at -	l2) Mouse r	nAb				<b>Cell Signaling</b> TECHNOLOGY®			
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For Research Use Only. Not for Use in Diagnostic Procedures.									
Applications: WB, IP	Reactivity: H M R Mk	Sensitivity: Endogenous	MW (kDa): 145 mature Met	Source/Isotype: Mouse IgG1	UniProt ID: #P08581	Entrez-Gene Id: 4233			

	H M K MK Endogenous	beta-subunit. 170 pro-Met.	Mouse 1981	#P00501	4233			
Product Usage Information	Application Western Blotting Immunoprecipitation			<b>Dilution</b> 1:1000 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 / Sensitiv	/ity Met (25H2) Mouse m/	Met (25H2) Mouse mAb detects endogenous levels of Met protein.						
Source / Purification		Monoclonal antibody is produced by immunizing animals with a synthetic peptide corresponding to residues surrounding Tyr1234 of human Met.						
Background	factor) is a disulfide-lir and the amino-termina spans the plasma mer Met with HGF results i signaling components for all of the biological Tyr1003 is essential fo the Met kinase domair domain provides a dire and/or tyrosine kinase	Met, a high affinity tyrosine kinase receptor for hepatocyte growth factor (HGF, also known as scatter factor) is a disulfide-linked heterodimer made of 45 kDa $\alpha$ - and 145 kDa $\beta$ -subunits (1,2). The $\alpha$ -subunit and the amino-terminal region of the $\beta$ -subunit form the extracellular domain. The remainder of the $\beta$ -chain spans the plasma membrane and contains a cytoplasmic region with tyrosine kinase activity. Interaction of Met with HGF results in autophosphorylation at multiple tyrosines, which recruit several downstream signaling components, including Gab1, c-Cbl, and Pl3 kinase (3). These fundamental events are important for all of the biological functions involving Met kinase activity. The addition of a phosphate at cytoplasmic Tyr1003 is essential for Met protein ubiquitination and degradation (4). Phosphorylation at Tyr1234/1235 in the Met kinase domain is critical for kinase activation. Phosphorylation at Tyr1349 in the Met cytoplasmic domain provides a direct binding site for Gab1 (5). Research studies have shown that altered Met levels and/or tyrosine kinase activities are found in several types of tumors, including renal, colon, and breast. Thus, investigators have concluded that Met is an attractive potential cancer therapeutic and diagnostic target (6,7).						
Background Refere	<b>nces</b> 1. Cooper, C.S. et al. ( 2. Bottaro, D.P. et al. ( 3. Bardelli, A. et al. (19 4. Taher, T.E. et al. (20 5. Schaeper, U. et al. ( 6. Eder, J.P. et al. (200 7. Sattler, M. and Salg	1991) Science 251, 8 997) Oncogene 15, 3 002) J Immunol 169, (2000) J Cell Biol 149 09) Clin Cancer Res	302-4. 103-11. 3793-800. 9, 1419-32. 15, 2207-14.	.118.				
Species Reactivity	Species reactivity is de	termined by testing in	n at least one approv	ved application (e.g., wes	tern blot).			
Western Blot Buffer	Western Blot Buffer IMPORTANT: For western blots, incubate membrane with diluted primary antibody in 5% w/v nonfa milk, 1X TBS, 0.1% Tween® 20 at 4°C with gentle shaking, overnight.							
Applications Key	WB: Western Blotting	IP: Immunoprecipita	tion					
Cross-Reactivity Ke	X: Xenopus Z: zebrafis	nouse <b>R:</b> rat <b>Hm:</b> hamster <b>Mk:</b> monkey <b>Vir:</b> virus <b>Mi:</b> mink <b>C:</b> chicken <b>Dm:</b> D. melanogaster zebrafish <b>B:</b> bovine <b>Dg:</b> dog <b>Pg:</b> pig <b>Sc:</b> S. cerevisiae <b>Ce:</b> C. elegans <b>Hr:</b> horse g <b>Rab:</b> rabbit <b>All:</b> all species expected						
Trademarks and Patents		Cell Signaling Technology is a trademark of Cell Signaling Technology, Inc. All other trademarks are the property of their respective owners. Visit cellsignal.com/trademarks for more information.						

## Met (25H2) Mouse mAb (#3127) Datasheet Without Images Cell Signaling Technology

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