

# HSP90

**PDB:**3IED

## Revision

**Revision Type:**created

**Revised by:**created

**Revision Date:**created

**Entry Clone Accession:**PF14\_0417

**Entry Clone Source:**

**SGC Clone Accession:**PF14\_0417:S97-E347:E10

**Tag:**N-terminal tag: mhhhhhssgrenlyfqg

**Host:**BL21-(DE3)-V2R-pRare2.

## Construct

**Prelude:**

**Sequence:**

SPVEKYNFKAENVKVMIDIIVNSLYTDKDVFLRELISNASDACDKKRIILENNKLIKDAEVVTNNEIKNETEKEKTENVNESTDKKEN  
VEEEKNDIKKLIKIKPDKEKKTLTITDNGIGMDKSELINNLGTIAQSGTAKFLKQIEEGKADSNLIGQFGVGFYSSFLVSNRVEVY  
TKKEDQIYRWSSDLKGSFSVNEIKKYDQYDDIKSGTKIILHLKEECDEYLEDYKLKELIKKYSEFIKFPTEIWSE

**Vector:**p15-mhl

## Growth

**Medium:**TB

**Antibiotics:**

**Procedure:**Plasmodium falciparum PP-HSP90(PF14\_0417) was expressed in E. coli BL21(λDE3) V2R pRare2 in TB growth media in the presence of carbenicillin/chloramphenicol (100 microgram/mL and 34 microgram/mL, respectively). A single colony was inoculated into 25 mL of LB with of carbenicillin/chloramphenicol (100 microgram/mL and 34 microgram/mL respectively) in a 50 mL Falcon tube and incubated with shaking at 250 rpm overnight at 37 degC. Then the culture was transferred into 900 mls of TB with 100 microgram/mL Carbenicillin and 34 microgram/ml chloramphenicol , 0.3 mL of antifoam (Sigma), 9 mls of 0.83 M MgSO4 and trace elements in a 1L bottle and cultured using the LEX system to an OD600 of 5, cooled to 15 degC, and induced with 0.5 mM isopropyl-1-thio-D-galactopyranoside (IPTG) overnight at 15 degC.

## Purification

**Procedure**

The cleared lysate was loaded onto a column prepacked with 10 g DE52 (Whatman) anion exchange resin (previously activated with 2.5 M NaCl and equilibrated with Binding Buffer); and

subsequently onto a 2mL Ni-NTA (Qiagen) column pre-equilibrated with Binding Buffer at approximately 1 - 1.5 mL/min. The volume of the Ni-NTA resin was pre-determined by the predicted protein yield from test expression analysis. After the lysate was loaded, the DE52 was further washed with 20 mL of Binding Buffer. The Ni-NTA column was then washed with 200 mL of Wash Buffer at 2 - 2.5 mL/min. After washing, the protein was eluted with 15 mL of Elution Buffer. 1 mM TCEP and 1 mM EDTA was added to the eluted PP-HSP90 (PF14\_0417).

The sample was then loaded onto a superdex 200 gel filtration column. The eluted protein ( in 10 mM Hepes, pH 7.5 and 500 mM NaCl) was concentrated using a 15 ml Amicon Ultra centrifugal filter device (Millipore) with a 10 kDa cutoff. PP-HSP90 (PF14\_0417) was concentrated to 33 mg/ml and stored at 4 degC. The protein was diluted to 15 mg/ml before use in crystal trials.

## **Extraction**

### **Procedure**

The culture was harvested by centrifugation. A pellet from 1 L of culture was 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)). The resuspended pellet , stored at -80 degC, was thawed overnight at 4 degC on the day before purification. Prior to mechanical lysis, the pellet from 1 L of culture was pretreated with 0.5 % CHAPS and 500 units of benzonase and protease inhibitor (1 mM benzamidine and 1 mM phenylmethyl sulfonyl fluoride (PMSF)) for 40 minutes at room temperature. Cells were mechanically lysed with a microfluidizer (Microfluidizer Processor, M-110EH) at approximately 18000 psi; and the cell lysate was centrifuged using a Beckman JA-25.50 rotor at ~75000 x g (24000 rpms) for 20 minutes at 10 degC.

**Concentration:**33 mg/ml.

### **Ligand**

#### **MassSpec:**

**Crystallization:**The protein was crystallized in 1.5 M NaCitrate, 0.1 M Na Cacodylate pH 5.5 at 20 degC using the sitting drop method. The ligand added at 2 mM AMPPNP, 4 mM MgCl<sub>2</sub>, 2 mM TCEP.

#### **NMR Spectroscopy:**

#### **Data Collection:**

#### **Data Processing:**