

#### **MONOCLONAL ANTIBODY**

For research use only. Not for clinical diagnosis.

Catalog No. CTB-LC3-2-IC

# Anti LC3 (Clone: LC3·1703)

#### **BACKGROUND**

LC3B is one of the mammalian Atg8 homologs and widely used as an autop hagosome marker. Immediately after synthesis, LC3 is processed by Atg4 and becomes LC3-I. Upon induction of autophagy, the C-terminal glycine of LC3-I is conjugated to phosphatidylehanolamine, resulting in formation of membrane-bound LC3-II. Most LC3-II is thought be present on autophagosome membrane. The autophagosome subsequently fuses with a lysosome, where inside materials, including LC3-II, are degraded. The expression level of LC3-II generally correlates with the number of autophagosome.

**Product type** Primary antibodies

**Host** Mouse

Source

Form Liquid

Protein G purified

PBS (pH7.4) with 1% BSA and less than 0.1% NaN 3 as a preservative.

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Antigen Human Recombinant LC3

Clone LC3 · 1703 lsotype lgG1

**Application notes** ICC, Immuno EM

Recommended use

**Recommended dilutions** 

ICC: 1/100

Immuno EM: 1/10

Optimal dilutions/concentrations should be determined by the end user.

**Staining Pattern** 

Cross reactivity Human, Mouse

Si I I

**Storage** Store below -20° C (below -70° C for prolonged storage).

Aliquot to avoid cycles of freeze/thaw.

**References**1) Kabeya, Y., Mizushima, N., Ue no, T., Yamamoto, A., Kirisako, T., Noda, T., Komi nami, E., Ohsumi, Y. and Yoshimori, T. LC3, a mammalian homologue of yeast Apg8p, is localized in autophagosome membranes after

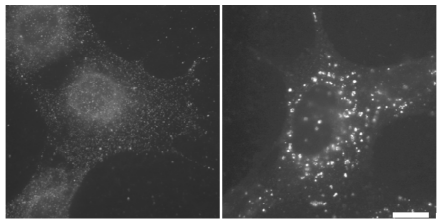
processing EMBO J. 19, 5720-5728. (2000)

2) Mizushima, N., Yoshimori, T. How to interpret LC3 immunoblotting Autophagy 3:542-545 (2007)

3) Mizushima, N., Yoshimori, T. and Levine, B. Me thods in mammalian autopha gy research. Cell 140; 313-326

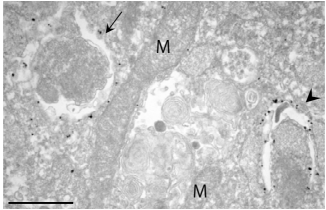
(2010)





(Fig 1) Immunofluorescence microscopy analysis of mouse embryonic fibroblasts (MEFs).

MEFs were cultured in regular DMEM supplemented with 10% FBS (left) or DMEM with out amino acids (right ) for 1 hr . Af ter fixation with 4% paraformaldehyde for 10 min at room temperature, they were permeabilized with 50  $\,\mu g/ml$  digitonin f or 5 mi n. Cell s were then subjected t o immunofluorescence mi croscopy u sing #1703 anti-LC3 a ntibody at 1:100 dilution. Scale bar, 20  $\,\mu m$ .



(Fig 2) Immuno-electron microscopy analysis of mouse embryonic fibroblasts (MEFs).

MEFs were culture d in DMEM without amino a cids for 2 hr. After fixation with 4% p araformaldehyde for 2 h r at room te mperature, they we re permeabilized with liquid nitrogen. Immuno -electron micro scopy analysis (pre-embedding method) of endogenous LC3 was performed using #1703 anti-LC3 antibody at 1:10 dilution. The gold labeling was intensified by using a silver e nhancement kit (HQ silver e nhancement kit, Nano probes, NY). Scale bar, 500 nm.

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