Recombinant Mouse DDR1 Kinase/MCK10 Protein (His &GST Tag)



Catalog Number: PKSM040294

Note: Centrifuge before opening to ensure complete recovery of vial contents.

Description

Synonyms 6030432F18;AI323681;Cak;CD167a;Nep;PTK3A

Species Mouse

Expression Host Baculovirus-Insect Cells

Sequence Leu444-Val874

AccessionQ03146-2Calculated Molecular Weight75.8 kDaObserved molecular weight68 kDaTagN-His-GST

Bioactivity The specific activity was determined to be 2 nmol/min/mg using synthetic modified

AXLtide peptide (modified-CKKSRGDYMTMQIG) as substrate.

Properties

Purity > 85 % as determined by reducing SDS-PAGE.

Endotoxin < 1.0 EU per µg of the protein as determined by the LAL method.

Storage Store at $< -20^{\circ}$ C, stable for 6 months. Please minimize freeze-thaw cycles.

Shipping This product is provided as liquid. It is shipped at frozen temperature with blue

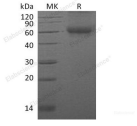
ice/gel packs. Upon receipt, store it immediately at < - 20°C.

Formulation Supplied as sterile solution of 20mM Tris, 500mM NaCl, pH 7.4, 10% glycerol,

2mM DTT

Reconstitution Not Applicable

Data



> 85 % as determined by reducing SDS-PAGE.

Background

Discoidin domain receptor family, member 1 (DDR1), also known as or CD167a (cluster of differentiation 167a), and Mammary carcinoma kinase 10 (MCK10), belongs to a subfamily of tyrosine kinase receptors with an extracellular domain homologous to Dictyostellium discoideum protein discoidin 1. Receptor tyrosine kinases play a key role in the communication of cells with their microenvironment. These kinases are involved in the regulation of cell growth, differentiation and metabolism. Expression of DDR1/MCK10/CD167 is restricted to epithelial cells, particularly in the kidney, lung, gastrointestinal tract, and brain. In addition, it has been shown to be significantly overexpressed in several human tumors. DDR1/MCK10/CD167 plays an important role in regulating attachment to collagen, chemotaxis, proliferation, and MMP production in smooth muscle cells. DDR1 functions in a feedforward loop to increase p53 levels and at least some of its effectors. Inhibition of DDR1 function resulted in strikingly increased apoptosis of wild-type

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p53-containing cells in response to genotoxic stress through a caspase-dependent pathway.

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