Spectrophotometry Measurements

Construct Group

Plasmid Group

Interlab

Cell Culture/Plating

Biobrick Group

Cyanobacteria Transformation Group

Experimental Verification

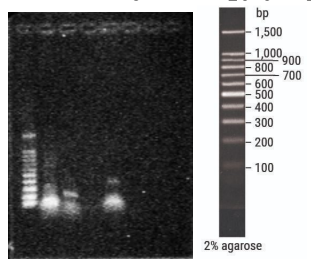
Plasmid & Construct Design Group

Week 9

July 30, 2018

Constructs Group (Natalie/Matthew/Karthik)

- Re-PCR Orig EYFP, isiAB sps, psbA2 sps, idiA sps, idiA



- 0.7% agarose gel
 - Lane 1: Promega 100 bp Benchtop ladder
 - Lane 2: idiA
 - Lane 3: psbA2
 - Lane 4: psbA2 sps (bad load)
 - Lane 5: idiA sps
 - Lane 6: psbA2 sps (bad load)
- PCR purify Q2 psbA2, Q2 rbc, opto EYFP
 - Nanodrop: only Q2 rbc was good at 48.8 ng/μL (rest had bad curves)
 - So re-PCR opto EYFP and psbA2 (psbA2 in gel above)

Cyanobacteria Transformation Group (Priya/Stephanie)

- Performed HiFi Assembly for better PCR cpc, cpc-560, and lone cscB
 - Plated transformants
- Inoculated four more colonies from the old cpc plate; five colonies from the weekend died in the 50 mL conical tube

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Plasmid Group

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Plasmid & Construct Design Group

July 31, 2018

Constructs Group (Karthik)

- Run PCR products orig EYFP. opto EYFP, isiAB sps (bad load) from yesterday on gel
- Re-PCR idiA, psbA2 sps, idiA sps, isiAB sps
- PCR purify psbA2
 - Nanodrop good (21.9 ng/μL) but low concentration
- PCR purify orig EYFP and Opto EYFP
 - Opto EYFP concentration 26.7 ng/μL
 - Orig EYFP had bad nanodrop curve

Cell Culturing/Plating (Natalie)

- Split the UTEX RM Temp. 7/23 (split) in half
 - Split 7/31 from UTEX room temp 7/23 = 37.5 mL of culture with 37.5 of BG-11 media
 - Supplemented the original culture (37.5 mL) with 37.5 mL of BG-11 media
- Split the UTEX Collier 5/19 room temperature in half
 - Split from room temp. Collier 7/31 = 25 mL of culture with 25 mL of BG-11 media
 - Supplemented the original culture (25 mL) with 25 mL of BG-11 media

Biobrick Group (Priya/Stephanie)

- Inoculated five colonies from new cpc, cpc-560, and cscB plates into aerated 15 mL culture tubes
- Dumped old cpc \rightarrow we had newer colonies/plates to work from and the O.D. was 0.000

August 1, 2018

Constructs Group (Woody/Matthew/Karthik)

- Purify orig EYFP
 - Nanodrop mediocre but concentration only 6.6 ng/µL, so save but try PCR again
- PCR troubleshooting

Spectrophotometry Measurements

Construct Group

Plasmid Group

Interlab

Cell Culture/Plating

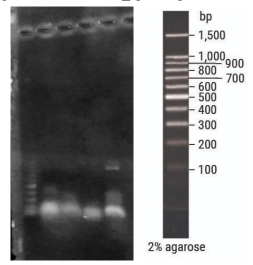
Biobrick Group

Cyanobacteria Transformation Group

Experimental Verification

Plasmid & Construct Design Group

- idiA_sps: primer concentration at ~200 ng/ μ L (half of expected) so increase volume to 2 μ L/reaction
 - Use results to adjust psbA2 sps PCR
- Q2 idiA: since primers working for other constructs, increased template volume to $2 \mu L/reaction$
- Re-PCR isiAB sps, orig EYFP using standard protocols
- idiA_sps, Q2 idiA, isiAB_sps, orig EYFP PCR products run on gel



- Lane 1: Promega 100 bp Benchtop ladder
- Lane 2: $idiA_sps \rightarrow eh$?
- Lane 3: isiAB sps \rightarrow bad
- Lane 4: Q2 idiA \rightarrow eh?
- Lane 5: orig EYFP \rightarrow good
- PCR purify orig EYFP, idiA sps, Q2 idiA
 - Only Q2 idiA gave good curve (concentration 37.8 ng/μL)
 - Other nanodrop curves were really bad
- PCR purify of Ladders using NEB kit with modifications: one with 6 uL MilliQ water and one with 6 uL MilliQ water and incubation at 37 deg C

