Protein Expression Protocol for the Catalytic Domain of an Adenylate Cyclase Membrane-Anchored by a Vibrio Quorum Sensing Receptor
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[Abstract] The direct regulation of a mycobacterial adenylate cyclase (Rv1625c) via exchange of its membrane anchor by the quorum sensing receptor CqsS (Vibrio harveyi) has recently been reported (Beltz et al., 2016). This protocol describes the expression and membrane preparation for these chimeric proteins.

Keywords: Adenylate cyclase (AC), Quorum sensing (QS), CqsS, Membrane protein, Protein expression, French press

[Background] Membrane-delimited mammalian adenylate cyclases (ACs) are class IIIa ACs. Regulation is indirectly via stimulatory (or inhibitory) Gα-proteins which are released intracellularly upon extracellular stimulation of G-protein-coupled-receptors (GPCR) by first messengers. ACs generate the universal second messenger cAMP using ATP as a substrate. The size of the two hexahelical membrane domains in vertebrate ACs by far exceeds the requirements for a simple membrane anchorage. Yet, regulatory features of this intrinsic membrane anchor/receptor domain are unknown. To investigate a potential function the canonical class IIIa AC Rv1625c from Mycobacteria was chosen which can be easily expressed in bacteria (Guo et al., 2001 and 2005) in contrast to mammalian AC isoforms. We replaced the hexahelical membrane anchor of the Rv1625c AC by the receptor domain of the hexahelical quorum sensing (QS) receptor from V. harveyi, CqsS, to examine whether we can confer a direct regulation of the AC by the QS-ligand ‘cholera autoinducer-1’, CAI-1. The design of the QS-receptor and the class IIIa membrane anchors are highly similar, i.e., minimal transmembrane α-helices and exceptionally short connecting loops.

We have demonstrated a direct regulation of a class IIIa AC by an extracellular signal. This considerably supports the hypothesis of a receptor function for the membrane anchor. Indeed, it raises the possibility that in addition to the well-established indirect GPCR-Gα-protein regulation of mammalian ACs a second, rather different set of signals directly impinge on this most important enzyme.

Materials and Reagents

A. Expression

1. Fisherbrand™ syringe filter, 25 mm, 0.22 μm (Thermo Fisher Scientific, Fisher Scientific, catalog number: 09719A)

Note: This product has been discontinued.
2. Glycerol stock of *Escherichia coli* BL21 (DE3) (F*ompT* hsdS*B* [rA'mB'] *gal dcm* [DE3]) transformed with plasmid pQE80L or pETDuet-3:
   a. pQE80L [QIAGEN, Δ*Xho*I, Δ*Nco*I]
      Encodes a lacIq repression module; N-terminal RGS-His6-tag
   b. pETDuet-3 (Figure 1)
      MCS1 of pETDuet-1 [Novagen] is converted to MCS of pQE30 [QIAGEN]
      MCS1: N-terminal RGS-His6-tag, MCS2: C-terminal S-tag

*Note: Expression of both plasmids is controlled by an IPTG-inducible T5 (pQE80L) or T7 (pETDuet-3) promoter. pQE80L has only one MCS. If you want to co-express a second protein from the same plasmid, pETDuet expression vectors are suitable.*

3. Ampicillin sodium salt (Carl Roth, catalog number: K029)
4. IPTG (isopropyl β-D-thiogalactopyranoside) (AppliChem, catalog number: A1008)
5. LB broth (Lennox) (Carl Roth, catalog number: X964)
6. Ampicillin solution (see Recipes)
7. IPTG solution (see Recipes)
8. LB-medium (see Recipes)
Figure 1. Comparison of the two MCS of pETDuet-1 (A) and pETDuet-3 (B)

Note: This sequence map is generated using the program DNA5 (https://pga.mgh.harvard.edu/web_apps/web_map/start).
B. Cell harvesting
   1. Culture from expression
   2. Tris (AppliChem, catalog number: A1086)
   3. HCl (37%)
   4. EDTA (Sigma-Aldrich, catalog number: E5134)
   5. Wash-buffer (see Recipes)

C. Membrane preparation
   1. 1.5 ml Eppendorf tube
   2. Frozen cell pellet from expression
   3. Thioglycerol (Sigma-Aldrich, catalog number: M1753)
   4. Sodium chloride, NaCl (EMD Millipore, catalog number: 106404)
   5. cOmplete™ EDTA-free protease inhibitor (Roche Diagnostics, catalog number: 05056489001)
   6. Glycerol (EMD Millipore, catalog number: 104094)
   7. Liquid N₂
   8. Lysis-buffer (see Recipes)
   9. Membrane-buffer (see Recipes)

