Phospho-MET (Tyr1234/1235) (6F11) rabbit mAb

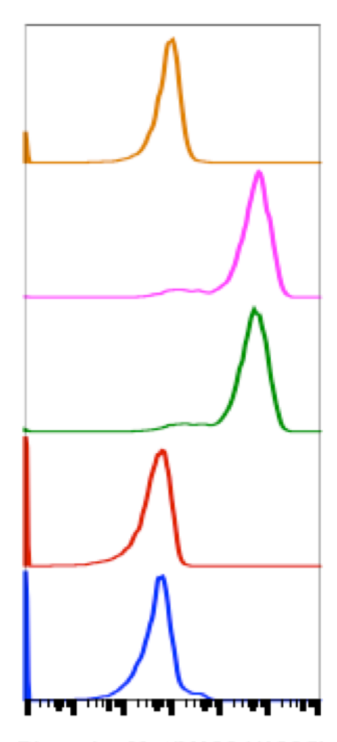
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For Research Use Only. Not For Use In Diagnostic Procedures.

Applications Flow Cytometry	Detection Anti-Rabbit IgG	Clonality Monoclonal	Isotype Rabbit IgGk
Format:	Unconjugated		
Cross Reactivity:	Predicted to work with mouse, rat and other homologues.		
Formulation:	1X PBS, 0.02% NaN3, 50% Glycerol, 0.1% BSA		
Preparation:	Protein A+G		
Reactivity:	Human,Mouse		
Recommended Usage:	$1\mu g/mL$ – $0.001\mu g/mL$. It is recommended that the reagent be titrated for optimal performance for each application. See product image legends for additional information.		
Immunogen:	A synthetic phospho-peptide corresponder Tyr1234/Tyr1235 of human phospho		nding
Description:	c-Met, also called tyrosine-protein kinase MET or hepatocyte growth factor receptor (HGFR), has tyrosine kinase activity (1). MET is a single pass tyrosine kinase receptor essential for embryonic development, organogenesis and wound healing. Normally, MET is expressed only in stems cells and progenitor cells but excessive expression of MET/HGFR and its autocrine activation by co-expression of hepatocyte growth factor (HGF) ligand are implicated in oncogenesis (2,3). Aberrantly activated MET leads to tumor growth, angiogenesis, and cancer metastasis and is correlated with poor prognosis. Abnormal activation of MET is observed in various human malignancies, such as kidney, liver, stomach, breast, and brain. MET activation by HGF induces MET kinase catalytic activity and leads to phosphorylation at Tyr 1234 and Tyr 1235.		
References:	Cooper CS (January 1992). "The met of transmembrane receptor for hepatocy Johnson M, Koukoulis G, Kochhar K, Ku "Selective tumorigenesis in non-parent hepatocyte growth factor transfection Johnson ME, Volpert O, Iyer AP (1995). transformation in NIH-3T3 cells transferactors. 12 (4): 303–13.	te growth factor". Oncog bo C, Nakamura T, lyer i chymal liver epithelial co ". Cancer Letters. 96 (1): "Evidence for autocrine	gene. 7 (1): 3-7. A (Sep 1995). ell lines by : 37-48. Kochhar KS, s basis of



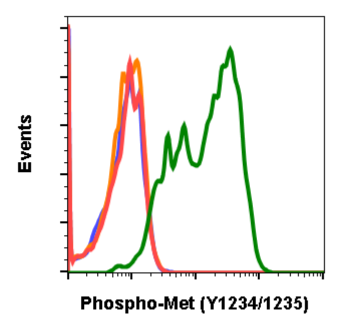


Phospho Met(Y1234/1235)