Cell Culturing/Plating (Natalie)

- Split the UTEX room temperature 7/23 #2 (split) in half

Spectrophotometry Measurements

Construct Group

Plasmid Group

Interlab

Cell Culture/Plating

Biobrick Group

Cyanobacteria Transformation Group

Experimental Verification

Plasmid & Construct Design Group

- Split 8/1 from UTEX room temperature 7/23 #2 (split) = 37.5 mL of culture and 37.5 mL of BG-11 media
- Supplemented original culture with 37.5 mL of BG-11 media
- Split the UTEX Collier 7/23 culture 1 in half
 - 8/1 split from UTEX collier 7/23 Culture 1 = 25 mL of culture and 25 mL of BG-11 media
 - Supplemented original culture with 25 mL of BG-11 media

Cyanobacteria Transformation Group (Natalie/Lin)

- Made sodium bicarbonate solution
 - 3.36 grams of sodium bicarbonate powder
 - 40 mL of autoclaved milli-Q water
 - Left under UV light for 30 minutes
- Made 6 BG-11 agar plates (low antibiotic concentration) with 80 mL of media
 - 1.2 g agar powder
 - 80 mL of BG-11 media
 - $3.2 \mu L$ of streptomycin and $3.2 \mu L$ of spectinomycin
 - 800 µL of sodium bicarbonate solution
- Made 1 more BG-11 agar plate (low antibiotic concentration) with 20 mL of media
 - .3 g agar powder
 - 20 ml of BG-11 media
 - .8 μL of streptomycin and .8 μL of spectinomycin
 - 200 µL of sodium bicarbonate solution
- Made 10 BG-11 agar plates (high antibiotic concentration) with 120 mL of media
 - 1.8 g of agar powder
 - 120 mL BG-11 media
 - $24 \mu L$ of streptomycin and $24 \mu L$ of spectinomycin
 - 1200 μL of sodium bicarbonate solution

Biobrick Group (Natalie/Lin)

- Made 8 LB and CAM plates with 110 mL of media
 - 2.75 g of LB powder

Spectrophotometry Measurements

Construct Group

Plasmid Group

Interlab

Cell Culture/Plating

Biobrick Group

Cyanobacteria Transformation Group

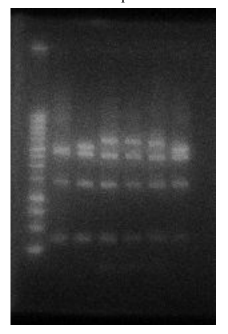
Experimental Verification

Plasmid & Construct Design Group

- 1.32 g of agar powder
- 110 mL of milli-Q water
- 110 μL of chloramphenicol
- Made 5 LB and CAM plates with 50 mL of media
 - 1.25 g of LB powder
 - .6 g of agar powder
 - 50 mL of milli-Q water
 - 50 μL of chloramphenicol

BioBrick Group (Priya/Stephanie/Dominika/Matt L)

- Digested lone cscB, linear 2991 with PvuII and ran gel
 - 1kb Promega Ladder
 - Lane 2 linear 2991, negative control/negative result
 - Lanes 3-8: Colonies 1-5
 - Colonies 1, 5 had negative results for the presence of cscB (appeared as if pAM2991 recircularized)
 - Colonies 2-4 had positive results



- Digested cpc and cpc-560 with KpnI

Spectrophotometry Measurements

Construct Group

Plasmid Group

Interlab

Cell Culture/Plating

Biobrick Group

Cyanobacteria Transformation Group

Experimental Verification

Plasmid & Construct Design Group

- Results inconclusive, need to use a different enzyme
 - Cause: re-circularization of 2991 vector in negative control has identical banding to positive result 2991
- Miniprepped cscB, cpc, cpc-560 DNA

August 2, 2018

Spectrophotometry Measurements (Natalie)

- Done at 750 nm with 1500 µL of culture
- UTEX room temp. 7/23 #1 (split) A=.588
- UTEX room temp. 7/23 #2 (split) A=.380
- Split 7/31 from UTEX room temp 7/23 #1 A= .685
- Split 8/1 from UTEX room temp. 7/23 #2 A=.819
- Split from UTEX Collier 7/23 culture 1 A=.970
- UTEX Collier 7/23 culture 1 A= .633
- Split from room temp. Collier 7/31 A= .991
- UTEX Collier 5/19 culture sup. With 25 mL BG-11 A=.913
- Syn. UTEX 2434 7/23 A=1.371

