

## *Division of Signal Transduction Therapy*

### **Standard Operating Procedure**

#### **Preparation of active Testis Specific Serine Kinase 1B [1 - 367]**

**Enzyme description:-** TSSK1B [1 - 367]

**Clone number:-** DU 34845

**Source:-** Recombinant

**Expression system:-** *E.coli*, co-expressed

**Tag:-** N-terminal GST

**Purification method:-** GSH Sepharose

**Calculated molecular mass:-**

Monoisotopic 68,397.88 daltons

Average Mass 68,442.10 daltons

[cysteines reduced, methionines have not been oxidised]

**Theoretical pI:-** 6.52

**Purity:-** >80 %

**Enzyme storage buffer:-**

50 mM Tris-HCl pH 7.5, 150 mM NaCl, 270 mM sucrose, 0.1 mM EGTA,  
0.1 % 2-mercaptoethanol, 0.03 % Brij-35, 1 mM benzamidine, 0.2 mM PMSF

**Storage temperature:-** -70 °C

**Assay buffer:-**

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

**Substrate:-**

CHKtide [KKKVSRSGLYRSPSPENLNRPR]

Final concentration: 300  $\mu$ M

*Division of Signal Transduction Therapy*

**Clone Data Sheet**

**TSSK1B [1 - 367]**

**Protein** TSSK1B [1 - 367]

**Clone number** DU 34845

**Species** Human

**Accession number** AAH22515.1

**Tags** N-terminal GST

**Bacterially  
expressed protein**

MSPILGYWKIKGLVQPTRLLEYLEEKYEEHLYERDEGDKWRNKKFEL  
GLEFPNLPYYIDGDVKLTQSMAIIRYIADKHNMLGGCPKERAIEISMLE  
GAVLDIRYGVSRIAYSKDFETLKVDFLSKLPPEMLKMFEDRLCHKTYLN  
GDHVTHPDFMLYDALDVVLYMDPMCLDAFPKLVCFKKRIEAIPOIDKY  
LKSSKYIAWPLQGWQATFGGGDHPKSDLEVLFGPPLGSMDDAAVLKR  
**RGYLLGINLGEESYAKVKSAYSERLKFNVAIKIDRKKAPADFLKFL**  
**PREIEILAMLNHCSI IKTYEIFETSHGKVYIVMELAVQGDLELEIKTR**  
**GALHEDEARKKFHQLSLAIKYCHDLDVVHRDLKCDNLLLDKDFNIKLS**  
**DFSFSKRCLRDDSGRMALSKTFCGSPAYAAPEVLQGIPIYQPKVYDIWS**  
**LGVILYIMVCGSMPYDDSNIKKMLRIQKEHRVNFPRSKHLTGECKDLI**  
**YHMLQPDVNRRLHIDEILSHCWMQPKARGSPSVAINKEGESSRGTEPL**  
**WTPEPGSDKKSATKLEPEGEAQPQAQPETKPEGTAMQMSRQSEILGFP**  
**SKPSTMETEEGPPQPPETRAQ**

**Native sequence** Amino acids M1 – Q367 of human TSSK1B.  
Residue M232 of the fusion protein is equivalent to M1 of the native  
enzyme. The GST tag is located at residues 1 – 220.

**Protease cleavage** Precission site (LEVLFGQP) at residues 221 - 228

**Cloning sites** *Bam*H1 and *Not*1 sites of pGEX-6P

*Division of Signal Transduction Therapy*

**Nucleotide  
sequence of insert**

ggatccATGGATGACGCTGCTGTCCTCAAGCGACGAGGCTACCTCCTG  
GGGATAAAATTTAGGAGAGGGCTCCTATGCAAAAGTAAAATCTGCTTAC  
TCTGAGCGCCTGAAGTTCAATGTGGCGATCAAGATCATCGACCGCAAG  
AAGGCCCCCGCAGACTTCTTGGAGAAATTCCTTCCCCGGGAAATTGAG  
ATTCTGGCCATGTTAAACCACTGCTCCATCATTAAGACCTACGAGATC  
TTTGAGACATCACATGGCAAGGTCTACATCGTCATGGAGCTCGCGGTC  
CAGGGCGACCTCCTCGAGTTAATCAAAACCCGGGGAGCCCTGCATGAG  
GACGAAGCTCGCAAGAAGTTCACCAGCTTTTCTTGGCCATCAAGTAC  
TGCCACGACCTGGACGTCGTCCACCGGGACCTCAAGTGTGACAACCTT  
CTCCTTGACAAGGACTTCAACATCAAGCTGTCCGACTTCAGCTTCTCC  
AAGCGCTGCCTGCGGGATGACAGTGGTGAATGGCCTTAAGCAAGACC  
TTCTGTGGGTCACCAGCGTATGCGGCCCCAGAGGTGCTGCAGGGCATT  
CCCTACCAGCCCAAGGTGTACGACATCTGGAGCCTAGGCGTGATCCTC  
TACATCATGGTCTGCGGCTCCATGCCCTACGACGACTCCAACATCAAG  
AAGATGCTGCGTATCCAGAAGGAGACCGCGTCAACTTCCCACGCTCC  
AAGCACCTGACAGGCGAGTGCAAGGACCTCATCTACCACATGCTGCAG  
CCCGACGTCAACCGGCGGCTCCACATCGACGAGATCCTCAGCCACTGC  
TGGATGCAGCCCAAGGCACGGGGATCTCCCTCTGTGGCCATCAACAAG  
GAGGGGGAGAGTTCCCGGGGAACCTGAACCCTTGTGGACCCCCGAACCT  
GGCTCTGACAAGAAGTCTGCCACCAAGCTGGAGCCTGAGGGAGAGGCA  
CAGCCCCAGGCACAGCCTGAGACAAAACCCGAGGGGACAGCAATGCAA  
ATGTCCAGGCAGTCGGAGATCCTGGGTTTCCCCAGCAAGCCGTCGACT  
ATGGAGACAGAGGAAGGGCCCCCCCCAACAGCCTCCAGAGACGCGGGCC  
CAGtgagcggccgc