PINK1 [MBP-tagged]

Kinase

Alternate Name: PTEN-Induced Putative Kinase 1

Cat. No.	66-0043-050
Lot. No.	30342

Quantity: 50 µg Storage: -70°C

NOT FOR USE IN HUMANS

FOR RESEARCH USE ONLY

Protein ubiquitylation and protein phos-

phorylation are two major post-trans-

lational modifications that regulate the functions of proteins in eukaryotic

cells. However, these 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 regulation and human disease.

Cloning of PTEN Induced putative Ki-

nase 1 (PINK1) was first described by Unoki and Nakamura *et al.* (2001).

PINK1 is a mitochondrial serine/threo-

nine kinase involved in the normal

function and integrity of mitochondria,

PINK1 reduces neuronal apoptosis

through a reduction in cytochrome c

release from mitochondria and sub-

sequent activation of caspase 3 (Petit et al., 2005). PINK1 has been shown

to phosphorylate Parkin at Ser65 - lo-

cated in its Ubl domain - which leads

to a marked activation in the activity of

the E3 ligase (Kondapalli et al., 2012).

PINK1 activation of Parkin catalyses

K63-linked polyubiquitylation and en-

hances parkin-mediated ubiquitin sig-

nalling through the I-kappa-B kinase/

nuclear factor kappa-B (NF-kappa-B)

pathway. It is thought that deregula-

tion of this pathway through Parkin-

son's Disease (PD)-linked mutations

in PINK1 is the cause of PD patho-

genesis (Sha et al., 2010). PINK1

Background

Physical Characteristics

Species: Tribolium castaneum

Source: E. coli

Quantity: 50 µg

Concentration: 1.9 mg/ml

Formulation: 50 mM Hepes pH 7.5, 150 mM NaCl, 2 mM DTT, 10% glycerol

Quality Assurance

Purity:

4-12% gradient SDS-PAGE InstantBlue™ staining Lane 1: MW markers Lane 2: 1 μg MBP-PINK1 Purity: >85% by InstantBlue™ SDS-PAGE

Molecular Weight: ~108.1 kDa

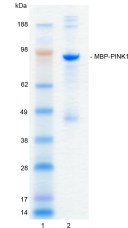
CERTIFICATE OF ANALYSIS Page 1 of 2

UBIQUIGEN

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

Protein Sequence: Please see page 2

Protein Identification: Confirmed by mass spectrometry.



0 0.5 1.0 2.0 2.0 MBP-PINK1 (D359A) (µg) 0 0.5 1.0 2.0 2.0 MBP-PINK1 WT (µg) kDa kDa 188 - MBP-PINK1 - Ub 188 = - MBP-PINK1 - Ub anti-PINK1 98 98 MBP-PINK1 - MBP-PINK1 62 62 anti-Parkin 2 3 Lanes 9 10

Kinase activity assay:

The activity of MBP-PINK1 was validated through its ability to phosphorylate Parkin thus activating and enabling Parkin-catalysed generation of MBP-PINK1-ubiguitin conjugates. MBP-PINK1 (0, 0.5, 1.0 and 2.0 µg) or MBP-PINK1 (D359A) (0, 0.5, 1.0 and 2.0 µg) were incubated in kinase assay buffer with Parkin (2.0 µg) in the presence or absence of ATP for 60 minutes at 30°C. The ubiquitylation reactions were then initiated through the addition His-UBE1, the E2 conjugating enzyme His-UBE2L3 (UbcH7) and ubiquitin and incubated for a further 60 minutes at 30°C. MBP-PINK1-ubiquitin conjugates were identified by Western blotting using an anti-PINK1 antibody (lanes 3, 4 and 9) which were observed in the presence of ATP and WT MBP-PINK1 predominantly over those observed in the presence of ATP and the kinase dead MBP-PINK1 (D359A) mutant (compare lanes 4 and 9 respectively). Ubiguitin conjugates were not observed in the absence of MBP-PINK1 (lanes 1 and 6) or ATP (lanes 5 and 10).

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Lot-specific COA version tracker: v1.0.0

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CERTIFICATE OF ANALYSIS Page 2 of 2

Background

Continued from page 1

controls Parkin E3 ligase activity not only by phosphorylating Parkin, but also by phosphorylating ubiquitin both at Ser65. It is thought that phosphorylation of Parkin serves to prime the E3 ligase enzyme for activation by ubiquitin (pSer65) (Kazlauskaite et al. 2014). USP30 (a deubiquitylase (DUB) localized to mitochondria) antagonizes mitophagy driven by Parkin and PINK1. Parkin ubiquitylates and tags damaged mitochondria for clearance. USP30 removes ubiquitin attached by Parkin onto damaged mitochondria and blocks Parkin's ability to drive mitophagy. Thus USP30 inhibition is potentially beneficial in Parkinson's disease by promoting mitochondrial clearance and guality control (Bingol et al. 2014).

