# SYSY HistoSure

## **Reference Protocol for Anti-Ig kappa light chain** (HS-346 011) Immunohistochemistry using DAB as Chromogen

Braun, Multigourmet

X0909

S2022

115-065-146

1.07298.0250

PK-4000

SK-4105

Agilent

Agilent

Jackson

Vectorlabs

Vectorlabs

Vectorlabs SP-2001

Merck

### **Tissue Fixation**

• 3.7% formaldehyde (24 h), 3.5 μM paraffin sections

### Materials and Reagents

- Food Steamer
- Staining Containers with slide holders (e.g. Tissue-Tek)
- Protein Block, Serum-Free
- Antibody diluent
- · Biotinylated anti-mouse antibody
- ABC HRP Kit, Standard
- ImmPACT DAB
- Hydrogen peroxide 30%
- PBS (pH 7.4)
- TBST (TBS, 0.05% Tween 20, pH 7.6)
- Antigen Retrieval buffer: Citrate Buffer (10 mM Citrate, 0.05% Tween 20, pH 6.0)
- Xylene, 100% ethanol, 90% ethanol, 80% ethanol and 70% ethanol, 2-propanol
- Optional: Hematoxylin Solution (Mayer's, Modified) or other nuclear counterstain
- · Optional: Avidin/Biotin Blocking Kit
- Non-aqueous mounting medium

### Method

#### 1. Deparaffinize and hydrate tissue sections

- a. Xylol 2 x 5 min 100% EtOH b. 2 x 2 min 90% EtOH 1 x 2 min C. 1 x 2 min d. 80% EtOH 70% EtOH 2 x 2 min е f. Deionized Water 1 x 20 sec
- g. PBS
  - 1 x 2 min

\*Keep the slides in PBS until ready to perform the Antigen Retrieval. Do not allow the slides to dry out\*

Synaptic Systems
Rudolf-Wissell-Str. 28a
37079 Göttingen, Germany

# SYSY | HistoSure

#### 2. Antigen Retrieval (AR) using a food steamer

- a. Heat the steamer with a suitable staining container filled with Antigen Retrieval buffer to ~97°C
- b. Transfer the sections into the staining box, wait until the temperature reaches 97°C
- c. Incubate the sections in the steamer for 30 min
- Remove the staining container from the steamer and allow the slides to cool down for 20 min (target end temperature ~60°C)
- 3. Wash slides in PBS, 3 x 1 min
- 4. Blocking endogenous peroxidase activity
  - a. Incubate the sections with 3% hydrogen peroxide in PBS (freshly prepared!) for  ${\bf 5}\ {\bf min}$
- 5. Wash slides in PBS, 2 x 1 min
- 6. Wash slides in TBST, 1 x 2 min
- 7. **Optional:** Perform Avidin-Biotin-Block according to manufacturer's instructions. *Note:*Certain tissues (e.g. liver, kidney) contain high levels of endogenous biotin. The Avidin-Biotin blocking step is recommended when using the ABC system for these tissues. If the background problem persists, consider trying a polymer-based detection system instead of biotinylated secondary antibody / ABC system.
- 8. Block in Protein Block, Serum-Free for 10 min
- 9. Drain slides (do not rinse)
- 10. Apply primary antibody diluted in Antibody Diluent and incubate in a humidified chamber for 1 h at room temperature

#### \*Suggested dilution: 1:5000 in Antibody Diluent\*

11. Wash slides in TBST, 3 x 2 min

#### 12. Apply secondary antibody diluted in Antibody Diluent for 30 min at room temperature. \*Suggested concentration: 5 μg/ml\* \*Perform step 13 in the interim\*

- 13. Prepare the ABC-reagent: 5 ml PBS + 1 drop A + 1 drop B, incubate for 30 min
- 14. Wash slides in TBST, 3 x 2 min

#### 15. Apply the ABC reagent for 30 min at room temperature

- 16. Wash slides in TBST, 3 x 2 min
- 17. Apply the DAB substrate, 1-10 min

\*Observe the staining with a microscope! Development times may differ depending upon the level of antigen\*

- 18. Stop the DAB reaction with deionized water
- 19. Optional: Counterstain
  - a. Follow the manufacturer's instructions for counterstaining and bluing
- 20. Wash slides in deionized water for 1 min

#### 21. Dehydrate tissue sections:

- a. 70% EtOH 2 x 10 sec
- b. 80% EtOH 1 x 10 sec
- c. 90% EtOH 1 x 10 sec
- d. 2-Propanol 2 x 1 min
- e. Xylol 3 x 2 min

#### 22. Mount slides in a suitable organic mounting medium and add coverslip

Note: The SYSY standard protocol generates good results in the SYSY labs and may be used as a reference. However, to achieve the highest

## SYSY | HistoSure

specific signal and lowest non-specific background signal, the best antigen retrieval condition, antibody concentration, incubation temperature, and incubation time must be determined individually. Please also refer to our general protocols.