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# Anti-Phospho-p62 (SQSTM1) (Ser351) mAb

CODE No.	M217-3
CLONALITY	Monoclonal
CLONE	5D5
ISOTYPE	Mouse IgG1 κ
QUANTITY	100 $\mu$ L, 1 mg/mL
SOURCE	Purified IgG from hybridoma supernatant
IMMUNOGEN	KLH conjugated synthetic peptide, CKEVDP(pS)TGELQSLQ (corresponding to amino acid residues 346-359 of mouse p62 (SQSTM1))
FORMULATION	PBS containing 50% Glycerol (pH 7.2). No preservative is contained.
STORAGE	This antibody solution is stable for one year from the date of purchase when stored at -20°C.

### **APPLICATIONS-CONFIRMED**

Western blotting0.5 μg/mL for chemiluminescence detection systemImmunohistochemistry1 μg/mL (paraffin section)Heat treatment for paraffin embedded section: microwave oven, for 20 min. in 10 mM citrate buffer (pH 6.3)Immunocytochemistry0.1 μg/mL

### SPECIES CROSS REACTIVITY on WB

Species	Human	Mouse	Rat	Hamster
Cells	huH-1	sodium arsenite-treated MEF, MEF <sup>Atg5-/-</sup>	Not tested	Not tested
Reactivity	+	+		

 Entrez Gene ID
 8878 (Human), 18412 (Mouse)

 REFERENCES
 1) Inoue, H., et al., Biomed Res. 38, 343-350 (2017) [WB]

 2) Santarino, I. B., et al., Sci. Rep. 7, 5812 (2017) [WB, IC]

 3) Watanabe, Y., et al., Autophagy 13, 133-148 (2017) [IC]

 4) Kageyama, S., et al., J. Biol. Chem. 289, 24944-24955 (2014)

5) Ichimura, Y., et al., Mol. Cell 51, 618-631 (2013)

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# **RELATED PRODUCTS**

	111020010
Antibodies	
PM074	Anti-Phospho-p62 (SOSTM1) (Ser351) pAb
D343-3	Anti-Phospho-p62 (SOSTM1) (Ser403) mAb (4F6)
D344-3	Anti-Phospho-p62 (SOSTM1) (Ser403) mAb (4C8)
PM045	Anti-p62 (SOSTM1) pAb
M162-3	Anti-p62 (SOSTM1) (Human) mAb (5F2)
M162-A48	Anti-p62 (SQSTM1) (Human) mAb
101102 1140	-Alexa Fluor <sup>®</sup> $488(5F2)$
M162-A59	Anti-n62 (SOSTM1) (Human) mAb
11102 1139	-Alexa Fluor <sup>®</sup> 594 (5F2)
M162-A64	Anti-n62 (SOSTM1) (Human) mAb
101102 1101	-Alexa Fluor <sup>®</sup> $647 (5F2)$
PM066	Anti-n62 C-terminal nAb
PM066-7	Anti-p62 C-terminal pAb-HRP-DirecT
PM036	Anti-I C3 nAb [WB IP IC IHC FCM]
M152 3	Anti I C3 mAb (AF12) [WB IP IC ECM EM]
M186-3	Anti-LC3 mAb $(4E12)$ [WB, II, IC, I CW, EW] Anti-LC3 mAb $(8E10)$ [WB]
M186 7	Anti I C3 mAb HPP DirecT (8E10)
PD014	Anti I C3 nAb [WB]
PD017	Anti Baclin 1 nAb
PM037	Anti GABABABAD
M125 2	Anti $CABADAD mAb$ (1E4)
PM038	Anti GATE 16 pAb
PD041	Anti-OATE-TO PAO
DM024	Anti-Atg2A pAb
FW1034 M122 2	Anti Ata2 mAb (2E8)
M124 2	Anti-Atg $J$ IIIAU (JE $\delta$ )
DM050	Anti-Atg4D IIIAU (9H3)
FINIU30 M152 2	Anti-Atg5 $pA0$
M133-3	Anti-Atg $3$ IIIA $0$ (4D $3$ )
PM059	Anti-Atg7 (Human) pAb
PD042	Anti-Atg9A pAD Anti-Atg10 (II $\rightarrow$ Atg(5A7)
M151-3	Anti-Atg10 (Human) mAb (SA/)
M154-5	Anti-Atg12 (Human) mAD (6E5)
PD036	Anti-Atg13 (Human) pAb
M183-3	Anti-Atg13 mAb (504)
PD020	Anti-Atg14 pAD $A_{11}(11) = A_{11}(11)$
M184-3	Anti-Atg14 (Human) mAD (4H8)
PM040	Anti-Atg16L pAb
M150-3	Anti-Atg16L mAb (1F12)
M160-3	Anti-UVRAG mAb (1H4)
PD027	Anti-Rubicon (Human) pAb
M1/0-3	Anti-Rubicon (Human) mAb (1H6)
PD037	Anti-Tel2 pAb
PM069	Anti-NRF2 pAb
M200-3	Anti-NRF2 mAb (1F2)
PM072	Anti-VMP1 pAb
PM0/6	Anti-Syntaxin-17 (Human) pAb
M212-3	Anti-Syntaxin-17 (Human) mAb (2F8)
M224-3	Antı-KEAPI mAb (KPI)
M230-3	Anti-Parkin mAb (Par6)

