# Lab Log

# Daniel Padfield

# 18/10/2017

- Autoclaved the soil again
- Autoclaved 1 L of DI millicule water
- Autoclaved 1 L of 0.6% soft agar
- Autoclaved a spoon to weigh out soil
- Got some  $SBW25\phi2$  from Floh. Around 1 mL at 7.26 x  $10^8$ .
  - Grow up overnight
  - Transfer 60  $\mu l$  of bacteria and 10  $\mu l$  of phage (around 7 x 10  $^6$  pfus) into 6 mL of KB agar
  - Should give a concentration of around 10<sup>8</sup> phage/mL
  - Done this in triplicate
- Grow up lacZ and WT strains overnight. Should give concentration of around ~10<sup>8</sup> cells in 60 µl.
- Do these in triplicate
- Added 60 μL of frozen overnight culture from first experiment (18/08/2017 lacZ and WT)

#### Retrospectively work out density of the overnight stocks and phage

### 19/10/2017

- Put 80g of soil into each 10cm x 10cm microcosm
  - Used autoclaved spoon
  - Placed scale in laminar flow hood (cleaned with ethanol before and after)
- Placed 5 mL ( $\sim 200~\mu$ l per microcosm) of lacZ and WT into separate 12 mL centrifuge tubes
  - Centrifuged for 15 minutes at max speed (~4500 r.p.m) on big centrifuge
  - Want to get to 5 mL per microcosm for inoculating (~ 125 mL in total)
  - Resuspended pellet into 2250  $\mu$ l, vortexed and placed 620  $\mu$ l, 620  $\mu$ l and 810  $\mu$ l into three different falcon tubes
  - Filled these three falcon tubes up to 40 mL, 40 mL and 45 mL respectively
    - \* This guaranteed the same concentration of sample in each falcon tube
  - Placed 5 mL of lacZ or WT strain into each microcosm
  - Froze (-80 °C) 900 μl of inoculate in 900 μl of glycerol (25% final concentration)
- In the no phage treatments, we added 5 mL of M9
- Added 5 mL of phage to phage treatments
  - Place 900 μl of bacteria + phage into three centrifuge tubes
  - Add 100 ul (10%) chloroform into each tube (under fume hood)
  - Vortex rigorously
  - Centrifuge for 2-3 minutes at full speed (minifuge)
  - Take out supernatant and placed in a single tube (took out 800 μl of each tube)
  - Put 40 mL of M9 into 6 tubes
  - Added 400 μl into each tube (~100 fold dilution from the initial stock)
  - Shake each tube and add 5 mL into each microcosm
- Place microcosms into the 26 °C incubator (Level 1 incubator room)

#### 22/10/2017

- setup 2 WT microcosms up for spot assays of phage
- used a crystal from T<sub>0</sub>
- autoclaved 2 L of KB agar. Put in small autoclave

#### 23/10/2017

- phage spot test  $10_{-1}$  to  $10_{-8}$ 
  - Floh's phage  $(7.26 \times 10^8)$
  - Phage we inoculated with
  - Extra phage tube (should have lower concentration)
  - blank plate
- plated  $T_0$  of WT and  $lacZ\ P$ . fluorescens

#### 24/10/2017

- phage spot test was no good (streaky)
- likely that we did not wait long enough for the spot to dry
- $\bullet$  set up more  $T_0$  bacteria overnight

# 25/10/2017

- phage spot assay again
- plating only 4 spots per plate instead of 8
- counted T<sub>0</sub> counts (plated 30 μl)
  - $lacZ @ 10^{-4}$ : 18
  - WT @ 10<sup>-4</sup>: 64

# 26/10/2017

- check phage spot, 10<sup>-6</sup> looks to be the correct dilution
- further counts are done by putting the 10 µl phage into the 1% bacteria soft agar
  - vortex and plate
  - leave to dry
  - incubate overnight

#### 30/10/2017

• grow up some T<sub>0</sub> bacteria overnight

# 31/10/2017

- serial dilution of phage to 10<sup>-6</sup>
- $50 \mu l bacteria + 10 \mu l phage + 5 ml soft agar$ 
  - vortex
  - pour
- set samples up for inoculated phage, floh's phage stock and our phage dregs (each in triplicate)
- in 28 °C incubator overnight

#### 01/11/2017

- looked at phage spots (10<sup>-6</sup> dilution)
- calculated how many phage are there

```
Phage\ concentration = \frac{number\ of\ plaques}{dilution\ factor\ \times\ volumeadded}
```

Volume added is almost always 10 µl. I wrote a mini function to calculate this

```
d <- data.frame(dregs = c(3,4,6), phage = c(6,2,4))

# back calculate number in there
# PFU/ml = plaque number / (dil fac * volume added)
plaq_num <- function(x, dilfac){return(x/(dilfac*0.01))}

d <- dplyr::mutate_at(d, c('dregs', 'phage'), plaq_num, dilfac = 10^-6)
knitr::kable(d)</pre>
```

ohage
e+08
e + 08
e+08

```
# concentration of phage added
mean(d$phage)

## [1] 4e+08

# concentration of other phage
mean(d$dregs)
```

