

# 33.m01398

**PDB:**3DZO

## Revision

**Revision Type:**created

**Revised by:**created

**Revision Date:**created

**Entry Clone Accession:**

**Entry Clone Source:**

**SGC Clone Accession:**33.m01398:F196-T575:F09

**Tag:**N-terminal: mgsshhhhhhsgrenlyfqg

**Host:**BL21-(DE3)-V2R-pACYC LamP

## Construct

**Prelude:**

**Sequence:**

FRGTDPGDVVIEELFNRIQPQANVRTTSEYMQSAADSLVSTSLWNTGQPFRVESELGERPRTLVRGTVLGQEDPYAYLEATDQETGES  
FEVHVVPYFTERPPSNAIKQMKEEVRLRLRGIKNQKQAKVHLRFIFPFDLVKDPQKKMIRVRLDERDMWVLSRFFLYPRMQSNLQ  
TFGEVLLSHSSTHKSLVHHARLQLTLQVIRLLASLHHYGLVHTYLRPVDIVLDQRGGVFTGFEHLVRDGASAVSPIGRGFAPPETT  
AERMLPFGQHHPTLMTFAFDWTGLAIYWIWCADLPNTDDAALGGSEWIFRSCKNIPQPVRALLEGFLRYPKEDRLLPLQAMETPE  
YEQLRTESAALPLYQTDGEPTREGGAPPSGTSQPDEAGAAEAVTAI

**Vector:**pET15-MHL

## Growth

**Medium:**TB

**Antibiotics:**

**Procedure:**TgROP2 was expressed in E. coli BL21-(DE3)-V2R-pACYC LamP strain in Terrific Broth (TB) in the presence of ampicillin/chloramphenicol (100 microgram/mL and 34 microgram/mL respectively). A single colony was inoculated into 100mL of LB with of ampicillin/chloramphenicol (100 microgram/mL and 34 microgram/mL respectively) in a 250 mL baffled flask and incubated with shaking at 250 rpm overnight at 37 °C. The culture was transferred into 4.0 L of TB with ampicillin/chloramphenicol (100 microgram/mL and 34 microgram/mL respectively) and 0.15 mL of antifoam (Sigma) in a 1 L bottle and cultured using the LEX system to an OD600 of 5-6, cooled to 15 °C, and induced with 0.5 mM isopropyl-1-thio-D-galactopyranoside (IPTG) overnight at 15 °C.

## Purification

**Procedure**

The cleared lysate was loaded onto a 1.0-2.5 mL Ni-NTA (Qiagen) column (pre-equilibrated with Binding Buffer) at approximately 1.5-2.0 mL/min. The Ni-NTA column was then washed with

150 mL of Wash Buffer at 2-2.5 mL/min. After washing, the protein was eluted with Elution Buffer. The eluted sample was applied to a Sephadex S200 26/60 gel filtration column pre-equilibrated with Gelfiltration Buffer. The fractions corresponding to the eluted protein peak were collected and further concentrated using a 15 mL Amicon Ultra centrifugal filter device (Millipore). The protein sample identity and purity were evaluated by mass spectroscopy and SDS-PAGE gel. The concentrated protein was stored at 4 degC. For long term storage, the protein was flash frozen and stored at -80 degC.

## **Extraction**

### **Procedure**

The culture was harvested by centrifugation. Pellets from 4 L of culture were resuspended to approximately 40 mL/L of cell culture in Binding Buffer with the addition of protease inhibitors (1 mM benzamidine and 1 mM phenylmethyl sulfonyl fluoride (PMSF)). Resuspended pellets stored at -80 degC were thawed overnight at 4 degC on the day before purification. Prior to sonication, each pellet from 1 L of culture was pretreated with 0.5 % CHAPS and 500 units of benzonase for 40 minutes at room temperature. After 10 minutes sonication, the cell lysate was centrifuged using a Beckman JLA-16.250 rotor at 15,500 rpms for 45 minutes at 4 degC.

### **Concentration:**

### **Ligand**

### **MassSpec:**

**Crystallization:** 22% PEG4000, 0.2M MgCl<sub>2</sub>, 0.1M Tris-HCl pH8.8 plus 10% Glycerol; Hanging Drop. 293K

### **NMR Spectroscopy:**

### **Data Collection:**

### **Data Processing:**