<u>Kits</u>

8485	Autophagy Ab Sampler Set
8486	Autophagy Watch
CY-7055	CycLex <sup>®</sup> Total p62 ELISA Kit
CY-7056	CycLex <sup>®</sup> Phospho-p62 Ser349 ELISA Kit
CY-7057	CycLex <sup>®</sup> Phospho-p62 Ser403 ELISA Kit
PM036-PN	Positive control for anti-LC3 antibody

WB: Western blotting IP: Immunoprecipitation IC: Immunocytochemistry IHC: Immunohistochemistry FCM: Flow cytometry EM: Immuno-electron microscopy

Other related antibodies and kits are also available. Please visit our website at <u>http://ruo.mbl.co.jp/</u>

## **SDS-PAGE & Western blotting**

- 1) Wash 1 x  $10^7$  cells 3 times with PBS and suspend with 1 mL of Laemmli's sample buffer, then sonicate briefly (up to 20 sec.).
- 2) Centrifuge the tube at 12,000 x g for 5 min. at 4°C and transfer the supernatant to another tube.
- 3) Boil the samples for 3 min. and centrifuge. Load 10 µL of the sample per lane in a 1-mm-thick SDS-polyacrylamide gel (10% acrylamide) for electrophoresis.
- 4) Blot the protein to a polyvinylidene difluoride (PVDF) membrane at 1 mA/cm<sup>2</sup> for 1 hr. in a semi-dry transfer system (Transfer Buffer: 25 mM Tris, 190 mM glycine, 20% MeOH). See the manufacturer's manual for precise transfer procedure.
- 5) To reduce nonspecific binding, soak the membrane in 10% skimmed milk (in PBS, pH 7.2) for overnight at 4°C.
- 6) Wash the membrane with PBS-T [0.05% Tween-20 in PBS] (5 min. x 3 times).
- 7) Incubate the membrane with primary antibody diluted with 1% skimmed milk (in PBS, pH 7.2) as suggested in the **APPLICATIONS** for 1 hr. at room temperature. (The concentration of antibody will depend on the conditions.)
- 8) Wash the membrane with PBS-T (5 min. x 3 times).
- 9) Incubate the membrane with 1:10,000 Anti-IgG (Mouse) pAb-HRP (MBL; code no. 330) diluted with 1% skimmed milk (in PBS, pH 7.2) for 1 hr. at room temperature.
- 10) Wash the membrane with PBS-T (5 min. x 3 times).
- 11) Wipe excess buffer on the membrane, then incubate it with appropriate chemiluminescence reagent for 1 min. Remove extra reagent from the membrane by dabbing with paper towel, and seal it in plastic wrap.
- 12) Expose to an X-ray film in a dark room for 3 min. Develop the film as usual. The condition for exposure and development may vary.

(Positive controls for Western blotting; sodium arsenite-treated MEF, MEF<sup>Atg5-/-</sup> and huH-1)



# Western blot analysis of Phospho-p62 (SQSTM1) (Ser351)

Lane 1: MEF, sodium arsenite-treated (10  $\mu$ M, 12 hr.) Lane 2: MEF Lane 3: MEF<sup>Atg5-/-</sup> Lane 4: huH-1 Lane 5: huH-1,  $\lambda$ -phosphatase-treated Lane 6: p62-knockout huH-1

Immunoblotted with Anti-Phospho-p62 (SQSTM1) (Ser351) mAb (M217-3)

Sodium arsenite-treated MEF and p62-knockout huH-1 were provided by Dr. Yoshinobu Ichimura<sup>1</sup> and Dr. Masaaki Komatsu<sup>2</sup>. (<sup>1</sup>Protein Metabolism Project, Tokyo Metropolitan Institute of Medical Science, <sup>2</sup>Department of Biochemistry, School of Medicine, Niigata University)

