

FnCpf1/Cas12a (Strain *U112*) (E7I2B) Rabbit mAb



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Applications: W, IF-IC, FC-FP	Reactivity: All	Sensitivity: Transfected Only	MW (kDa): 152	Source/Isotype: Rabbit IgG	UniProt ID: #A0Q7Q2
Product Usage		Application			Dilution
Information		Western Blotting			1:1000
		Immunofluorescence (Imn	nunocytochemistry)		1:400
		Flow Cytometry (Fixed/Per	meabilized)		1:100
Storage		Supplied in 10 mM sodium HEPES (pH 7.5), 150 mM NaCl, 100 μg/ml BSA, 50% glycerol and less than 0.02% sodium azide. Store at –20°C. Do not aliquot the antibody.			
Specificity/Sensitivity		FnCpf1/Cas12a (Strain <i>U112</i>) (E7I2B) Rabbit mAb recognizes transfected levels of total FnCpf1/Cas12a (Strain <i>U112</i>) protein. This antibody does not cross-react with Cas9 (<i>S. pyogenes</i>), Cas9 (<i>S. aureus</i>), and AsCpf1/Cas12a (Strain <i>BV3L6</i>) proteins.			
Source / Purification		Monoclonal antibody is produced by immunizing animals with a synthetic peptide corresponding to residues surrounding Ile841 of Cpf1/Cas12a from <i>Francisella tularensis subsp. novicida</i> (Strain <i>U112</i>) protein.			
Background		CRISPR-Cas (clustered regularly interspaced short palindromic repeats and CRISPR-associated proteins) are RNA-guided nuclease effectors that are utilized for precise genome editing in mammalian systems (1). Cpf1/Cas12a (CRISPR from <i>Prevotella</i> and <i>Francisella</i>) proteins are members of the Class 2 CRISPR system (2). Class 2 CRISPR systems, such as the well characterized Cas9, rely on single-component effector proteins to mediate DNA interference (3). Cpf1/Cas12a endonucleases, compared to Cas9 systems, have several unique features that increase the utility of CRISPR-based genome editing techniques: 1) Cpf1/Cas12a-mediated cleavage relies on a single and short CRISPR RNA (crRNA) without the requirement of a trans-activating crRNA (tracrRNA), 2) Cpf1/Cas12a utilizes T-Rich protospacer-adjacent motif (PAM) sequences rather than a G-Rich PAM, and 3) Cpf1/Cas12a generates a staggered, rather than a blunt-ended, DNA double-stranded break (2). These features broaden the utility of using CRISPR-Cas systems for specific gene regulation and therapeutic applications. Several Cpf1/Cas12a bacterial orthologs have been characterized for CRISPR-mediated mammalian genome editing (2,4).~FnCpf1 (Strain <i>U112</i>)/Cas12a is a Cpf1/Cas12a enzyme derived from <i>Francisella novicida U112</i> (5).			
Background References		1. Cong, L. et al. (2013) <i>Science</i> 339, 819-23. 2. Zetsche, B. et al. (2015) <i>Cell</i> 163, 759-71. 3. Horvath, P. and Barrangou, R. (2010) <i>Science</i> 327, 167-70. 4. Zhang, Y. et al. (2017) <i>Sci Adv</i> 3, e1602814. 5. Zetsche, B. et al. (2015) <i>Cell</i> 163, 759-71.			
Species Reactivity	y	Species reactivity is determ	nined by testing in at	least one approved applic	cation (e.g., western blot).
Western Blot Buffer		IMPORTANT: For western blots, incubate membrane with diluted primary antibody in 5% w/v BSA, 1X			

TBS, 0.1% Tween® 20 at 4°C with gentle shaking, overnight.

Applications Key

W: Western Blotting IF-IC: Immunofluorescence (Immunocytochemistry) FC-FP: Flow Cytometry (Fixed/Permeabilized)

Cross-Reactivity Key

All: All Species Expected

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