

Reference Protocol for Anti-IBA1 (HS-234 013) Immunohistochemistry using DAB as Chromogen

Tissue Fixation

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

Materials and Reagents

Food Steamer	Braun, Multigourmet	
• Staining Containers with slide holders (e.g. Tissue-Tek)		
Protein Block, Serum-Free	Agilent	X0909
Antibody diluent	Agilent	S2022
 Biotinylated anti-rabbit antibody 	Jackson	111-065-144
ABC HRP Kit, Standard	Vectorlabs	PK-4000
ImmPACT DAB	Vectorlabs	SK-4105
Hydrogen peroxide 30%	Merck	1.07298.0250
• 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	Vectorlabs SP-2001	

• Non-aqueous mounting medium

Method

1) Deparaffinize and hydrate tissue sections

- a) Xylol 2 x 5 min b) 100% EtOH 2 x 2 min
- c) 90% EtOH 1 x 2 min
- d) 80% EtOH 1 x 2 min
- e) 70% EtOH 2 x 2 min
- 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

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
- d) 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 5 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: 1:100 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 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.