Spectrophotometry Measurements

Construct Group

Plasmid Group

Interlab

Cell Culture/Plating

Biobrick Group

Cyanobacteria Transformation Group

Experimental Verification

Plasmid & Construct Design Group

Week 6

July 9, 2018

Constructs Group (Matt)

- Resuspension of primers (following IDT protocol)
 - 2991 cscB combo end \rightarrow 964 μ L of autoclaved milli-Q water
 - 2991 sps combo start \rightarrow 732 μ L of autoclaved milli-Q water
- PCR of Q1 combo cscB (CSCB Cycle)
- PCR of Q1 lone cscB (CSCB Cycle)

Plasmid Group (Priya/Stephanie/Jennifer)

- Nanodrop of 1579 purified (used 2 μL)
 - $1579 \text{ PUR} = 4.9 \text{ ng/}\mu\text{L}$
- Gel purification again of 1579 (Monarch New England Biolab Kit)

Cell Culture/Plating (Matt/Elon/Karthik)

- Made glycerol stock
 - 2x in snap cap tube, 2x in screw cap tube for each 1414, 1579, 2991
 - each tube has 500 μL of 50% glycerol and 500 μL of E. coli.

July 10, 2018

Spectrophotometer Measurement (Natalie/Lukas)

- Done at 750 nm with 1500 μL of culture
- 6/13 split from 6/6 10 mL/ 100 BG-11 (left) A= .728
- 6/6/18 10 mL cyano per 100 BG-11 A=.813
- 6/6/18 redone 5/28 C A= 1.194
- $6/6/18\ 10\ \text{mL} \rightarrow 5/19\ \text{C A} = .677$
- UTEX Room temperature 6/13 (split) A= 1.066
- UTEX Room temperature 6/26 (split) A= 1.240
- UTE Collier 5/19 Culture sup. with 20 mL BG-11 A=1.492

Cell Culture/Plating (Matt/Elon)

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- Made 40 mL stock solution of 10% methanol and BG-11
 - 4 mL of methanol
 - 36 mL of BG-11
- Froze 4 tubes of cyanobacteria in the -80 °C freezer
 - Added 450 μLof 10% methanol and BG-11
 - Added 450 µL of UTEX room temperature 6/13 (split) into each of 4 tubes
- Plated 50 μ L of 6/13 split from 6/6 10 mL/100 BG-11 (left) on tryptic soy agar plates to see if there was contamination
- Plated 50 μ L of 6/6/18 10 mL \rightarrow 5/19 C on tryptic soy agar plates to see if there was contamination

July 11, 2018

Interlab (Jenn/Steph)

- Made 8 LB media agar plates with 80 mL of LB and agar
 - Added 2g of LB and .96 g of agar
 - 80 μL of chloramphenicol
 - 80 mL of milli-Q water
- Made 13 LB media agar plates with 115 mL of LB and agar
 - Added 2.75 g of LB media and 1.32 g of Agar
 - 115 μL of chloramphenicol
 - 115 mL of milli-Q water

Constructs Group (Karthik/Natalie)

- Ran a .7% agarose gel
 - Buffer of 350 mL of 1X TAE = 35 mL of 10X TAE and 315 mL of milli-Q water
 - 1. MW ladder 15 μL
 - 2. Q3 cscB 25 μL of product and 5 μL of loading dye
 - 3. Q3 sps 25 μL of product and 5 μL of loading dye
- Made a diamond dye dilution
 - 200 mL of 1X TAE
 - 20 mL of 1X TAE and 180 mL of milli-Q water
 - 20 μL of diamond dye

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- Added 20 mL ethanol to the wash buffer to gel extraction kit
- Gel extraction of Q3 cscB and Q3 sps (Monarch NEB Gel Extraction Kit)
 - Q3 cscB= .07 g = 70 mg \rightarrow added 280 μ L of gel dissolving buffer
 - Q3 sps = .07 g = 70 mg \rightarrow added 280 μ L of gel dissolving buffer
 - Nanodrop (1.5 µL used)
 - Q3 sps = $6.0 \text{ ng/}\mu\text{L}$
 - Q3 cscB = $15.8 \text{ ng/}\mu\text{L}$

Plasmid Group (Stephanie/Jenn/Priya/Sara)

- Checked if DNA evaporated from stocks of plasmid DNA
- Nano dropped all of the stock DNA
 - $1579 = 51.8 \text{ ng/}\mu\text{L}$
 - $1414 = 564.6 \text{ ng/}\mu\text{L}$ (DNA evaporated left with about 3 μL)
 - 2991 = 178.6 ng/ μ L
- Made 19 plates for HIFI assembly
 - Spectinomycin plates = 125 mL of LB and Spec
 - Ampicillin/Kanamycin plates = 55 mL of LB and Amp/Kan
 - 2991 5 plates of Spectinomycin
 - 1414 8 plates of Spectinomycin
 - 1579 6 plates of Ampicillin/Kanamycin

Cell Culture/Plating (Jenn/Stephanie)

- 6/13 split from 6/6 10 mL/100 BG-11 (left) and 6/6/18 10 mL \rightarrow 5/19 C plates showed contamination
- Plated the rest of the stock solutions to see if they were also contaminated

July 12, 2018

Constructs Group (Karthik/Natalie/Woody)

- Ran a .7% agarose gel with Diamond nucleic acid dye
 - None of the bands showed up

Plasmid Group (Stephanie)

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- Labeled plates made yesterday for HIFI assembly

Cell Culture/Plating (Stephanie)

Out of the cyanobacteria stock solution that were plated, contaminated cyanobacteria were discarded

Interlab (Lin/Matt/Natalie/Dominika)

- Started competent cell test (following iGEM protocol)
 - $10 \text{ pg/}\mu\text{L}$
 - $100 \text{ pg/}\mu\text{L}$
 - Incubated 2 plates at 20:00 at 37 °C and 200 rpm
- Started transformation (following iGEM protocol)
 - 8 plates with 100 μL of cells unspun
 - 8 plates with 100 μL of cells spun
 - Incubated 16 plates at 20:00 at 37 °C and 200 rpm

July 13, 2018

Constructs Group (Karthik/Matthew/Natalie)

- Made a gel and added 5 μL of diamond dye to the gels
 - 1. MW ladder
 - 2. MW ladder
 - Both ladders showed up but were very smeary
- Made another gel with a ladder and diluted diamond dye and shook it
 - Nothing showed up
- PCR of Q3 cscB and Q3 sps using NEB Q5 polymerase
 - Used 0.5 μLof each template, 1 μL of each primer

Interlab (Natalie/Stephanie/Lin)

- Made 160 mL of LB Media
 - 4 g of LB powder
 - 160 mL of milli-Q water
- Put around 10 mL per falcon tube

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- Transferred 2 colonies from each plates into liquid media
- Incubated the colonies overnight at 37 °C and 220 rpm