#### ## [1] 433333333

- sampled all soil microcosms (after 13 days)
  - $-\sim 2g$  of soil into a 12 mL centrifuge tube (with  $\sim 7$  glass beads)
    - $\ast\,$  sample using the big end of a 5 mL pipette and weigh on scales under hood
  - add 10 mL of M9  $\,$
  - vortex for  $\sim 1$  minute
  - add 900 μl of sample to 900 μl of 50% glycerol and freeze at -80 °C (bacteria + phage)
  - make phage suspension for each tube
    - \* add 900 µl of sample to centrifuge tube
    - \* add 100  $\mu$ l of chloroform under fume hood
    - \* vortex
    - \* centrifuge at full speed for 4 minutes
    - \* put supernatant into separate tube
      - · samples 1-17 800 μl, samples 18-48 750 μl
- names of samples
- 1-12: WT no phage
- 13-24: lacZ no phage
- 25-36: WT + phage
- 37-48: lacZ + phage
- Autoclaved things
  - 2 empty 500 mL bottles
  - M9 salts x10 (500 mL)
    - \* converted the weight of  $Na_2HPO_4$  to the amount of  $Na_2HPO_4.7H_2O$  needed.
    - \* MW of  $Na_2HPO_4.7H_2O$  is 268. MW of  $Na_2HPO_4$  is 142.
    - \* to convert from weight of  $Na_2HPO_4$  to  $Na_2HPO_4.7H_2O$  is to multiply by  $\frac{268}{142}$

- centrifuge tubes.

sample	weight_g
1	2.1
2	2.0
3	1.9
4	2.0
5	2.0
6	2.0
7 8	2.0
9	2.0 2.1
10	1.9
11	2.2
12	2.1
13	2.1
14	1.9
15	2.2
16	2.2
17	1.9
18	2.2
19	2.0
20 21	2.3
$\frac{21}{22}$	2.3 2.2
23	1.9
24	2.1
25	2.1
26	2.2
27	1.9
28	2.0
29	2.2
30	2.3
31	2.0
32	2.2
33 34	$2.2 \\ 2.2$
$\frac{34}{35}$	2.0
36	2.0
37	2.3
38	1.9
39	2.2
40	1.9
41	2.2
42	2.0
43	2.0
48	
44 45 46 47	2.2 2.1 2.2 2.2 2.1

#### 02/11/2017

- set up phage spot tests of all soil phage suspensions against ancestral WT on soft agar.
  - just checking for presence/absence of phage
  - 5 mL per plate, 4 spots per plate, 12 spots in total
  - important to let the spot dry before moving
- poured plates (all Xgal)

#### 03/11/2017

- all no phage treatments had no phage and all phage treatments had phage
- plated all the bacteria + phage treatments at  $10^{-4}$  and  $10^{-5}$  (30 µl)
  - left on bench over the weekend

#### 06/11/2017

- counted all the plates
  - all still had bacteria in and were not obviously contaminated (Yays)
- picked 20 colonies from each sample and placed in KB media in 48 well plates
  - $-750 \mu l$  of KB media
  - used matchsticks
- grown statically for 2 days at 26  $^{\rm o}{\rm C}$

# 08/11/2017

- put 500 µl of 50% glycerol into each well (final concentration 20%)
- froze in the -80  $^{\circ}$ C
- autoclaved
  - -2x 800 mL hard agar
  - -2x 800 mL soft agar

#### 13/11/2017

• prep for sampling soil microcosms tomorrow. Labelled samples. Booked downstairs hood with power access.

#### 14/11/2017

- sampled all 48 microcosms
- froze 900 µl of bacteria + phage in 900 µl of 50% glycerol (final concentration equals 25%)
- phage sample in fridge
  - 1-12: 750 µl
  - 13-48: 600 µl
- setup soft agar plates of ancestral bacteria and did phage spot assays

sample	$weight\_g$
1	2.2
2	2.2
3	2.3
4	2.1

sample	weight_g
5	2.2
6	2.1
7	2.1
8	2.0
9	2.0
10	2.0
11	2.2
12	2.1
13	2.0
14	2.2
15	2.2
16	2.0
17	2.1
18	2.3
19	2.0
20	2.0
21	2.1
22	2.0
23	2.1
24	2.2
25	2.0
26	2.0
27	2.2
28	2.0
29	2.0
30	2.0
31	2.3
32	2.1
33	2.2
34	2.2
35	2.1
36	2.1
37	2.1
38	2.3
39	2.0
40	2.0
41	2.3
42	2.0
43	2.2
44	2.2
45	2.0
46	2.2
47	2.0
48	2.0

# 15/11/2017

• all phageless samples are still phageless, all phage samples still contain phage

# 16/11/2017

- plated all  $T_2$  samples  $-10^{-3} \text{ and } 10^{-4}$   $-30 \text{ } \mu \text{l of on plate}$
- used undiluted M9 x10 on Tuesday!!!
  made up 400 mL of M9 salts x10 for autoclave