

## Certificate of Analysis

### B-Raf (V599E), active

Also designated B-Raf (V600E)

(Recombinant enzyme expressed in Sf21 insect cells)

Item # 14-557, 14-557-K, 14-557M

Parent Lot # 28439U

The data presented in this document apply to the parent lot shown above and to all pack sizes derived from subsequent vialling runs of this parent lot. An alphabetical suffix after the parent lot number is used to denote each vialling run.

**Product Description:** N-terminal GST-tagged recombinant human B-Raf residues 416-end, containing a V599E mutation. Expressed by baculovirus in Sf21 insect cells. Purified using glutathione agarose. Purity 44% by SDS-PAGE and Coomassie blue staining. MW = 67.3kDa.

**Specific Activity (Parent lot# 28439U):** 167914U/mg, where one unit of B-Raf, active = 1 unit of MAPK2(m) activity which in turn is equivalent to 1nmol phosphate incorporated into 0.33mg/ml myelin basic protein per minute at 30°C with a final ATP concentration of 100µM. Note the activity is determined by a triple linked assay which involves the activation of MEK1(h) by B-Raf, followed by the subsequent activation of MAPK2(m) by the activated MEK1(h).

**Formulation:** 0.823mg/ml of enzyme in 50mM Tris/HCl pH7.5, 150mM NaCl, 0.1mM EGTA, 0.03% Brij-35, 270mM sucrose, 1mM benzamidine, 0.2mM PMSF, 0.1% 2-mercaptoethanol. Frozen solution.

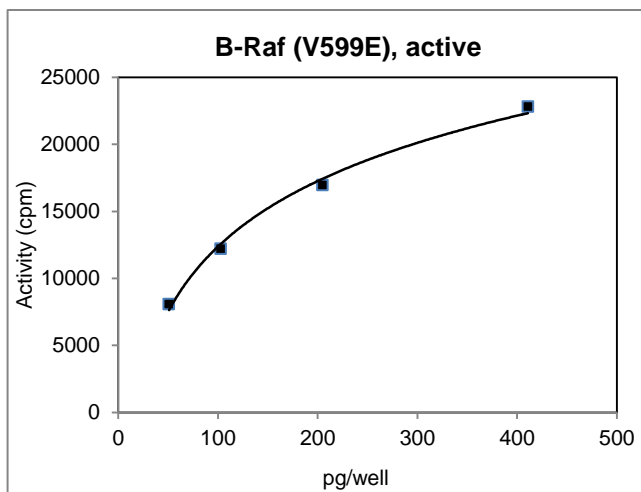
**Storage and Stability:** On receipt of material store at -70°C. Unopened reagent is stable for a minimum of 1 year from date of shipment when stored at recommended storage temperature. Avoid repeat freeze/thaw cycles. For maximum recovery of product, centrifuge original vial prior to removing the cap.

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

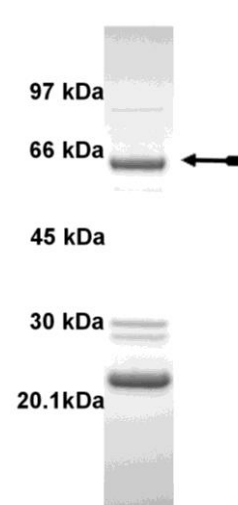
**FOR IN VITRO RESEARCH USE ONLY  
NOT FOR USE IN HUMANS OR ANIMALS**

### Quality Control Testing

**Kinase Assay:** 51-411pg of this lot of enzyme was used to activate 0.2µM MEK1(h) in the assay described on page two. Assay background was subtracted from the actual counts to yield the results.



**MS Tryptic Fingerprint:** Confirmed product identity as B-Raf with the translated sequence listed on page three.



**SDS-PAGE and Coomassie Stain:** Purity was assessed by SDS-PAGE and Coomassie blue staining using 3µg of B-Raf, active.

