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Certificate of Analysis

PI3 Kinase (p120 γ)

(Recombinant enzyme expressed in Sf21 insect cells)

Catalogue # 14-558

Lot # 31017U

Product Description: N-terminal 6His-tagged recombinant full-length human PI3 kinase (p120 γ) expressed by baculovirus in Sf21 insect cells. Purified using Ni²⁺/NTA-agarose. Purity 74% by SDS-PAGE and Coomassie blue staining. MW = 130kDa.

Specific Activity (lot# 31017U): 39U/mg, where one unit of PI3 Kinase (p120 γ) activity is defined as 1nmol phosphatidylinositol 3,4,5-trisphosphate formed per minute at 22°C with a final ATP concentration of 100 μ M.

Formulation: 20 μ g of enzyme in 8.2 μ l of 50mM Tris/HCl pH7.5, 150mM NaCl, 10% glycerol, 0.1mM EGTA, 2mM DTT.

Storage and Stability: Store at -70°C from date of shipment. For maximum recovery of product, centrifuge original vial prior to removing the cap.

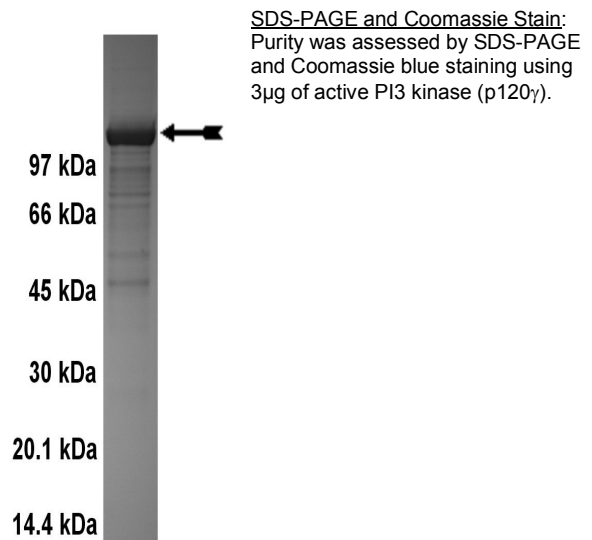
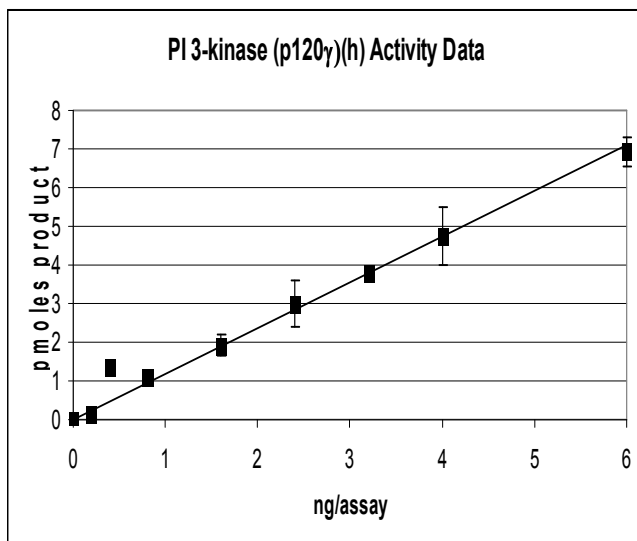
Handling Recommendations: Rapidly thaw the vial under cold water and immediately place on ice. Aliquot unused material into pre-chilled microcentrifuge tubes and immediately snap-freeze the vials in liquid nitrogen prior to re-storage at -70°C.

**FOR IN VITRO RESEARCH USE ONLY
NOT FOR USE IN HUMANS OR ANIMALS**

Quality Control Testing

Kinase Assay: 0–6ng of this lot of PI3 Kinase (p120 γ) was assayed in a 30 minute reaction using the PIPProfiler™ protocol (see <http://www.upstate.com/features/pipprofiler>).

MS Tryptic Fingerprint: Confirmed identity as PI3 Kinase (p120 γ) with 41% amino acid coverage of the translated sequence listed on page three.



