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# Expression and purification of recombinant UvsX recombinase

Forked from Expression and purification of recombinant RT-MMLV

lucero.merino.c<sup>1</sup>, lucero.mascaro.r<sup>1</sup>

<sup>1</sup>Universidad Peruana Cayetano Heredia (UPCH, Peru)



#### **ABSTRACT**

The UvsX is a enzyme that is part for an isothermal DNA amplification based on the recombination process, the RPA reaction.

RPA uses 4 enzymes: UvsX, UvsY, Bsu and Gp32. It's an isothermal amplification technique that can run at 37°C. In this protocol we are producing a recombinant UvsX that has a 6xHIS-tag using a *E. coli* expression system.

The protocols for the production of the other proteins are also available in protocols.io.

#### **MATERIALS**

#### Binding buffer, pH 7.2

[M] 50 millimolar (mM) Tris-HCl, pH 7.2

[M] 20 millimolar (mM) Imidazole, pH 7.2

[M] 500 millimolar (mM) KCl

[M] 5 % (v/v) Glycerol

[M] 0.1 millimolar (mM) PMSF

[M] 0.03 % (v/v) 2-Mercaptoethanol (BME)

#### Elution buffer, pH 7.2

[M] 50 millimolar (mM) Tris-HCl, pH 7.2

[M] 500 millimolar (mM) Imidazole, pH 7.2

[м] 500 millimolar (mM) KCl

[M] 5 % (v/v) Glycerol

[M] 0.03 % (v/v) BME

#### Storage buffer, pH 8

[M] 20 millimolar (mM) Tris-HCl, pH 8

[M] 400-500 millimolar (mM) KCl

[M] 20 % (v/v) Glycerol

[M] 1 millimolar (mM) DTT

#### Ladder:

Pageruler Prestained Protein Ladder **Thermo Fisher Scientific Catalog** #26616

#### **Equipment:**

Sonicator OMNI Sonic Ruptor 400

Protein purification system FPLC AKTA START

### **DAY1: Transformation of competent cells**

1d

- 1 Quantify the plasmid containing the UvsX recombinase gene and determine the volume that contains A 100 ng of the plasmid.
- Defrost the aliquot of BL21(DE3) chemically competent cells On ice . Softly pipette

  L 100 ng of the plasmid in the aliquot and let the tube rest On ice for 00:30:00

30m

Incubate the tube at 42 °C for 00:00:30 .

30s

4 Quickly return the tube 8 On ice and incubate for 00:05:00

5m

- Add the mixture to a microcentrifuge tube with Δ 800 μL SOC medium and incubate at 37 °C for 00:45:00 .
- 45m

- 6 Centrifuge the tube 3 4500 rpm, Room temperature, 00:08:00

8m

- 7 Discard  $\perp$  800  $\mu$ L of the supernatant and gently resuspend the pellet with the rest.
- Add the resuspension to a LB agar plate with recently transformed cells. Incubate plate Overnight at \$37 °C.

### **DAY2: Preparation of pre-inoculum**

1d

- 9 For verification that the colonies in the plate contain the desired plasmid with the protein sequence, perform a PCR colony using universal T7 primers and the PCR protocol for Phusion DNA Polymerase
  - Phusion High-Fidelity DNA Polymerase 500 units **New England Biolabs Catalog** #M0530L
  - . Use the following thermocycling procedures for the UvsX recombinase plasmid:

Step	Temperature (°C)	Time	Cycles
Initial Denaturation	98	3 min	1
Denaturation	98	30 sec	25
Annealing	60	30 sec	
Extension	72	42 sec	
Final Extension	72	5 min	1
Hold	4	∞	

Run the PCR product in a 1% agarose gel and verify if there is a band of the desired weight (UvsX insert = 1387 bp).

Select an isolated bacterial colony from the plate and inoculate a test tube with

△ 10 mL LB medium and IMI 0.05 mg/mL Kanamycin . Incubate the tube ♦ Overnight at €5 220 rpm, 37°C .

### **DAY 3-A:Protein expression in small scale**

2d

- Inoculate  $\square$  50  $\mu$ L from the pre-inoculum to an Erlenmeyer flasks with  $\square$  50 mL LB medium and  $\square$  1. Incubate at  $\square$  2. Incubate at  $\square$  2. Incubate at hours).
- Add IPTG to a final concentration of [M] 0.5 millimolar (mM) and incubate at \$\tilde{\mathcal{O}}\$ 220 rpm, 18°C, 16:00:00 .
- Centrifuge the cell culture 8000 rpm, 4°C, 00:05:00. Discard the supernatant. At this point, you may store the cells pellet at -20°C until you are ready to run the purification.

5m

### **DAY 4-A:Protein purification in resin**

1d

Resuspend the cell pellet in  $\Delta$  5 mL Binding buffer . Then add lysozyme to a final concentration of [M]  $0.1 \, \mu g/\mu L$  .