MEF<sup>Atg5-/-</sup> was provided by Dr. Noboru Mizushima. (Department of Biochemistry and Molecular Biology, Graduate School and Faculty of Medicine, The University of Tokyo)

### **Immunohistochemistry**

- 1) Deparaffinize the sections with Xylene 3 times for 5 min. each.
- 2) Wash the slides with Ethanol 3 times for 5 min. each.
- 3) Wash the slides with PBS 3 times for 5 min. each.
- 4) Remove the slides from PBS and heat-treated with 10 mM Citrate buffer (pH 6.3) for 20 min. using microwave.
- 5) Let the slides cool down at room temperature in the Citrate buffer.
- 6) Wash the slides with running water for 5 min., then wash with PBS for 5 min.
- 7) Remove the slides from PBS and inactivate endogenous peroxidase with 3% H<sub>2</sub>O<sub>2</sub> in PBS for 10 min.
- 8) Wash the slides 2 times in PBS for 5 min. each.
- 9) Remove the slides from PBS, wipe gently around each section and cover tissues with blocking buffer [1% BSA/20 mM HEPES/135 mM NaCl (pH 7.4)] for 5 min. at room temperature to block non-specific staining. Do not wash.
- 10) Tip off the blocking buffer, wipe gently around each section and cover tissues with primary antibody diluted with the blocking buffer as suggested in the **APPLICATION**. (The concentration of antibody will depend on the conditions.) Incubate the sections for 1 hr. at room temperature.
- 11) Wash the slides 2 times in PBS for 5 min. each.
- 12) Wipe gently around each section and cover tissues with Histostar<sup>TM</sup> (Ms + Rb) (MBL; code no. 8460). Incubate for 30 min. at room temperature.
- 13) Wash the slides 2 times in PBS for 5 min. each.
- 14) Visualize by reacting for 5 min. with Histostar<sup>TM</sup> DAB Substrate Solution (MBL; code no. 8469). \*DAB is a suspect carcinogen and must be handled with care. Always wear gloves.
- 15) Wash the slides in water for 5 min.
- 16) Counterstain in hematoxylin for 1 min., wash the slides 3 times in water for 5 min. each, and then immerse the slides in PBS for 5 min.
- 17) Dehydrate by immersing in Ethanol 3 times for 3 min. each, followed by immersing in Xylene 3 times for 3 min. each. Now ready for mounting.

(Positive control for Immunohistochemistry; Human liver carcinoma)



*Immunohistochemical detection of Phospho-p62 (SQSTM1) (Ser351) in human liver carcinoma* 

Brown: Anti-Phospho-p62 (SQSTM1) (Ser351) mAb (M217-3) Blue: Hematoxylin

## **Immunocytochemistry**

- 1) Spread the cells on a glass slide, then incubate in a  $CO_2$  incubator for one night.
- 2) Remove the culture supernatant by careful aspiration.
- 3) Wash the slide 2 times with PBS.
- 4) Fix the cells with 4% paraformaldehyde (PFA)/PBS for 10 min. at room temperature (20~25°C).
- 5) Wash the slide 2 times with PBS.
- 6) Permeabilize the cells with 100  $\mu$ g/mL of Digitonin/PBS for 10 min. at room temperature.
- 7) Wash the slide 2 times with PBS.
- 8) Add 200 μL of the primary antibody diluted with PBS as suggested in the **APPLICATIONS** onto the cells. Incubate for 1 hr. at room temperature. (Optimization of antibody concentration or incubation condition is recommended if necessary.)
- 9) Wash the slide 2 times with PBS.
- Add 200 μL of 1:500 Alexa Fluor<sup>®</sup>488 anti-mouse IgG (Invitrogen; code no. A11001) diluted with PBS onto the cells. Incubate for 30 min. at room temperature. Keep out light by aluminum foil.
- 11) Wash the slide 2 times with PBS.
- 12) Counter stain with DAPI for 5 minutes at room temperature.
- 13) Wash the glass slide 2 times with PBS.
- 14) Wipe excess liquid off the slide but take care not to touch the cells. Never leave the cells to dry.
- 15) Promptly add mounting medium onto the slide, then put a cover slip on it.

(Positive control for Immunocytochemistry; huH-1)



*Immunocytochemical detection of Phospho-p62 (SQSTM1) (Ser351)* Green: Anti-Phospho-p62 (SQSTM1) (Ser351) mAb (M217-3) Blue: DAPI