Cyanobacteria Transformation Group (Elon/Stephanie)

- Combined Split 7/31 from UTEX room temp 7/23 #1 with split 8/1 from UTEX room temp. 7/23 #2 into one flask
- Started transformation of lone cscB colonies 2-4, linear 2991, and negative control (Water) following Golden Protocol

Biobrick Group (Stephanie/Natalie)

- Performed HiFi Assembly with idiA, optimized EYFP, original EYFP constructs
- Plated idiA, psbA2, rbc, optimized EYFP, original EYFP constructs and negative control
- Digested cpc and cpc-560 with EcoR1, Kpn1
 - Cpc-560
 - Lane 1: 1kb Promega Ladder
 - Lanes 2-6: Cpc-560 colonies 1-5 all had a positive result for presence of insert

Spectrophotometry Measurements

Construct Group

Plasmid Group

Interlab

Cell Culture/Plating

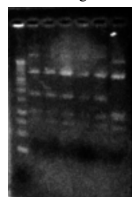
Biobrick Group

Cyanobacteria Transformation Group

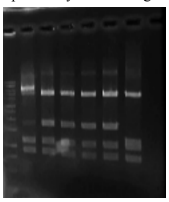
Experimental Verification

Plasmid & Construct Design Group

- Lane 7 - Digest of 1414-DNA, negative result (as expected)



- Cpc
 - Lane 1: 1kb Promega Ladder
 - Lanes 2-6: cpc colonies 1-5
 - Lane 7 Digest of Linear 1414, negative result (as expected)
 - Cpc colony 1 had a negative result, 2-5 were positive for presence of insert



- Miniprepped cpc and cpc-560 DNA

Constructs Group (Karthik/Matt/Woody)

- PCR troubleshooting for idiA sps/psbA2 sps, isiAB sps
 - idiA sps: increase template to 2 μL with increased primer at 2 μL
 - If it works, do same for psbA2_sps later
 - isiAB sps: increase primer to 2 μL

Spectrophotometry Measurements

Construct Group

Plasmid Group

Interlab

Cell Culture/Plating

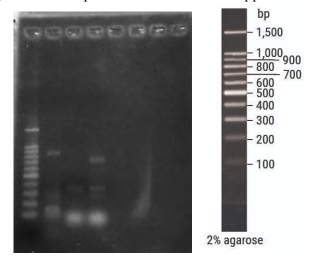
Biobrick Group

Cyanobacteria Transformation Group

Experimental Verification

Plasmid & Construct Design Group

- If doesn't work increase template next reaction
- Re-PCR orig EYFP
- Run gel with PCR products and see what happens



- Lane 1: Promega 100 bp Benchtop ladder
- Lane 2: isiAB sps \rightarrow good
- Lane 3: $idiA sps \rightarrow bad$
- Lane 4: orig EYFP \rightarrow good
- PCR purify isiAB sps and orig EYFP
 - Good nanodrop results for both
 - Concentrations: isiAB_sps 15.6 ng/μL, orig EYFP 26.2 ng/μL

August 3, 2018

Cyanobacteria Transformation Group (Natalie/Lin)

- Made 13 BG-11 Agar plates with 170 mL of media
 - 2.55 g agar
 - 170 mL BG-11
 - 34 μL streptomycin and 34 μL of spectinomycin
 - 1700 μL sodium bicarbonate solution

Cyanobacteria Transformation Group (Stephanie/Elon/Priya)

Spectrophotometry Measurements

Construct Group

Plasmid Group

Interlab

Cell Culture/Plating

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Cyanobacteria Transformation Group

Experimental Verification

Plasmid & Construct Design Group

- Plated cyanobacteria transformed from day before (lone cscB colonies 2-4, linear 2991, negative control / water)

Biobrick Group (Stephanie/Priya/MattL)

- Inoculated five colonies from rbc, idiA, psbA2, optomized eyfp, original eyfp

Cell Culture/Plating (Elon/Natalie)

- Made a 2% culture of cyanobacteria with sodium bicarbonate from UTEX Collier 7/23 Culture 1
 - No bicarb UTEX room temp #1= 1 mL of culture + 50 mL of BG-11 media
 - 5 mM bicarb UTEX room temp #1 = 1 mL of culture + 50 mL of BG-11 media + 250 μ L of sodium bicarbonate solution
 - 10 mM bicarb UTEX room temp #1 = 1 mL of culture + 50 mL of BG-11 media + 500 μ L of sodium bicarbonate solution
 - 20 m bicarb UTEX room temp #1 = 1 mL of culture + 50 mL of BG-11 media + 1000 μL of sodium bicarbonate solution