P13 Kinase (p120 γ) Sequence Information

Protein	human P13 Kinase (p120 γ)
Tags	N-terminal 6His
Native sequence	M34 of the recombinant protein is equivalent to M1 of human P13 Kinase (p120 γ)
Accession number	GenBank AF327656. The recombinant protein contains the amino acid substitution R459Q with respect to this accession number. This conflict is reported in GenBank XM_027912. The residue coordinates in the native sequence are given.

Recombinant P13 Kinase (p120 γ) amino acid sequence:

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1 MSYYHHHHHH DYDIPTTENL YFQGAMGSGI RPCMELENYK QPVVLRDNC RRRRRMKPRS
61 AAASLSSMEL IPIEFVLPTS QRKCKSPETA LLHVAGHGNV EQMKAQVWLR ALETSSVAADF
121 YHRLGPHHFL LLYQKKGQWY EIYDKYQVVQ TLDCLRYWKA THRSFGQIHL VQRHPPSEES
181 QAFQRQLTAL IGYDVTDVSN VHDDELEFTR RGLVTPRMAE VASRDPKLYA MHPWVTSKPL
241 PEYLWKKIAN NCIFIVIHRS TTSQTIKVSP DDTFGAILQS FFTKMAKKKS LMDIPESQSE
301 QDFVLRVCGR DEYLVGETPI KNFQWVRHCL KNGEEIHVVV DTPDPALDE VRKEEWPLVD
361 DCTGVTGYHE QLTIHGKDHE SVFTVSLWDC DRKFRVKIRG IDIPVLPNT DLTVFVEANI
421 QHGQQVLCQR RTSPKPFTEE VLWNVWLEFS IKIKDLPKGA LLNLQIYCGK APALSSKASA
481 ESPSSESKGK VQLLYYVNL LIDHRFLLRR GEYVLHMWQI SGKGEDQGSF NADKLT SATN
541 PDKENSMSIS ILLDNYCHPI ALPKHQTPD PEGDRVRAEM PNQLRKQLEA I IATDPLNPL
601 TAEDKELLWH FRYESLKHPK AYPKLFSSVK WGQQEIVAKT YQLLARREVW DQSALDVGLT
661 MQLLDCNFSD ENVRAIAVQK LESLEDDDLV HYLLQLVQAV KFEPYHDSAL ARFLLRGLR
721 NKRIGHFLFW FLRSEIAQSR HYQQRFAVIL EAYLRGCGTA MLHDFTQQVQ VIEMLQKVTL
781 DIKLSAEKY DVSSQVISQL KQKLENLQNS QLPESFRVPY DPGLKAGALA IEKCKVMASK
841 KKPLWLEFKC ADPTALSNET IGIIFKHGDD LRQDMLILQI LRIMESIWET ESLDLCLLPY
901 GCISTGDKIG MIEIVKDATT IAKIQQSTVG NTGAFKDEVL NHWLKEKSPT EEKFQAAVER
961 FVYSCAGYCV ATFVLGIGDR HNDNIMTET GNLFHIDFGH ILGNYKSFLG INKERVPFVL
1021 TPDFLFVMTG SGKKTSPHFQ KFQDICVKAY LALRHHTNLL IILFSMMLMT GMPQLTSKED
1081 IEYIRDALTV GKNEEDAKKY FLDQIEVCRD KGWTVQFNWF LHLVLGIKQG EKHSA
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Recombinant P13 Kinase (p120 γ) nucleotide sequence:

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1 atgtcgact accatcacca tcaccatcac gattacgata tcccaacgac cgaaaacctg
61 tattttcagg gcgccatggg atccggaatt cgcccttgca tggagctgga gaactataaa
121 cagcccgtgg tgctgagaga ggacaactgc cgaaggcgcc ggaggatgaa gccgcgcagt
181 gctgcggcca gcctgtcctc catggagctc atccccatcg agttcgtgct gccaccagc
241 cagcgcaaat gcaagagccc cgaaacggcg ctgctgcacg tggccggcca cggcaactg
301 gagcagatga aggccaggt gtggctgca gcgctggaga ccagcgtggc ggcggacttc
361 taccaccggc tgggaccgca tcaactcctc ctgctctatc agaagaagg gtagtggtac
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Reviewed and approved by site quality representative.

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