## AMPKα (23A3) Rabbit mAb



Orders: 877-616-CELL (2355)

orders@cellsignal.com

Support: 877-678-TECH (8324)

Web: info@cellsignal.com

cellsignal.com

3 Trask Lane | Danvers | Massachusetts | 01923 | USA

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

	eactivity: I M R Mk	<b>Sensitivity:</b> Endogenous	<b>MW (kDa):</b> 62	Source/Isotype: Rabbit IgG	<b>UniProt ID:</b> #Q13131, #P54646	Entrez-Gene Id 5562, 5563
Product Usage Information		<b>Application</b> Western Blotting			<b>Dilution</b> 1:1000	
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.				
		For a carrier free (BSA and azide free) version of this product see product #30127.				
Specificity/Sensitivity		AMPKa (23A3) Rabbit mAb detects endogenous levels of AMPKa protein. The antibody detects both the $\alpha 1$ and $\alpha 2$ isoforms of the catalytic subunit.				
Source / Purification		Monoclonal antibody is produced by immunizing animals with a synthetic peptide corresponding to the amino-terminal sequence of human AMPK $\alpha$ .				
Background		key role in the regular catalytic α subunit an genes (α1, 2; β1, 2; γ1 environmental stress, association with acces activation loop, and the phosphorylated at The significance of these part translationally modificance of the section of the secti	tion of energy home dregulatory β and ν , 2, 3) (2). The kinas such as heat shock assory proteins STRA his phosphorylation r258 and Ser485 (for bosphorylation event by myristoylation er182 (6,7). Phosphe e phosphorylation a subunits have been Bateman domains), ccumulation in head egulates the metabo	eostasis (1). AMPK is a a subunits, each of whe is activated by an elect, hypoxia, and ischem D and MO25, phosphois required for AMPK or α1; Ser491 for α2). The is have yet to be elunt and multi-site phosphorylation at Ser108 of at Ser24/25 and Ser18; identified, most of whe Mutations at these site or skeletal muscle (1) olism of fatty acids and	yeast to plants and anii heterotrimeric complexich is encoded by two divated AMP/ATP ratio dia (1). The tumor supprorylates AMPKα at Thr1 activation (3-5). AMPKα he upstream kinase and cidated (6). The β1 subhorylation including Sethe β1 subunit seems to affects AMPK localizatich are located in the pies lead to reduction of ,2). Accumulating evided glycogen, but also moys, as well as blood flow	c composed of a rethree distinct ue to cellular and essor LKB1, in 72 in the is also dependent of the biological unit is post-r24/25, Ser96, o be required for ion (7). Several utative AMP/ATP AMPK activity ence indicates dulates protein
Background Refere	nces	2. Carling, D. (2004) 7. 3. Hawley, S.A. et al. (' 4. Lizcano, J.M. et al. (5. Shaw, R.J. et al. (2006) Woods, A. et al. (2006)	(2004) J Cell Sci 117, 5479-87. 004) Trends Biochem Sci 29, 18-24. et al. (1996) J Biol Chem 271, 27879-87. et al. (2004) EMBO J 23, 833-43. al. (2004) Proc Natl Acad Sci USA 101, 3329-35. al. (2003) J Biol Chem 278, 28434-42. . et al. (2001) Biochem J 354, 275-83.			
Species Reactivity		Species reactivity is de	etermined by testin	n in at least one annro	ved application (e.g., w	estern blot)

**Species Reactivity** 

**Applications Key** 

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.

125, 51176 11156115 25 46 1 5 1116

W: Western Blotting

Cross-Reactivity Key H: Human M: Mouse R: Rat Mk: Monkey

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