

# CYP11A1

**PDB:**3NA0

## Revision

**Revision Type:**created

**Revised by:**created

**Revision Date:**created

**Entry Clone Accession:**AT51-H8 (BC032329)

**Entry Clone Source:**MGC

**SGC Clone Accession:**

**Tag:**N-terminal: ferredoxin - MGC: BC010284; C-terminal: 6His-tag

**Host:***E.coli* JM109 (Stratagene)

## Construct

**Prelude:**

**Sequence:**

STRSPRPFNEIPSPGDNGWLNLYHFWRETGTHKVHLHHVQNFQKYGPITYREKLGNVESVYVIDPEDVALLFKSEGNPERFLIPPWV  
AYHQYYQRPIGVLLKKSAAWKKDRVALNQEVMPEATKNFLPLLDVSRDFVSVLHRRIKKAGSGNYSGLDSDLFRFAFESITNVI  
FGERQGMLEEVNPEAQRFDIAIYQMFHTSVPMLNLPDLFRLFRTKTWKDHVAAWDVIFSKADIYTQNFYWELRQKGSVHHDYRGI  
LYRLLGDSKMSFEDIKANVTEMLAGGVDTTSMTLQWHL YEMARNLKVQDMLRAEVLAARHQAQGD MATMLQLVPLLKASIKETLR LH  
PISVTLQRYLVNDLVLRDYMIPAKTLVQVAIYALGREPTFFFDPENFDPTRWLSKDKNITYFRNLGFGWGVQRCLGRRIAELEMTIF  
LINMLENFRVEIQHLSVDVGTTFNLILMPEKPISFTFWPFNQEATQQ

**Vector:**pCW-LIC-29

## Growth

**Medium:**

**Antibiotics:**

**Procedure:**CYP11A1 was co-expressed with GroEL/ES in *E.coli* JM109 in TB medium. Cells were grown at 37°C to an OD<sub>600</sub> of 1.0 and induced by 0.5mM IPTG and 4mg/ml of arabinose and in the presence of 0.5mM  $\delta$ -aminolevulinic acid and incubated 48 hours at 26°C.

## Purification

**Procedure**

The lysate was centrifuged at 60 000g for 60min. The supernatant was loaded onto 5ml NiHiTrap column (Amersham Biosciences) equilibrated with Buffer A. The column was washed with buffer A and protein was eluted using a linear gradient of 5-100% Buffer B. The protein was further purified by ion-exchange chromatography on SourceQ column (Amersham Biosciences), equilibrated with 5 mM KPi, pH 7.4, 20% glycerol and 4 mM CHAPS and eluted with linear gradient of Buffer C.

## **Extraction**

### **Procedure**

Collected/resuspended cells in extraction buffer were disrupted in a high-pressure Microfluidizer (Microfluidics Corp.) at 18,000 psi. CHAPS was added to a final concentration 16 mM and lysate was incubated at 4°C for 60 min.

**Concentration:** 20 mg/ml

### **Ligand**

### **MassSpec:**

**Crystallization:** Purified CYP11A1 was crystallized in presence of cholesterol, 22R-hydroxycholesterol, 20S-hydroxycholesterol, and 20R,22R-dihydroxycholesterol using hanging drop vapor diffusion method drop at 18°C by mixing 1 µl of the protein solution with 1 µl of the reservoir solution containing 10% PEG 8K, 0.2 M calcium acetate, and 0.1 M HEPES pH 7.5.

### **NMR Spectroscopy:**

### **Data Collection:**

### **Data Processing:**