References:

Bingol B, Tea JS, Phu L, Reichelt M, Bakalarski CE, Song Q *et al.* (2014) The mitochondrial deubiquitinase USP30 opposes parkinmediated mitophagy. *Nature* **510**, 370-5.

Kazlauskaite A, Kondapalli C, Gourlay R, Campbell DG, Ritorto MS, Hofmann K et al. (2014) Parkin is activated by PINK1-dependent phosphorylation of ubiquitin at Ser65. *Biochem J* **460**, 127-139.

Kondapalli C, Kazlauskaite A, Zhang N, Woodroof HI, Campbell DG, Gourlay R, Burchell L, Walden H, MacCartney TJ, Deak M, Knebel A, Alessi DR and Muqit MM (2012) PINK1 is activated by mitochondrial membrane potential depolarization and stimulates PARKIN E3 ligase activity by phosphorylating Serine 65. *Open Biology* **5**, 120080.

Petit A, Kawarai T, Paitel E, Sanjo N, Maj M, Scheid M, Chen F, Gu Y, Hasegawa H, Salehi-Rad S, Wang L, Rogaeva E, Fraser P, Robinson B, St George-Hyslop P, Tandon A (2005) Wild-type PINK1 prevents basal and induced neuronal apoptosis, a protective effect abrogated by Parkinson disease-related mutations. *J Biol Chem* **280**, 34025-34032.

Sha D, Chin LS, Li L (2010) Phosphorylation of parkin by Parkinson disease-linked kinase PINK1 activates parkin E3 ligase function and NF-kappa-B signaling. *Hum Molec Genet* **19**, 352-363.

Unoki M, Nakamura, Y (2001) Growth-suppressive effects of BPOZ and EGR2, two genes involved in the PTEN signaling pathway. *Oncogene* **20**, 4457-4465.

Physical Characteristics

Continued from page 1

Protein Sequence: MKIEEGKLVIWINGDKGYNGLAEVGKKFEKDT GIKVTVEHPDKLEEKFPQVAATGDGPDIIF WAHDRFGGYAQSGLLAEITPDKAFQDKLYP FTWDAVRYNGKLIAYPIAVEALSLIYNKDLLP **NPPKTWEEIPALDKELKAKGKSALMFNLQEPY FTWPLIAADGGYAFKYENGKYDIKDVGVDNA** GAKAGLTFLVDLIKNKHMNADTDYSIAEAAF NKGETAMTINGPWAWSNIDTSKVNYGVTVLPT **FKGOPSKPFVGVLSAGINAASPNKELAKE FLENYLLTDEGLEAVNKDKPLGAVALKSY EEELVKDPRIAATMENAOKGEIMPNIPOMSAF WYAVRTAVINAASGRQTVDEALKDAQTNS** SSNNNNNNNNLGDDDDKVPEFLEVLFQG PGSMSVRAVGSRLFKHGRSLIQQFCKRDLNT TIGDKINAVSQATAAPSSLPKTQIPKNFAL RNVGVOLGLOARRILIDNVLNRVTNSL SAELRKKATRRILFGDSAPFFALVGVSIAS **GTGILTKEEELEGVCWEIREAISKIKWOYY** DIDESRFESNPITLNDLSLGKPIAKGTNGV VYSAKVKDDETDDNKYPFALKMMFNYDIQSNS MEILKAMYRETVPARMYYSNHDLNNWEIELAN RRKHLPPHPNIVAIFSVFTDLIQELEGSKD LYPAALPPRLHPEGEGRNMSLFLLMKRYDCN LQSFLSTAPSTRTSLLLLAQLLEGVAHMTAH GIAHRDLKSDNLLLDTSEPESPILVISDFGC CLADKTNGLSLPYTSYEMDKGGNTALMAPEI ICQKPGTFSVLNYSKADLWAVGAIAYEIF NCHNPFYGPSRLKNFNYKEGDLPKLPDEVPT VIQALVANLLKRNPNKRLDPEVAANVCQLFL WAPSTWLKPGLKVPTSGEILQWLLSLTTKVL CEGKINNKSFGEKFTRNWRRTYPEYLLISSFL CRAKLANVRNALHWIQENLPELD

Tag (**bold text**): N-terminal MBP Protease cleavage site: PreScission[™] (<u>LEVLFQ▼GP</u>) PINK1 (regular text): Start **bold italics** (amino acid residues 1-570) Accession number: XP_968367.1



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