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### Kinase Assay Protocol

#### Stock Solutions:

1. **10 x Reaction Buffer:** 500mM Tris/HCl pH7.5, 1mM EGTA, 1mM Na<sub>3</sub>VO<sub>4</sub>, 1% 2-mercaptoethanol, 0.1% Brij-35.
2. **MEK1(h), unactive:** Use at a final assay concentration of 0.2μM (0.0126mg/ml). Prepare a 0.126mg/ml stock and add 2.5μl stock per assay point.
3. **MAPK2(m), unactive:** Use at a final assay concentration of 2μM (0.14mg/ml). Prepare a 0.7mg/ml stock and add 5μl of stock per assay point.
4. **Myelin Basic Protein (MBP):** Use at a final assay concentration of 0.33mg/ml. Prepare a 3.33mg/ml stock and add 2.5μl stock per assay point.
5. **B-Raf, active:** Dilute with 25mM Tris/HCl pH7.5, 0.1mM EGTA, 1mg/ml BSA. Use 51-411pg per assay point.
6. **[γ-<sup>33</sup>P]ATP:** 2.5 x magnesium acetate/[γ-<sup>33</sup>P]ATP cocktail: 25mM MgAc and 0.25mM ATP to which is added [γ-<sup>33</sup>P]ATP (specific activity approximately 500 - 800cpm/pmol as required.)

#### Assay Procedure (96 well plate format):

1. Add 2.5μl of 10 x reaction buffer to wells.
2. Add 2.5μl of **MEK1(h), unactive**.
3. Add 2.5μl of **MAPK2 (m), unactive**.
4. Add 2.5μl of **MBP**.
5. Add **5μl (51–411pg) B-Raf, active**.
6. Add 10μl of diluted [γ-<sup>33</sup>P] ATP mixture.
7. Incubate for 10 minutes at 30°C.
8. Stop the reaction by adding 5μl 3% phosphoric acid to each well.
9. Using a multichannel pipette, spot 10μl onto the appropriate area of a **P30 Filtermat**.
10. Wash the filtermat twice for 5 minutes with 75mM phosphoric acid.
11. Wash the filtermat once for 2 minutes with methanol.
12. Once dry, transfer the filtermat to a sealable plastic bag and add 4ml of scintillation cocktail.
13. Read in a scintillation counter. Compare cpm of enzyme samples with cpm of control samples that contain all assay components plus 1μl of 30% phosphoric acid.

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## B-Raf (V599E) Sequence Information

<u>Protein</u>	Human B-Raf (V599E)
<u>Tags</u>	N-Terminal GST
<u>Native sequence</u>	Q237 of the recombinant protein is equivalent to Q416 of human B-Raf
<u>Accession number</u>	GenBank NM_004333. The V599E mutation is thought to mimic phosphorylation of the native enzyme, resulting in a protein with high activity and leading to constitutive ERK activation. As B-Raf is commonly activated by somatic point mutation in human cancer, it may provide a new therapeutic approach to malignant melanoma. (Davies H., et al., <i>Nature</i> , 2002. 417: 949-54, and Mercer KE., & Pritchard CA., <i>Biochim Biophys Acta</i> . 2003. 1653: 25-40)

#### **Recombinant B-RAF (V599E) amino acid sequence:**

1 MSPILGYWKI KGLVQPTRLL LEYLEEKYEE HLYERDEGDK WRNKKFELGL EFPNLPYYID  
61 GDVKLTQSMA IIRYIADKHN MLGGCPKERA EISMLEGAVL DIRYGVSRIA YSKDFETLK  
121 DFLSKLPEML KMFEDRLCHK TYLNGDHVTB PDFMLYDALD VVLYMDPMCL DAFFKLVCFK  
181 KRIEAIQPQID KYLKSSKYIA WPLQGWQATF GGGDHPPKSD LEVLFQGPEF KGLRRQQKSP  
241 GPQRERKSSS SSEDRNRMKT LGRRDSSDDW EIPDGQITVG QRIGSGSFGT VYKGKWHGDV  
301 AVKMLNVTAP TPQQLQAFKN EVGVLRKTRH VNILLFMGYS TKPQLAIVTQ WCEGSSLYHH  
361 LHIIETKFEM IKLIDIARQT AQGMDYLHAK SIIHRDLKSN NIFLHEDLTV KIGDFGLATE  
421 KSRWSGSHQF EQLSGSILWM APEVIRMQDK NPYSFQSDVY AFGIVLYELM TGQLPYSNIN  
481 NRDQIIFMVG RGYLSPDLSK VRSNCPKAMK RLMAECLKKK RDERPLFPQI LASIELLARS  
541 LPKIHRSAE PSLNRAGFOT EDFSLYACAS PKTPIQAGGY GAFPVH