- Add  $\sim \pm 100 \,\mu\text{L}$  of glass beads and shake vigorously in a vortex for  $\odot 00:20:00$  at room temperature. You can do this by fixing a 15 mL tube to the vortex rubber platform with tape.
- 17 Centrifugate at (5 13500 rpm, 4°C, 00:07:00). Collect the supernatant and label it as a Soluble fraction. The pellet is the Insoluble fraction. Collect small fractions of each one to run an acrylamide gel afterwards.
- Prepare the resin. Homogenize resin with its storage buffer by shaking the bottle and transfer it to a new tube. You will use A 330 µL of resin for each A 1 mL of soluble fraction. Let the slurry sediment or spin it down. Remove the storage buffer and wash the resin in **Binding** buffer. Wash the resin with the same volume as the obtained soluble fraction. Repeat this wash step 3 times.
- Add the soluble fraction to the resin. Homogenize the mixture gently in an orbital shaker for 20 min (~60 RPM) at room temperature.
- Let the resin sediment for 10 minutes. Collect a small fraction of the supernatant to run an acrylamide gel afterwards, and discard the remainder. Resuspend resin with 1 mL of **Binding buffer.** Homogenize the tube gently with finger taps. Don't flip the tube (1st washing step).
- 21 Spin down for a few seconds and discard supernatant. Resuspend resin with 1 mL of **Binding** buffer. Homogenize the tube gently with finger taps. Don't flip the tube (2nd washing step).
- Spin down for a few seconds and discard supernatant. Resuspend resin with 1 mL of **Elution buffer (50mM Imidazole).** Homogenize the tube gently with finger taps. Don't flip the tube. Incubate for 00:10:00.

10m

23 Spin down for a few seconds and collect the supernatant. Resuspend resin with 1 mL of **Elution** 

**buffer (500mM Imidazole).** Homogenize the tube gently with finger taps. Incubate for 00:10:00. Collect small fractions of elutions to run an acrylamide gel afterwards.

Run a 12% acrylamide gel at 200 V to evaluate all the samples you just generated: Lysis sample, Soluble fraction, Insoluble fraction, Flowthrough, 1st washing step, 2nd washing step and Eluted fraction.

### DAY 3-B: Protein expression in medium scale

2d

10m

Inoculate 2.5 mL from the pre-inoculum to an Erlenmeyer flask with 250 mL LB medium and [M] 0.05 mg/mL Kanamycin, use 4 flasks to obtain 1L of cell culture. Incubate at 3 220 rpm, 37°C until OD<sub>600</sub> = 0.5 - 0.6 (3-4 hours).

- Add IPTG to a final concentration of [M] 0.5 millimolar (mM) to each flask and incubate at \$\cdot 220 \text{ rpm, } 18^{\cdot C}, 16:00:00 \cdot \text{.}
- Centrifuge the cell culture 4000 rpm, 4°C, 00:20:00. Discard the supernatant. At this point, you may store the 1-2 grams of cell pellet at -20°C until you are ready to run the purification.

20m

### **DAY 4-B: Cells Lysis**

- Incubate the cells at 5 220 rpm, Room temperature , 00:20:00
- Sonicate on ice until the lysate turns translucid. Use 5 cycles of . Then 00:15:00 power OFF, with the tube on ice.

Centrifuge 6000 rpm, 4°C, 00:20:00 to separate the insoluble fraction (pellet) from the soluble fraction. Transfer the soluble fraction to a new and clean tube on ice. Collect small fractions of each one to run an acrylamide gel afterwards.

#### **DAY 4-B: Protein Purification with FPLC**

1d

- Prepare the 5 mL HisTrap column in the FPLC system. Wash the tubes, pumps system and the column with 7 column volumes (c.v.) of distilled and filtrated water. Then equilibrate the column with 7 c.v. of **Binding buffer**.
- Load the soluble fraction to the FPLC system at a flow of 1 mL/min. Collect a small fraction of each step and signal change to run an acrylamide gel afterwards. Wash the column with 5 c.v. of **Binding buffer**, until the UV and conductivity signal stabilizes.
- Washing: Load the column with 17% of pump B (Elution Buffer), which is equivalent to ~100 mM Imidazole, until the signal stabilizes.
   Elution: Load the column with 48% of pump B (Elution Buffer), which is equivalent to ~250 mM

**Elution:** Load the column with 48% of pump B (**Elution Buffer**), which is equivalent to ~250 mM Imidazole, until the signal stabilizes. Collect the elution in 8 mL tube fractions and in case the UV signal changes. Then load the column with 3 c.v. of 100% of pump B (**Elution Buffer**), which is equivalent to 500 mM Imidazole, until the signal stabilizes.

- Wash the column for storage. Wash the FPLC system with distilled and filtrated water. Load the column with 7 c.v of distilled and filtrated water. To storage the column, load it with 5 c.v. of ethanol 20% and storage it at 4°C. Finally, remove the rest of the water from the system with ethanol 20% and keep the system with it until next use.
- Determine the fractions with the UvsX recombinase by running a SDS-PAGE in a 10% acrylamide gel. The UvsX recombinase weights ~44 kDa.
- 36 Concentrate the eluted fractions with the protein with an
  - Amicon Ultra-15 Centrifugal Filter Unit **Emd Millipore Catalog #UFC910024**Reconstitute the concentrate so it is stored with the components detailed in **Storage Buffer** to decrease the Imidazol to 20 mM or less. Add glycerol to a 20%, homogenize, make aliquots of Δ 400 μL of the protein and storage them at -80°C.

