Report for project Pseudomonas (MIRA)

Experiment created on 13.04.2018 14:56.

ARES0001

Extraction of DNA from pseudomonas bacterial culture utilizing chemical lysis and phenol:chloroform separation

Task created on 13.04.2018 19:39.

■ Bead Beating & PC DNA Extraction

No due date Completed on 13.04.2018 20:11

Completed 02.28.18

Task tags: dnaextraction beadbeating

Created by Emily Reesey on 13.04.2018 20:11.

- ◆ Step 1: Materials Uncompleted
 - TE buffer
 - 20% Sarkosyl solution
 - 10mg/mL RNase A
 - 10mg/mL proteinase K
 - Phenol:chloroform:isoamyl alcohol: 25:24:1
 - Chloroform:isoamyl alcohol, 24:1
 - 3 M sodium acetate
 - 70% and 100% ethanol
 - Phase-lock light gel tubes

Created by Emily Reesey on 13.04.2018 20:11.

Step 2: Bacterial Culture Uncompleted

- 1. From a frozen stock, plate pseudomonas on MacConkey agar and grow overnight.
- 2. Select an individual colony and plate onto a new MacConkey plate. Grow overnight.

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Step 3: Lyse Bacteria Uncompleted

- 1. Scrape plates with sterile loop and add into tube with 300uL TE buffer.
- 2. Add 50uL of 20% Sarkosyl solution.
- 3. Vortex bead beating tubes for 5 minutes on high speed.
- 4. Spin at full speed for 5 minutes to pellet bacteria & beads.
- 5. Remove supernatant and add to a new tube.
- 6. Add 15uL RNase A (10mg/mL) and 15uL of proteinase K (10mg/mL). Briefly vortex to mix. Incubate at 37C for 30 minutes.
- 7. Use TE to bring the volume up to 600uL and transfer to a 2mL phase lock tube.

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Step 4: Phenol:Chloroform Extraction Uncompleted

- 1. Add 600uL of phenol phenol:chloroform:isoamyl alcohol to phase lock tube.
- 2. Gently mix on rotator until emulsion forms do not vortex.

- 3. Spin at high speed for 10 minutes.
- 4. Remove aqueous phase to new tube phase lock tube.
- Add 600uL TE to original phase-lock gel tube. Mix until emulsion forms and centrifuge at high speed for 10 minutes.
- 6. Add aqueous phase into the new phase lock tube from step 4.
- 7. Add 600uL of chloroform:isoamyl alchocol (24:1) to extracted aqueous phase.
- 8. Gently mix on rotator until emulsion forms, then spin at high speed for 10 minutes.
- 9. Remove aqueous phase to new tube.

Created by Emily Reesey on 13.04.2018 20:11.

Step 5: Precipitate DNA Uncompleted

- 1. Add 50uL of 3M sodium acetate (pH 5.2) and 3 volumes of cold 100% ethanol.
- 2. Spin at full speed for 5 minutes.
- 3. Decant off supernatant then wash pellet with 500uL of 70% EtOH
- 4. Spin at full speed for 5 minutes.
- 5. Remove ethanol and dry pellet in 40C dry block for 10 minutes do not over dry the pellet.
- 6. Resuspend DNA in $50-100 \mu$ of TE or EB buffer and store at 4C-do not freeze DNA.
- 7. Quantify DNA on Qubit and Nanodrop.