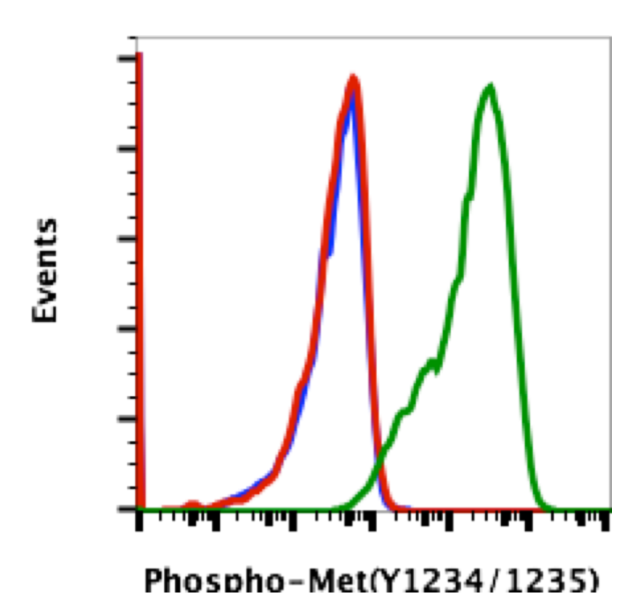
\$WELLID	Treatment	Median : BL1-A
F11+PP	FBS+PV	839
F11+NP	FBS+PV	49352
F11 0.01 ug/mL	FBS+PV	43691
F11 0.01 ug/mL	Ctrl	429
2'Ab	Ctrl	478

Flow cytometric analysis of Ramos cells secondary antibody only negative control (blue) untreated (red) treated with FBS + pervanadate (green) treated plus blocked by non-phospho peptide (violet) or treated plus blocked by phospho-peptide (brown) using 0.01 ug/mL Phospho-MET(Y1234/1235) antibody METY12341235-6F11 0.01 ug/mL. Cat. #2216.

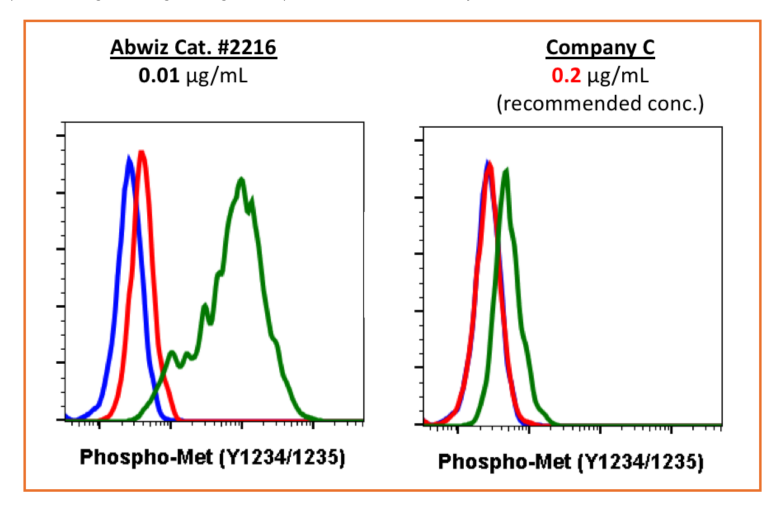




Flow cytometric analysis of NIH3T3 cells secondary antibody only negative control (blue) or $0.1~\mu g/mL$ of isotype control Cat. #2141 (orange) or treated with imatinib (red) or with pervanadate (green) using Phospho-MET(Y1234/1235) antibody METY12341235-6F11 at $0.1~\mu g/mL$. Cat #2216.



Flow cytometric analysis of Ramos cells secondary antibody only negative control (blue) or untreated (red) or treated with pervanadate (green) using 0.005 ug/mL Phospho-MET(Y1234/1235) antibody METY12341235-6F11. Cat. # 2216.



Flow cytometric analysis of Ramos cells secondary only negative control (blue) or untreated (red) or treated with FBS + pervanadate (green) using Phospho-Met (Tyr1234/1235) antibody MetY12341235-6F11 (Abwiz Cat. #2216) at 0.01ug/mL or Company C antibody at 0.2ug/mL (manufacturer's recommended concentration).