Spectrophotometry Measurements at 19:09 (Lin/Natalie)

- Done at 750 nm with 1500 µL of culture
- No bicarb UTEX room temp #1 A=0.000
- 5 mM bicarb UTEX room temp #1 A= 0.000
- 10 mM bicarb UTEX room temp #1 A= 0.000
- 20 mM bicarb UTEX room temp #1 A= 0.009

August 4, 2018

Spectrophotometer Measurements at 10:35 (Natalie)

- Done at 750 nm with 1500 μL of culture
- UTEX room temp 7/23 #1 (split) A= 0.726
- UTEX room temp 7/23 # 2 (split) A= 0.608
- 8/1 split from UTEX Collier 7/23 culture 1 A= 0.988
- UTEX Collier 7/23 Culture 1 A= 0.852
- Split from room temp. Collier 7/23 A= 1.034

Spectrophotometry Measurements

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Cyanobacteria Transformation Group

Experimental Verification

Plasmid & Construct Design Group

- UTEX Collier room temp. 5/19 A= 1.060

Spectrophotometry Measurements at 18:37 (Lin)

- Done at 750 nm with 1500 μL of culture
- No bicarb UTEX room temp #1 A=0.134
- 5 mM bicarb UTEX room temp #1 A= 0.136
- 10 mM bicarb UTEX room temp #1 A= 0.135
- 20 mM bicarb UTEX room temp #1 A= 0.127

Cell Culture/Plating at 11:00 (Natalie)

- Split the 8/1 split from UTEX Collier 7/23 Culture 1 in half
 - 8/4 from 8/1 (UTEX Collier 7/23 culture 1) = 25 mL of culture and 75 mL of BG-11 media
 - Supplemented original culture with 25 mL of BG-11 media
- Split the flask with split from room temp Collier 7/31 in half
 - 8/4 from 7/31 UTEX room temp. Collier = 25 mL of culture and 75 mL of BG-11 media
 - Supplemented original culture with 25 mL of BG-11 media
- Split UTEX Collier 5/19 culture (room temp.) in half
 - 8/4 room temp Collier (5/19) = 25 mL of culture and 75 mL of BG-11 media
 - Supplemented original culture with 25 mL of BG-11 media
- Supplemented UTEX rm. Temp 7/23 #2 (split)
 - Added 25 mL of BG-11 media to the culture
- Supplemented UTEX Collier 7/23 Culture 1
 - Added 15 mL of BG-11 media to the culture

Biobrick Group (Lin/Natalie/Matt/Priya/Stephanie)

- idiA, psbA2, rbc digested with EcoR1, Kpn1
 - Ran Gel Electrophoresis:
 - idiA colonies 1-5 all negative for presence of idiA

Spectrophotometry Measurements

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Plasmid Group

Interlab

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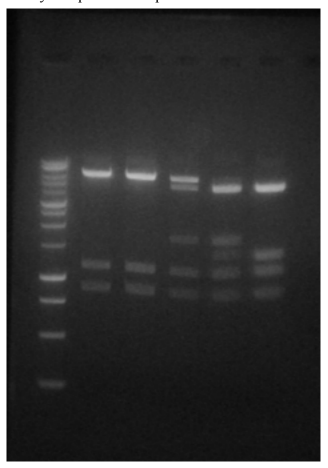
Biobrick Group

Cyanobacteria Transformation Group

Experimental Verification

Plasmid & Construct Design Group

- psbA2 colonies 1-2 negative for psbA2, 3-4 have a questionable result, colony 5 is positive for psbA2



- Rbc colonies 1-5 all positive for presence of rbc
- Original and optimized EYFP, 2991 digested with EcoR1, Spe1
 - Ran gel electrophoresis:
 - Failure, gels possibly punctured, will try again
 - Used the 1500 bp band from the promega ladder
- Gel Purification Troubleshooting
 - Original Protocol
 - 15μ L elution with modified protocol (lid taken off)

Spectrophotometry Measurements

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- $15 \mu L$ elution and incubate at 37 for a minute instead of room temp (modified protocol)

Cyanobacteria Transformation (Stephanie)

- Used cpc and cpc560 colonies 2-3
- Transformed all cyanobacteria from UTEX room temp 7/23 #1 (split)