### **Protocol: Isolation of Bacterial Membrane Vesicles**

# Protocol Notes:1

- Grow pseudomonas from frozen stock to provide supernatant for membrane vesicles
- Purpose is to isolate the bacterial membrane vesicles using ultracentrifugation
- This protocol is suitable for serum, BALF, and most other biological fluids.
- Threads of ultracentrifuge buckets must be coated with Beckman Spinkote, and the gaskets must be coated with Beckman Vacuum grease.
- Must refrigerate SW 41 Ti rotor buckets at 4 degrees C prior to running.

#### Materials:

- 1. Pseudomonas aeruginosa ATCC #27853, Location: Freezer -80 #2, Shelf 4
- 2. Gentamicin stock 50 mg/ml, Location: Refrigerator #20
- 3. Mueller Hinton Broth (MHB), Location: Refrigerator #20 (prepared)
  - Sigma-Adlrich Mueller Hinton Broth, Cat#70192, Lot#
  - Store prepared media at 4 C in dark
  - Store dehydrated powder in dry location, Location:
- 4. Flask 250 mL, clear sterile plastic
- 5. Inoculating loops, sterile plastic, single use
  - Fisherbrand, Cat#13-075-1
- 6. 0.45~&~0.22 micron cellulose acetate membrane filters
  - Osmonics?
- 7. 50-mL polystyrene conical tubes (Falcon tube)
- 8. Beckman SW 41 Ti rotor, serial# 97U 10110 (swing-bucket rotor)
- 9. Beckman SW 41 Ti 104.9 buckets (6)
- 10. Beckman Spinkote, Cat# 306812
- 11. Beckman Vacuum Grease Silicone, Cat# 335148
- 12. Beckman Coulter Polyallomer Thin-walled Centrifuge Tubes,  $14 \times 89$  mm, 13.2 mL capacity
  - Lot# Z11208SCA, Reorder# 331372

<sup>&</sup>lt;sup>1</sup>Authors: Created by Patil Injean on 2014-06-30; Modified by ALG 2014-07

- can sterilize acid or alkali, not alcohols
- 13. Beckman Optima L-70k Ultracentrifuge
- 14. Sorval Legend XTR Centrifuge
- 15. Sorval rotor F13-14-50cy
- 16. New Brunswick Scientific Incubator Shaker G25
- 17. Ice
- 18. HEPES buffer 50 mM, pH 6.8, prepared on N/A by N/A, Location: refrigerator #? in 250-mL glass reagent bottle
  - Sigma
- 19. NanoDrop ND-1000
- 20. (Micro) BCA protein assay reagent kit
  - Pierce
  - mirco optimized for 0.5-20 mcg protein ml<sup>-1</sup>
- 21. Trypticase Soy Agar (TSA) plate, Location: Refrigerator #20, BD Cat#B11043
  - 1. Dissovle 20 g into 500 mL ddH2O. Mix by swirling, microwave 3 minutes, mix by swirling, repeat 3-4 times until powder completely dissolved. Autoclave at 121 degrees C for 15 minutes. (IMED autoclave settings: easy cycle, liquid, 15 minutes) At this time, turn waterbath on to 55 degree C.
  - 2. Remove from autoclave, mix solution by swirling to avoid clump/chunks. Place in 55 degree C waterbath, lay out dishes in hood (save bags), pipette 20-25 mL per plate (make sure to swirl solution around to avoid solidifying). Let dry 10-20 minutes, cover, flip, store in original bags at 4 degrees C.

## Methods:

- 1. Plate frozen stock cultures of *Ps. aeruginosa* on trypticase soy agar (TSA) plates. Grow *Ps. aeruginosa* at 37 degrees C for approximately 24 hours.
- 2. Prepare and autoclave 1 liter of cation-adjusted Mueller-Hinton Broth medium (containing 1.23 mM calcium chloride dihydrate and 1.03 mM magnesium chloride hexahydrate).
- 3. Add 200 mL of Mueller-Hinton Broth to sterile 250-mL flasks
- 4. Using a sterile 10 mcl inoculation loop, scrape a 2 colonies of *Ps. aeruginosa* growth from agar plates (from Day 1) and transfer into 250-mL flask with MHB in it (Step 3). Grow broth cultures at 37 degree C for approximately 6 h with gentle shaking (New Brunswick Scientific Incubator Shaker G25)

