## IRE1α (14C10) Rabbit mAb



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## For Research Use Only. Not for Use in Diagnostic Procedures.

<b>Applications:</b> W, W-S, IP	<b>Reactivity:</b> H M R	<b>Sensitivity:</b> Endogenous	<b>MW (kDa):</b> 130	<b>Source/Isotype:</b> Rabbit IgG	<b>UniProt ID:</b> #075460	Entrez-Gene Id: 2081
Product Usage Information	•	<b>Application</b> Western Blotting Simple Western™ Immunoprecipitation			<b>Dilution</b> 1:1000 1:10 - 1:50 1:50	
Storage		Supplied in 10 mM sodium HEPES (pH 7.5), 150 mM NaCl, 100 $\mu$ g/ml BSA, 50% glycerol and less than 0.02% sodium azide. Store at –20°C. Do not aliquot the antibody.				
Specificity/Sensitivity		IRE1 $\alpha$ (14C10) Rabbit mAb detects endogenous levels of total IRE1 $\alpha$ protein.				
Source / Purification		IRE1 $\alpha$ (14C10) Rabbit mAb is produced by immunizing rabbits with a synthetic peptide corresponding to residues surrounding His963 of human IRE1 $\alpha$ .				
Background		reticulum (ER) after the folded. Disruptions of I developed an adaptive protein folding (1). One transmembrane serine UPR that transmits the this kinase, IRE1a, was Upon activation of UPR mechanism using its elactivator that induces in the folder.	eir synthesis. Inside ER homeostasis lea mechanism called of the players in l /threonine kinase unfolded protein later identified an la, IRE1a splices X-b ndoribonuclease a many UPR respons mRNAs based on	membrane proteins tran e the ER, they are post-trads to the accumulation I unfolded protein respo JPR, IRE1, was first ident (2-4). This kinase was prosignal across the ER mer d shown to be ubiquitou ox binding protein (XBP) ctivity (6). This converts a ive genes (6). Recently, I the ER-localization and p in UPR (7).	anslationally modit of unfolded proteir nse (UPR) to counte ified in <i>Saccharom</i> oposed to be a pro nbrane (3,4). A hun sly expressed in hu I) mRNA by an unce KBP1 into a potent RE1a was shown to	ried and properly as (1). The ER has eract compromised yces cerevisiae as a ximal sensor for man homolog of aman tissues (5). Conventional transcriptional mediate the rapid
Background Re	eferences	<ol> <li>Kaufman, R.J. et al. (2002) Nat Rev Mol Cell Biol 3, 411-421.</li> <li>Nikawa, J. and Yamashita, S. (1992) Mol. Microbiol. 6, 1441-1446.</li> <li>Cox, J.S. et al. (1993) Cell 73, 1197-1206.</li> <li>Mori, K. et al. (1993) Cell 74, 743-756.</li> <li>Tirasophon, W. et al. (1998) Genes Dev. 12, 1812-1824.</li> <li>Lee, K. et al. (2002) Genes Dev. 16, 452-466.</li> <li>Hollien, J. and Weissman, J.S. (2006) Science 313, 104-107.</li> </ol>				

**Species Reactivity** Species reactivity is determined by testing in at least one approved application (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 **W-S:** Simple Western™ **IP:** Immunoprecipitation

**Cross-Reactivity Key** H: Human M: Mouse R: Rat

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