#### **Recombinant B-RAF (V599E) nucleotide sequence:**

1 atgtccccta tacttaggtt ttggaaaatt aaggcccttg tgcaaccac tcgacttc  
61 ttggaatatc ttgaagaaaa atatgaagag catttgtatg agcgcgatga aggtgataaa  
121 tggcgaaaca aaaagtttga attgggtttg gagtttcca atcttcctta ttatattgt  
181 ggtgatgtta aattaacaca gtctatggcc atcatacggtt atatacgta caagcacaac  
241 atgttggtt gttgtccaaa agagcgtgca gagatttcaa tgcttgaagg agcggttttgg  
301 gatatttagat acggtgtttc gagaattgca tatagtaaag actttgaaac tctcaaagtt  
361 gatttctta gcaagctacc tgaatgctg aaaatgttcg aagatcgat ttatgtcataaa  
421 acatatttaa atgggtatca tgtaaccat cctgacttca tggtgtatga cgctcttgat  
481 gttgtttat acatggaccc aatgtgcctg gatgcgttcc caaaaattagt ttgttttaaa  
541 aaacgttattt aagctatccc acaaattgtat aagtacttga aatccagcaa gtatatacg  
601 tggccttgc agggctggca agccacgtt ggtggtggcg accatcctcc aaaatcgat  
661 ctggaagttc tgttccaggg gcccgaaattc aaaggccctac gtcgacaaca gaaatctcca  
721 ggacctcagc gtgaaaggaa gtcatcttca tcctcagaag acaggaatcg aatgaaaaca  
781 cttggtagac gggactcgag tgatgattgg gagattcctg atgggcagat tacagtggga  
841 caaagaattt gatctggatc atttggaaaca gtctacaagg gaaagtggca tggtgatgt  
901 gcagtaaaaa tggtgaatgt gacagcacct acacctcagc agttacaagc cttaaaaat  
961 gaagttaggag tactcaggaa aacacgacat gtgaatatcc tactcttcat gggctattcc  
1021 acaaagccac aactggctat tgttacccag tgggtgtgagg gctccagctt gtatcaccat  
1081 ctccatatca ttgagaccaa atttgagatg atcaaactta tagatattgc acgacagact  
1141 gcacagggca tggattactt acacgccaag tcaatcatcc acagagacct caagagtaat  
1201 aatatatttc ttcatgaaga cctcacagta aaaataggtg attttggct agctacagag  
1261 aaatctcgat ggagtggtc ccatcagttt gaacagttgt ctggatccat tttgtggatg  
1321 gcaccagaag tcatcagaat gcaagataaa aatccataca gctttcagtc agatgtatata  
1381 gcatttggaa ttgttctgtt tgaattgtatc actggacagt taccttattt aaacatcaac  
1441 aacaggggacc agataatttt tatggtggga cgaggatacc tgtctccaga tctcagtaag  
1501 ctacqqaqta actqtccaaa aqccatqaaq aqattaatqq cagagtqccct caaaaagaaaa

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1561 agagatgaga gaccactttt tccccaaattt ctcgcctcta ttgagctgct ggcccgctca  
1621 ttgccaaaaaa ttcacccgcag tgcatcagaa ccctccttga atcgggctgg tttccaaaca  
1681 gaggatttttta gtctatatgc ttgtgcttct ccaaaaacac ccatccaggc agggggatat  
1741 ggtgcgtttc ctgtccactg a

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