# MARK3 [6His-tagged]

Kinase

Alternate Names: MAP/Microtubule Affinity-regulating Kinase 3; MARK3; Microtubule-associated Protein/Microtubule Affinity-regulating Kinase 3. CDC25C-associated Protein Kinase 1; CTAK1

Cat. No.	66-0004-050
Lot. No.	2135

Quantity: 50 µg Storage: -70°C

FOR RESEARCH USE ONLY

NOT FOR USE IN HUMANS



### **CERTIFICATE OF ANALYSIS Page 1 of 2**

Protein Sequence: Please see page 2

### Background by Sir Philip Cohen

Protein ubiquitylation and protein phosphorylation are the two major mechanisms that regulate the functions of proteins in eukaryotic cells. However, these different posttranslational modifications do not operate independently of one another, but are frequently interlinked to enable biological processes to be controlled in a more complex and sophisticated manner. Studying how protein phosphorylation events control the ubiquitin system and how ubiquitylation regulates protein phosphorylation has become a focal point of the study of cell requlation and human disease. Cloning of human Microtubule Affinity Regulating Kinase 3 (MARK3) was first described by Peng et al. (1998). MARK3 is a member of the subfamily of protein kinases that include the AMP-activated protein kinase (AMPK) and, like AMPK itself, is activated by the tumour suppressor kinase LKB1 (Lizcano et al., 2004). The physiological roles of the MARK subfamily of protein kinases include the phosphorylation of microtubule associated proteins and the requlation of cell polarity. Members of the MARK sub-family also phosphorylate tau at sites that induce its dissociation from tubulin. Enhanced phosphorylation of these sites is an early hall mark of Alzeheimer's disease and is followed by abnormal aggregation of tau to paired helical filaments that are found in people with Alzheimer's disease (Marx et al., 2010). MARK3 con-

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## **Physical Characteristics**

Species: human

Source: Sf21 insect cell-baculovirus expression

Quantity: 50 µg

Concentration: 1 mg/ml

Formulation: 50 mM Tris/HCl pH 7.5, 0.1 mM EGTA, 150 mM NaCl, 270 mM sucrose, 0.03% Brij, 0.1%  $\beta$ -Mercapto-ethanol, 1 mM Benzamidine, 0.2 mM PMSF

Molecular Weight: ~86.1 kDa

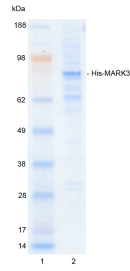
Purity: >70% by InstantBlue™ SDS-PAGE

**Stability/Storage:** 12 months at -70°C; aliquot as required

## **Quality Assurance**

### **Purity:**

4-12<sup>%</sup> gradient SDS-PAGE InstantBlue™ staining Lane 1: MW markers Lane 2: 1 μg His-MARK3



Protein Identification:

Confirmed by mass spectrometry.

### Activity Assay:

The specific activity of His-MARK3 was determined using the method described by Hastie *et al.* (2006) with the enzyme being assayed at several concentrations. His-MARK3 was incubated for 10 minutes at 30°C in kinase reaction buffer in the presence of CHKtide substrate (300  $\mu$ M) and [ $\gamma$ -<sup>32</sup>P]ATP (100  $\mu$ M). Duplicate reactions were stopped by spotting the assay mixture onto Whatman P81 paper – capturing the phosphorylated substrate. The radioactivity incorporated was measured on a scintillation counter and the enzyme's mean specific activity was calculated.

His-MARK3 specific activity:

282.0 Units/mg (282.0 Units/ml)

1 Unit = 1 nmole of phosphate incorporated into the substrate in 1 minute

Substrate: CHKtide (KKKVSRSGLYRSPSMPENLNRPR)



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Lot-specific COA version tracker: v1.0.0

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## **Physical Characteristics**

### Continued from page 1

Background

tains a ubiquitin-like domain adjacent to the kinase catalytic domain and, similar to MARK4, undergoes Lys29/ Lys33-linked polyubiquitylation that may inhibit its activity. It may also interact with the deubiquitylase USP9X (Al-Hakim *et al.*, 2008).

by Sir Philip Cohen

### **References:**

Al-Hakim AK, Zagorska A, Chapman L, Deak M, Peggie M, Alessi DR (2008) Control of AMPK-related kinases by USP9X and atypical Lys(29)/Lys(33)-linked polyubiquitin chains. *Biochem J* 411, 249-60.