D. Data analysis
   1. Tris (AppliChem, catalog number: A1086)
   2. HCl (37%)
   3. Sodium dodecyl sulfate, SDS (Sigma-Aldrich, catalog number: 71729)
   4. β-mercaptoethanol (EMD Millipore, catalog number: 805740)
   5. Glycerol (EMD Millipore, catalog number: 104094)
   6. Bromophenol blue (Sigma-Aldrich, catalog number: B8026)
   7. RGSHis-antibody (QIAGEN, catalog number: 34650)
   8. S-tag-antibody (EMD Millipore, catalog number: 71549)
   9. Protein marker IV prestained (VWR, catalog number: 27-2110)
   10. Protein marker I unstained (VWR, catalog number: PEQL27-1010)
   11. ECL Plex goat-anti-mouse IgG-Cy3-antibody (GE Healthcare, catalog number: PA43010)
   12. 4x SDS-loading dye (see Recipes)

Equipment

A. Expression
   1. 1 L Erlenmeyer flask
   2. Eppendorf Biophotometer
   3. Cuvettes
   4. Shaker incubator (37 °C and 22 °C)
B. Cell harvesting

1. Centrifuges (supplier: Thermo Fisher Scientific)
   a. Sorvall RC5B Plus
      Rotor: Kontron-Hermle A6.14 (Sorvall, catalog number: 202200)
   b. Heraeus Megafuge 1.0R
      Rotor: Heraeus Sepatech BS4402/A (Heraeus, catalog number: 3360)
2. 250 ml Sorvall metal rotor tubes (Sorvall, catalog number: 522)
3. 50 ml centrifuge tube (Greiner Bio One, catalog number: 227261)
4. Vortex Genie-2 (VWR, catalog number: 4445900)
5. Box with water-ice-mix
6. Freezer (-80 °C)

C. Membrane preparation

1. Vortex Genie-2 (Scientific Industries, model: Vortex-Genie 2)
2. French® Pressure Cell Press (SLM Instruments, SLM Aminco®, model: FA-078-E1)
   Alternative supplier: Glen Mills Inc. (USA) or G.Heinemann Ultraschall- und Labortechnik (Germany)
3. Aminco® French Pressure Cell (SLM Instruments, SLM Aminco®, catalog number: FA-073, serial number: 9110668)
   Alternative supplier: Glen Mills Inc. (USA) or G.Heinemann Ultraschall- und Labortechnik (Germany)
4. Centrifuges
   a. Heraeus Megafuge 1.0R
      i. Rotor: Heraeus Sepatech BS4402/A (Heraeus, catalog number: 3360)
   b. Beckmann L-60
      i. Rotor: Beckman Coulter, model: Type 50.2 Ti
5. Box with water-ice-mix
6. Polycarbonate ultracentrifuge-tubes
7. 7 ml Dounce Tissue Grinder (WHEATON, catalog number: 357542)
8. Freezer (-80 °C)

D. Data analysis

1. Ettan DIGE Imager (GE Healthcare, catalog number: 63005642 or 29-0834-61)

Software

1. Program DNA5 (https://pga.mgh.harvard.edu/web_apps/web_map/start)
Procedure

A. Expression
1. Inoculate a flask containing 200 ml LB-medium and 200 µl ampicillin (final concentration: 100 µg/ml) with approximately 5 ml overnight culture with the desired construct (see Materials and Reagents A1. point) to an OD_{600} of 0.1.
2. Incubate the culture under shaking (200 rpm) at 37 °C up to an OD_{600} of 0.2-0.3 (approx. 45-90 min).
3. Lower temperature to 22 °C (shaker incubator).
4. At an OD_{600} of 0.4-0.6 induce expression by 500 µM IPTG (100 µl 1 M IPTG/200 ml culture).

B. Cell harvesting
1. Harvest the cells at an OD_{600} of 2.0-2.8 (120-150 min after induction).
   Collect cells at 3,200 x g for 10 min at 4 °C (Sorvall centrifuge).
2. Add 25 ml of wash-buffer (4 °C) to the pellet, suspend by vortexing and pellet at 4,300 x g for 30 min at 4 °C (Heraeus centrifuge).
3. Discard supernatant and store cells at -80 °C or continue with the membrane preparation.

