Phospho-MEK1/2 (Ser221) (D3) rabbit mAb APC Conjugate

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

Flow Cytometry N/A Monoclonal Rabbit IgGk	Applications	Detection	Clonality	Isotype
Flow Cytometry N/A Monocional Nappicing Control of the Cytometry	Flow Cytometry	N/A	Monoclonal	Rabbit IgGk

Format: APC

Cross Reactivity: Predicted to work with mouse, rat and other homologues.

Formulation: 1X PBS, 0.09% NaN3, 0.2% BSA

Preparation: Protein A+G

Reactivity: Human, Mouse

Recommended

Usage:

For flow cytometric staining, the suggested use of this reagent is 5 μ L per million cells or 5 μ L per 100 μ L of staining volume. 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 corresponding to residues surrounding Ser221 of

human phospho MEK1/2.

Description: Mitogen-activated protein kinase (MAPK) is the main building block of the

intracellular signaling network. The signals are initiated by activation of a small G protein (e.g., Ras), followed by a sequential activation of several sets of protein kinases. The extracellular signal-regulated kinase (ERK) pathway is stimulated by a large number of extracellular stimuli as well as various internal processes. This cascade regulates processes related to proliferation, differentiation, development, and oncogenic transformation. The signaling is usually initially started by activation of small G proteins (e.g. Ras), which in turn recruits to the membrane and activates the MAP3K (Raf kinase). The cascade leads to the activation of MAP kinase kinases

(MEKs). MEKs are phosphorylated at Ser218 and Ser222 in MEK1. The

phosphorylation activates MEKs. Phosphorylation of MEK at other sites regulates its activity as well. For example, phosphorylation at Ser386 by ERKs may either inhibit

ERKs activity or under certain condition, facilitate the activation of ERKs by

enhancing the MEK1 binding to Grb10 scaffold protein. Phosphorylation of MEK1 at Ser298 by p21-activating protein (PAK1) may lead to its activation whereas this can be inhibited by a feedback phosphorylation of MEK1 at Thr292 by ERKs. Protein Ser/Thr phosphatase inactivates MEK by dephosphorylation of pSer218 and

pSer222. Other phosphatases my also regulate MEK activity. Upon activation, MEKs behaves as a dual kinase and phosphorylate key Tyr and Thr residues of ERKs, thus

causing their activation.

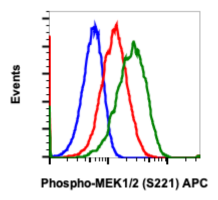
References: Raman M. and Cobb MH., (2003) Curr Biol 13:R886-888.

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Flow cytometric analysis of HeLa cells treated with imatinib and unstained as negative control (blue) or treated with imatinib (red) or with pervanadate (green) and stained using Phospho-MEK1/2 (S221) antibody MEK12S221-D3 APC conjugate. Cat. #2279.