

Division of Signal Transduction Therapy

Standard Operating Procedure

Preparation of active EPH A2 [591 – 976]

Enzyme description:- EPH A2 [591 - 976]

Clone number:- DU 4438

Source:- Recombinant

Expression system:- *E.coli*

Tag:- N-terminal GST

Purification method:- GSH Sepharose

Expression level:- 6 mg/L

Calculated molecular mass:-

Monoisotopic 70, 354.92 daltons

Average Mass 70, 400.53 daltons

[cysteines reduced, methionines have not been oxidised]

Theoretical pI:- 6.94

Purity:- 90 %

Activation protocol:- Constitutively active

Enzyme storage buffer:-

50 mM Tris-HCl pH 7.5, 270 mM Sucrose, 150 mM NaCl, 0.1 mM EGTA,
0.1 % 2-mercaptoethanol, 0.02 % Brij-35, 1 mM benzamidine, 0.2 mM PMSF

Storage temperature:- -80 °C

Assay:- Standard filter binding assay

Assay buffer:-

50 mM Tris-HCl pH 7.5, 0.1 mM EGTA, 0.1 % 2-mercaptoethanol, 10 mM MgAc

Substrate:-

Poly Glu Tyr (4:1) Final concentration: 0.1 mg/ml

Specific activity range:- To be determined

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Clone Data Sheet

EPH A2 [591 - 976]

<u>Protein</u>	EPH A2 [591 - 976]
<u>Clone number</u>	DU 4438
<u>Species</u>	Human
<u>Accession number</u>	NM_004431
<u>Tags</u>	N-terminal GST
<u>Bacterially expressed protein</u>	MSPILGYWKIKGLVQPTRLLEYLEEKYEEHLYERDEGDKWRNKKFELG LEFPNLPYYIDGDVKLTQSMAIIRYIADKHNMLGGCPKERAETSMLEGA VLDIRYGVSR IAYSKDFETLKVDFLSKLPEMLKMFEDRLCHKTYLNGDH VTHPDFMLYDALDVLYMDPMCLDAFPKLVCFKKRIEAIPOIDKYLKSS KYIAWPLQGWQATFGGGDHPPKSDLEVLFGGPLGSPHTYEDPNQAVLKF TTEIHPSCVTRQKVI GAGEFGEVYKGMLKTSSGKKEVPVAIKTLKAGYT EKQRVDFLGEAGIMGQFSHHNIIRLEGVISKYKPMMIITEYMENGALDK FLREKDGEFSVLQLVGMLRGIAAGMKYLANMNYVHRDLAARNILVNSNL VCKVSDFGLSRVLEDDPEATYTTSSGKIPIRWTAPEAISYRKFTSASDV WSFGIVMWEVMTYGERPYWELSNHEVMKAINDGFR LPTPMDCP SAIYQL MMQCWQQERARRPKFADIVSILDKLIRAPDSLKTLADFDPRVSIRLPST SGSEGVPFRTVSEWLESIKMQQYTEHFMAAGYTAIEKVVMQMTNDDIKRI GVRLPGHQKRIAYSLLGLKDQVNTVGIP I
<u>Native sequence</u>	Amino acids P591 – I976 (end) of human EPH A2. Residue P232 of the fusion protein is equivalent to P591 of the native enzyme. The GST tag is located at residues 1 – 220.
<u>Protease cleavage</u>	PreScission (<u>LEVLFQGPL</u>) residues 221 - 229
<u>Cloning sites</u>	<i>Bgl</i> III and <i>Sal</i> I site of pGEX 6P-1

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Nucleotide Sequence

atgtcccctatactagggttatttgaaaaattaagggccttgtgcaaccca
ctcgacttccttttggaaatccttgaagaaaaatatgaagagcatttgta
tgagcgcgatgaaggtgataaatggcgaaacaaaaagtttgaattgggt
ttggagtttcccaatccttccttattatattgatgggtgatgttaaattaa
cacagtctatggccatcatacgttatatagctgacaagcacacaacatgtt
gggtgggttgtccaaaagagcgtgcagagatttcaatgcttgaaggagcg
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ACTACCGAGATCCATCCATCCTGTGTCACTCGGCAGAAGGTGATCGGAG
CAGGAGAGTTTGGGGAGGTGTACAAGGGCATGCTGAAGACATCCTCGGG
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GAGAAGCAGCGAGTGGACTTCCTCGGCGAGGCCGGCATCATGGGCCAGT
TCAGCCACCACAACATCATCCGCCTAGAGGGCGTCATCTCCAAATACAA
GCCCATGATGATCATCACTGAGTACATGGAGAATGGGGCCCTGGACAAG
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CGCCCCGGAGGCCATTTCTTACCGGAAGTTCACCTCTGCCAGCGACGTG
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CCTACTGGGAGTTGTCCAACCACGAGGTGATGAAAGCCATCAATGATGG
CTTCCGGCTCCCCACACCCATGGACTGCCCCCTCCGCCATCTACCAGCTC
ATGATGCAGTGCTGGCAGCAGGAGCGTGCCCGCCGCCCAAGTTCGCTG
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TGCCATCGAGAAGGTGGTGCAGATGACCAACGACGACATCAAGAGGATT
GGGGTGCGGCTGCCCCGCCACCAGAAGCGCATCGCCTACAGCCTGCTGG
GACTCAAGGACCAGGTGAACACTGTGGGGATCCCCATctgagtcgac