- 5. Obtain Ps. aeruginosa cultures from the shaking incubator and remove a 1-ml aliquot from each 250/500-ml culture to check the respective optical densities. Late exponential phase Ps. aeruginosa is used ( $\sim 10^9$  cfu ml<sup>-1</sup>), usually at approximately six hours of incubation.
  - Based on this cfu, the OD would be 0.900
- 6. Add stock concentration gentamicin to 250/500-mL bacterial culture flasks to final concentration of 8 mcL/mL. Incubate gentamicin in the culture flask x30 minutes on orbital shaker at 37 degrees C.
  - 32 mcL for 200 mL culture volume
  - 16 mcL for 100 mL culture volume
- 7. Using a serological pipet, remove entire volume of supernatant required at one time, transferring to sterile 50-mL polystyrene conical tubes. Balance tubes with cold 50 mM HEPES using a scale.
  - 1. Fill tubes leaving only a few millimeters of empty space.
  - 2. Tubes must be weighed within 0.1 0.2 g of one another
- 8. Spin samples in Sorval Legend XTR, 6,000 g (6000 RPM with F13-14-50cy) for 30 minutes at 4 degrees C.
  - 1. This is saved as Program? on the Sorval Legend XTR centrifuge
  - 2. Supernatant after this step contains the membrane vesicles.
  - 3. Discard the pellet.
- 9. Filter supernatant sequentially through 0.45 and then 0.22 micron cellulose acetate filters to remove any bacteria.
- 10. Using a serological pipet, transfer filtered supernatant to ultracentrifuge tubes. Balance tubes with cold 50 mM HEPES using a scale.
  - 1. Fill tubes leaving only a few millimeters of empty space.
  - 2. Tubes must be weighed within 0.1 0.2 g of one another
- 11. Spin samples in ultracentrifuge, 150,000 g (?? RPM with SW 41 Ti) for 90 minutes at 4 degrees C.
  - 1. This is saved as Program? on the Optima L-70k Ultracentrifuge
  - 2. Pellet after this step contains the membrane vesicles. Mark location of pellet with pen to facilitate extraction.
  - 3. Discard the supernatant.
  - 4. Place volume in excess of 72 mL into 4 degrees C at once
  - 5. Note that only 72 mL can be processed at one time in the SW 41 Ti rotor
  - 6. Multiple passes may need to be done to process all of the supernatant

- 12. Use a serological pipet to carefully aspirate the supernatant from the samples. Use a pipetman to remove as much supernatant from the pellet as possible.
- 13. Resuspend membrane vesicle pellet in 50 mM HEPES buffer and using serological pipet transfer the suspension to 50-mL conical tubes. Balance tubes with cold 50 mM HEPES using a scale.
  - 1. We do a two spin wash of the vesicle pellet
  - 2. Fill tubes leaving only a few millimeters of empty space.
  - 3. Tubes must be weighed within 0.1 0.2 g of one another
- 14. Spin samples in Sorval Legend XTR, 16,000 g (9,750 RPM with F13-14-50cy) for 30 minutes at 4 degrees C.
  - 1. This is saved as Program? on the Sorval Legend XTR centrifuge
  - 2. Pellet after this step contains the membrane vesicles.
  - 3. Discard the supernatant.
  - 4. Wash 1 completed
- 15. Again, use a serological pipet to carefully aspirate the supernatant from the samples. Use a pipetman to remove as much supernatant from the pellet as possible.
- 16. Resuspend membrane vesicle pellet in 50 mM HEPES buffer and using serological pipet transfer the suspension to 50-mL conical tubes. Balance tubes with cold 50 mM HEPES using a scale.
  - 1. Fill tubes leaving only a few millimeters of empty space.
  - 2. Tubes must be weighed within 0.1 0.2 g
- 17. Spin samples in Sorval Legend XTR, 16,000 g (9,750 RPM with F13-14-50cy) for 30 minutes at 4 degrees C.
  - 1. This is saved as Program? on the Sorval Legend XTR centrifuge
  - 2. Pellet after this step contains the membrane vesicles.
  - 3. Discard the supernatant.
  - 4. Wash 2 completed
- 18. Using serological pipet, resuspend pellet in 200-400 mcL of 50 mM HEPES
- 19. Determine membrane vesicle protein content using a (Micro) BCA protein assay reagent kit using the NanoDrop.
- 20. Plate approximately 1/10 of each final pellet (~50 mcg from each pellet) on TSA plate and incubate at 37 degrees C x48h to ensure no bacterial colonies form.
  - If no colonies form, then membrane vesicle pellet can be considered sterile
- 21. Freeze membrane vesicles at -80 degree C in freezer #2, shelf #4.

## **References:**

- 1. Renelli2004 DNA-containing membrane vesicles of Pseudomonas aeruginosa PAO1 and their genetic transformation potential. (main protocol)
- 2. Ellis2010 Naturally produced outer membrane vesicles from Pseudomonas aeruginosa elicit a potent innate immune response via combined sensing of both lipopolysaccharide and protein components. (SDS-PAGE for LPS)
- 3. Bauman2006 Purification of outer membrane vesicles from Pseudomonas aeruginosa and their activation of an IL-8 response.
- 4. Kadurugamuwa1995 Virulence factors are released from Pseudomonas aeruginosa in association with membrane vesicles during normal growth and exposure to gentamicin: a novel mechanism of enzyme secretion. (original isolation technique, gel)
- 5. Huntley, J.F., Robertson, G.T., Norgard, M.V. Method for the Isolation of Francisella tularensis Outer Membranes. J. Vis. Exp. (40), e2044, doi:10.3791/2044 (2010). (isolation of membrane proteins themselves)