Hastie CJ, McLauchlan HJ, Cohen P (2006) Assay of protein kinases using radiolabeled ATP: a protocol. *Nat Protoc* **1**, 968-71.

Lizcano JM, Goransson O, et al. (2004) LKB1 is a master kinase that activates 13 kinases of the AMPK subfamily, including MARK/PAR-1. EMBO J 23, 833-43.

Marx A, Nugoor C, Panneerselvam S, Mandelkow E (2010) Structure and function of polarity-inducing kinase family MARK/ Par-1 within the branch of AMPK/Snf1-related kinases. *FASEB* J 24, 1637-48.

Peng CY, Graves PR, Ogg S, Thoma RS, Byrnes MJ, 3rd, Wu Z, Stephenson MT, Piwnica-Worms H (1998) C-TAK1 protein kinase phosphorylates human Cdc25C on serine 216 and promotes 14-3-3 protein binding. *Cell Growth Differ* **9**, 197-208.

Background kindly written by:

Sir Philip Cohen FRS, FRSE University of Dundee

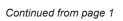
Director of the Medical Research Council Protein Phosphorylation Unit (1990-2012)

Director of the Scottish Institute for Cell Signalling incorporating the Protein Ubiquitylation Unit (2008-2012)

Co-Director of the Division of Signal Transduction Therapy (1998-2012)

Deputy Director of the Division of Signal Transduction Therapy (from July 2012)

Professor Cohen's research group is studying the interplay between protein phosphorylation and protein ubiquitylation in the regulation of innate immunity.



### **Protein Sequence:**

 $\texttt{MSYYHHHHHHDYDIPTT} \underline{\texttt{ENLYFQG}} \texttt{AMGS}$ GIQRPTSTSSLVAAA**S**TRTPLPTVNERD TENHTSHGDGRQEVTSRTSRSGARCRN SIASCADEQPHIGNYRLLKTIGKGN FAKVKLARHILTGREVAIKIIDKTQLNPT SLQKLFREVRIMKILNHPNIVKLFEVI ETEKTLYLIMEYASGGEVFDYLVAHGRM **KEKEARSKFROIVSAVOYCHOKRIVHRDL** KAENLLLDADMNIKIADFGFSNEFTVG GKLDTFCGSPPYAAPELFQGKKYDGPE VDVWSLGVILYTLVSGSLPFDGONLKEL RERVLRGKYRIPFYMSTDCENLLKRFLV LNPIKRGTLEQIMKDRWINAGHEE DELKPFVEPELDISDQKRIDIMVGM GYSQEEIQESLSKMKYDEITATYLLL GRKSSELDASDSSSSSNLSLAKVRPSS DLNNSTGQSPHHKVQRSVSSSQKQRRYSD HAGPAIPSVVAYPKRSQTSTADGDLKEDG ISSRKSSGSAVGGKGIAPASPMLG NASNPNKADIPERKKSSTVPSSNTAS GGMTRRNTYVCSERTTADRHSVIONG **KENSTIPDQRTPVASTHSISSAATP** DRIRFPRGTASRSTFHGQPRERRTATYN GPPASPSLSHEATPLSQTRSRGSTNLF SKLTSKLTRSRNVSAEQKDENKEAK PRSLRFTWSMKTTSSMDPGDMMREIRKV LDANNCDYEQRERFLLFCVHGDGHAEN LVQWEMEVCKLPRLSLNGVRFKRISGT SIAFKNIASKIANELKL

Tag (**bold text**): N-terminal His Protease cleavage site: TEV (<u>ENLYF ▼QG</u>) MARK3 (regular text): Start **bold italics** (amino acid residues 2-729) Accession number: AAC15093.1

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