

# PKM2

**PDB:**3ME3

## Revision

**Revision Type:**created

**Revised by:**created

**Revision Date:**created

**Entry Clone Accession:**BC007952

**Entry Clone Source:**MGC: AU36-H1

**SGC Clone Accession:**HPC002-A01

**Tag:**N-terminal His6-tag, not removed

**Host:**BL21-V2R

## Construct

**Prelude:**PANK3: M1-P531

**Sequence:**

mgsshhhhhssglvprgsMSKPHSEAGTAFIQTQQLHAAMADTFLEHMCRLDIDSPITARNTGIICTIGPASRSVETLKEMIKSG  
MNVARLNFSGHGTHEYHAETIKNVRTATESFASDPILYRPVAVALDTKGPEIRTGLIKSGTAEVELKKGATLKITLDNAYMEKCDEN  
ILWLDYKNICKVVEVGSKIYVDDGLISLQVKQKGADFLVTEVENGGSLGSKKGVNLPGAAVDLPVSEKDIQDLKFGVEQDVMVFA  
SFIRKASDVHEVRKVLGEKGKNIKIISKIENHEGVRRFDEILEASDGMVARGDLGIEIPA EKVF LAQKMMIGRCNRAGKPVICATQ  
MLESMIKKPRPTRAEGSDVANAVLDGADCIMLSGETAKGDYPLEAVRMQH LIAREAEAAIYHLQLFEELRR LAPITSDPTEATAVGA  
VEASFKCCSGAIIVLTKSGRSAHQVARYRPRAPIIAVTRNPQTARQAHL YRGIFPVLCKDPVQEAWAEDVDLRVNFAMNVGKARGFF  
KKGDVVIVLTGWRPGSGFTNTMRVVPVP

**Vector:**pET28a-LIC

## Growth

**Medium:**

**Antibiotics:**

**Procedure:**The seeds were grown in 80 mL Luria-Bertani broth media supplemented with 50 µg/mL kanamycin at 37 °C overnight. The following morning, all of the seeds were inoculated 1800 mL of Terrific Broth media supplemented with 50 µg/mL kanamycin, 8 g/l glycerol and approximately 500 µl antifoam in glass flasks in the Large Scale Expression System (LEX). Cells were grown at 37 °C until OD<sub>600nm</sub> of 4.0 and were then induced by addition of IPTG to a concentration of 0.5 mM. Protein expression was allowed to continue over night at 18 °C.

## Purification

**Procedure**

The supernatant was passed through DE52 (Whatman) column equilibrated with the binding buffer and then loaded onto 3 mL Ni-NTA column (Qiagen) equilibrated with the same binding buffer at 4 °C. The Ni-NTA column was washed with 150 mL of the wash buffer and the protein

was eluted with 15 mL of the elution buffer. The eluate was further purified by size-exclusion chromatography (Superdex 200) equilibrated with 10 mM HEPES pH 7.5, 150mM KCl, 2mM TCEP, 5% glycerol, and 5mM MgCl<sub>2</sub>. The protein was concentrated using an Amicon Ultra centrifugal filter to the concentration of 50 mg/mL.

## **Extraction**

### **Procedure**

Cultures were centrifuged and the cell pellets were suspended in 100 mL of the binding buffer with a protease inhibitor cocktail (0.1 mM M benzamidine-HCl and 0.1 mM phenylmethyl sulfonyl fluoride) and flash frozen. The thawed cell pellet was lysed by a combination of 0.5% CHAPS (Sigma) and sonication. The lysate was centrifuged at 15000 rpm for 30 min and the supernatant was used for subsequent steps of purification.

**Concentration:** 50 mg/mL

### **Ligand**

3- {[4-(2,3-dihydro-1,4-benzodioxin-6-ylsulfonyl)- 1,4-diazepan-1-yl]sulfonyl} aniline **MassSpec:**

**Crystallization:** Crystallization trials were set up using the vapor diffusion method and the protein drop was equilibrated against a reservoir solution with 1:1 volume ratio. Prior to crystallization, the purified protein was incubated overnight at 4 °C in the presence of the activator (to 5 ~10 mM final concentration). Crystals of activator bound PKM2 were grown at 25% PEG-3350, 0.1M ammonium sulfate, 0.1M Bis-Tris, pH6.5.

### **NMR Spectroscopy:**

#### **Data Collection:**

#### **Data Processing:**