

Human/Mouse Myl9/12 (F6) rabbit mAb

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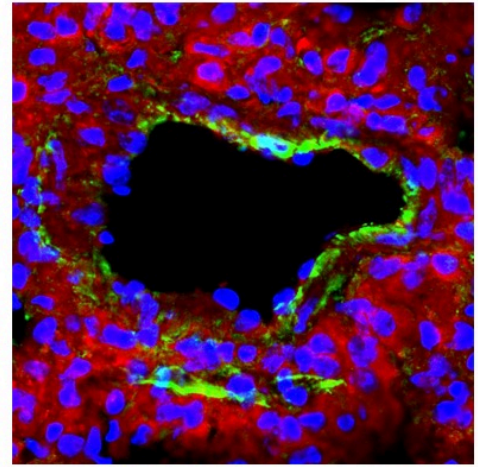
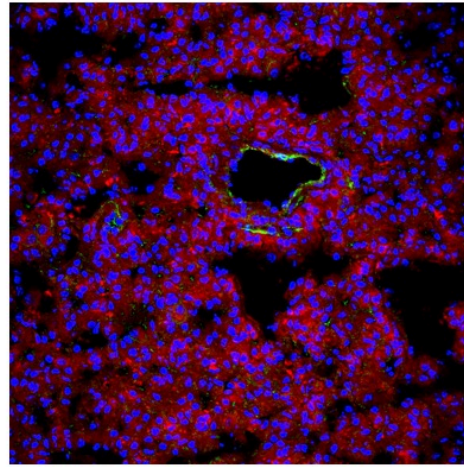
Store at: -20°C

For Research Use Only. Not For Use In Diagnostic Procedures.

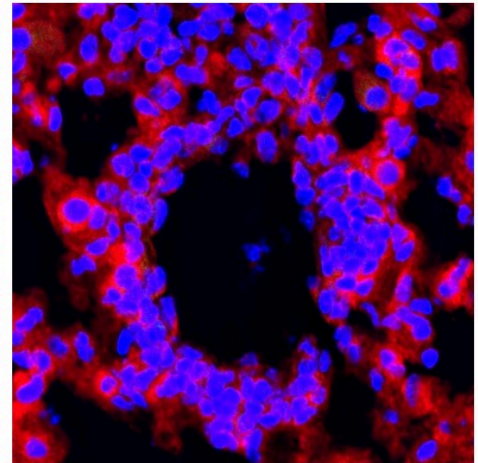
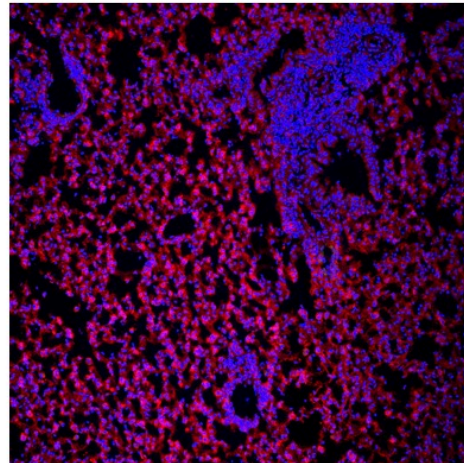
Applications	Detection	Clonality	Isotype
Functional Assay,IHC,ELISA	Anti-Rabbit IgG	Monoclonal	Rabbit IgGk

Format:	Unconjugated
Cross Reactivity:	Antibody may react with the same target protein from other species sharing the same sequence.
Formulation:	1X PBS, 0.02% NaN ₃ , 50% Glycerol, 0.1% BSA
Preparation:	Protein A+G
Reactivity:	Human,Mouse
Recommended Usage:	1µg/mL – 0.001µg/mL. It is recommended that the reagent be titrated for optimal performance for each application. See product image legends for additional information.
Immunogen:	N-terminal peptide of Myl9
Description:	<p>Myosin regulatory light chain (Myl) 9 is a regulatory subunit of the ATPase myosin protein. Myl9 regulates actin rearrangement to direct cellular migration, shape, and adhesion. Myl9 itself is regulated by post-translational modifications, including phosphorylation, acetylation and methylation. Phosphorylation of Myl9 at Thr18 and Ser19 promotes myosin ATPase activity and interaction with actin. Nα-acetylation of Myl9 has been shown to increase Ser19 phosphorylation and cytoplasmic activity, while Nα-methylation promotes DNA binding in the nucleus. Myl9, Myl12a, and Myl12b (Myl9/12) have been identified as functional ligands for CD69 in inflamed lungs, playing a major role in chronic inflammatory disorders such as chronic rhinosinusitis. Homozygous deletion in the MYL9 gene in humans has been identified as a putative molecular basis of the disease megacystis-microcolon-intestinal hypoperistalsis (MMIHS) syndrome, especially considering Myl9's role in contracting smooth muscle cell.</p>
References:	<p>Hayashizaki K, Kimura M, Tokoyoda K, et al. (2016) Science Immunology. 1: eaaf9154.</p> <p>Nevitt C, Tooley JG, and Tooley CES. (2018) Biochemical Journal. 475:3201-3219.</p> <p>Morena CA, Sobreira N, Pugh E, Zhang P, Steel G, Torres FR, and Cavalcanti DP. (2017) European Journal of Human Genetics. 26:669-675.</p>

F-6



Isotype



MyI9/12
CellMask
TO-PRO3

Anti-human/mouse MyI9/12 Abwiz antibody AWBMyI9F6 (Cat. #1151) shows strong and specific tissue staining by immunohistochemistry.