 2 x 5 minutes.
- 12. Counterstain with hematoxylin. Rinse with deionized water.

# [Contact Information]



# Shanghai Long Island Antibody Diagnostica Inc.

Add: 100A, Building #1, No. 288 Hangnanzhi Road, Zhuanghang Industrial Zone, Fengxian District, 201415 Shanghai, People's Republic of China Tel: +86 21-64910505; 400-920-0015 Fax: +86 21-57469228 Email: sales@longislandab.com Web: www.longislandab.com