C. Membrane preparation
1. Thaw frozen cells on ice and suspend in 25 ml of lysis-buffer (4 °C) by vortexing.
   Notes:
   a. The cell pellet should be completely dissolved to avoid clogging the outlet of the Aminco® French Pressure Cell.
   b. Add always the cOmplete™ EDTA-free protease inhibitor tablet just before using the lysis-buffer.
2. Lyse cells mechanically by French press (1,100 psi) twice (Figure 2).
   Notes:
   a. Aminco® French Pressure Cell should be kept pre-cooled at 4 °C.
   b. Open outlet of the Aminco® French Pressure Cell such that a flow drop by drop is visible.
   c. Make sure that samples are continuously cooled in ice-water.
3. Centrifuge homogenate for 30 min, 4 °C, 4,300 x g (Heraeus centrifuge) and discard pellet (cell debris).
4. Transfer supernatant to an ultracentrifuge-tube and pellet membranes at 100,000 x g, 4 °C for 60 min (Beckman L-60 centrifuge).
5. Decant supernatant, take up membranes (pellet) in 1-2 ml membrane-buffer and gently suspend in a homogenizer (Dounce Tissue Grinder). Transfer the membrane preparation into a 1.5 ml Eppendorf tube.
Note: The amount of membrane-buffer to be used depends on the size of the pellet. Suspend a pellet of approximately 1 cm in diameter at the bottom of the centrifuge tube in 2 ml membrane-buffer.

6. Freeze membrane preparation in liquid N₂ and store at -80 °C.

Figure 2. Short French press protocol. A. Equipment of the French Pressure Cell; B. Aminco French Pressure Cell (taken out of the fridge 4 °C); C. Attention! Before use: Grease the O-rings and back-up rings of the stamp with glycerol! Insert the stamp into the cell body so that the inner cell body surface is covered with glycerol as well! D. Place the cell body with the stamp upside down into the stand with the opening facing upwards. E. Place the flow valve into the hole. F. Fill in the cell solution (looks milky). G. Put the lid on top. H. Close the flow valve. I. Place the French Pressure Cell into the gadget and… (J) close the bracket. K. Start the French press by turning the hand gear on ‘high’ and (L) switch on the pump. M. When the French press set up 1,100 psi… (N) open carefully the flow valve such that… (O) a flow drop by drop is visible. P.
Attention! Stop in time so you can easily open the bracket and the stamp does not hit the ground!

Q. When you are done, flip the switch to ‘down’ and turn on the pump again. The hydraulic lift moves down and you can take out the French Pressure Cell. R. Your cells are lysed properly when you can see the scale through the tube. S. Put your sample back on ice for the next step.

**Data analysis**

The expression of the proteins is verified by SDS-PAGE (Laemmli, 1970) and Western blot. The isolated membranes are incubated in 4x SDS-loading dye at room temperature for at least 30 min prior to application to SDS-PAGE (do not boil sample). Dilute the sample (membrane preparation) in MilliQ H₂O to get 2.5-5 µg of protein in a final volume of 15 µl. Add 5 µl 4x SDS-loading dye. Load 20 µl of the mixture into one slot of the SDS-PAGE. The membranes are incubated for 1 h with each antibody (first antibody at 4 °C, second antibody at room temperature). The first antibody is either the RGSHis- (Figures 3A and 3C) or the S-tag-antibody (Figure 3B). In both cases, the ECL Plex goat-anti-mouse IgG-Cy3-antibody is used as a secondary antibody (dilution 1:2,500). Western blot evaluation is carried out using an Ettan DIGE Imager.

*Note: In contrast to soluble proteins, membrane proteins are NOT boiled (95 °C, 5-10 min).*
**Recipes**

1. **Ampicillin solution (100 mg/ml)**
   
   Dissolve 100 mg ampicillin in 1 ml MilliQ H₂O (filter to sterilize)

2. **1 M IPTG solution**
   
   Dissolve 238.3 mg IPTG in 1 ml MilliQ H₂O (filter to sterilize)

3. **LB-medium**
   
   Dissolve 20 g LB in 1 L demineralized H₂O (autoclave 20 min at 121 °C)
4. Wash-buffer
   50 mM Tris/HCl (pH 8.0 at room temperature)
   1 mM EDTA

5. Lysis-buffer
   50 mM Tris/HCl (pH 8.0 at room temperature)
   2 mM thioglycerol
   50 mM NaCl
   1 tablet cOmplete™ EDTA-free protease inhibitor/50 ml lysis-buffer

6. Membrane-buffer
   40 mM Tris/HCl (pH 8.0 at room temperature)
   1.6 mM thioglycerol
   20% glycerol (85%)

7. 4x SDS-loading dye
   130 mM Tris/HCl (pH 6.8)
   10% SDS
   10% β-mercaptoethanol
   20% glycerol (85%)
   0.06% bromophenol blue

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References


