Turner Lab Notebook

Callum Malcolm

Contents

Wednesday 15-03-2023 Cell Culture	3
N2 Growth Curve	
Γhursday 16-03-2023	4
N2/BLLW Growth Curve Day 1	4
Friday 17-03-2023	4
N2/BLLW Growth Curve Day 2	
Split Cells	4
Saturday 18-03-2023	5
N2/BLLW Growth Curve Day 3	5
Sunday 18-03-2023	5
N2/BLLW Growth Curve Day 4	
Monday 20-03-2023	5
N2 Growth Curve Day 5	
Split Cells	
Seed BLLW	
Fuesday 21-03-2023	5
Friday 24-03-2023	5
Seed Cells - N2/RBL1/JIRE	5
Monday 27-03-2023	6
Split Cells	
Wednesday 29-03-2023	6
Split Cells	
•	
Friday 31-03-2023	6
Split Cells	6
Monday 03-04-2023	6
Seed Cells - N2/RBL1/JIRE	
Seed EC50	6
Wednesday 05-04-2023	6
Seed Cells - N2/RBL1/JIRE	

Thursday 06-04-2023 Collect EC50 Plate	6 6
Tuesday 11-04-2023 Seed EC50_230411	7
Wednesday 12-04-2023 Cell Culture	
Tuesday 02-05-2023 Seeding Cells - N2/RBL1/RAMOS	8
Wednesday 03-05-2023 Split Cells	8
Thursday 04-05-2023	8
Monday 19-06-2023 Split Cells Seed EC50	
Saturday 20-05-2023	9
Tuesday 20-06-2023 RNA Extraction for SP Test 3 Sample Tapestation_230620 Reverse Transcription PCR Amplification of cDNA	11 11
Wednesday 21-06-2023 Tapestation_230619_SPTest3_cDNA	13
Thursday 22-06-2023 Collect EC50 Plate	13
Friday 23-06-23 Split Cells	
Monday 26-06-23 Collect EC50 Plate	14 14
Monday 24-07-2023 Seeding Cells - RAMOS/FDC	
Tuesday 25-07-2023 Seed Cells - N2-BC4/RBL1/BLLW/RAMOS-BC1	16
Wednesday 26-07-2023 Cell Culture	16
Thursday 27-07-2023 Split Cells - RAMOS	

Friday 28-07-2023	17
Friday 06-10-2023	17
Cell Culture	17
Made New Media: AR-2	17
Monday 09-10-2023	17
Cell Culture	17
Seeded A20	17
Seeded FDC	17
Tuesday 10-10-2023	18
Cell Culture	18
Seeded N4	
Seeded RAMOS-BC	18
Checked A20	18

Wednesday 15-03-2023

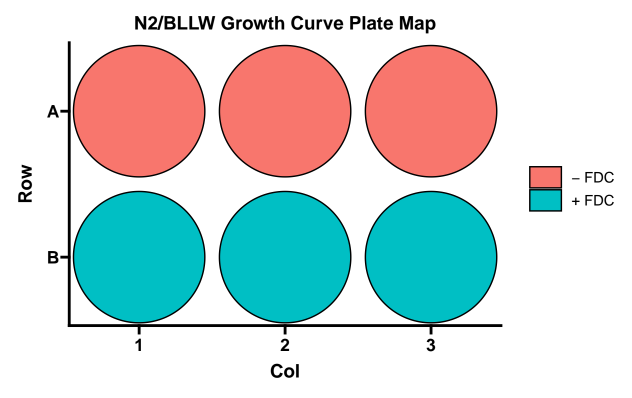
Cell Culture

- Given N2/RBL1/BLLW PDX line from Chris Steele

N2 Growth Curve

Looking to compare N2 growth on and off feeders 1. Seeded 2mL of N2/BLLW cell suspension at $0.25 \times 10^6 \text{cells/mL}$

96 Well Plate Map



Thursday 16-03-2023

N2/BLLW Growth Curve Day 1

- Cells gently mixed
- 30 μL cell suspension taken and counted
- Only counted 1 square and multiplied by 4

Cell Line	FDC Status	Count	Cell Number
N2	+ FDC	25	2000000
N2	+ FDC	26	2080000
N2	+ FDC	12	960000
N2	- FDC	15	1200000
N2	- FDC	14	1120000
N2	- FDC	6	480000
BLLW	+ FDC	6	480000
BLLW	+ FDC	6	480000
BLLW	+ FDC	5	400000
BLLW	- FDC	3	240000
BLLW	- FDC	3	240000
BLLW	- FDC	6	480000

Friday 17-03-2023

N2/BLLW Growth Curve Day 2

- $\bullet\,$ Cells gently mixed
- 30 μL cell suspension taken and counted
- Only counted 1 square and multiplied by 4

Cell Line	FDC Status	Count	Cell Number
N2	+ FDC	36	2880000
N2	+ FDC	28	2240000
N2	+ FDC	38	3040000
N2	- FDC	61	4880000
N2	- FDC	38	3040000
N2	- FDC	16	1280000
BLLW	+ FDC	3	240000
BLLW	+ FDC	5	400000
BLLW	+ FDC	5	400000
BLLW	- FDC	1	80000
BLLW	- FDC	0	0
BLLW	- FDC	1	80000

Split Cells

Cell Line	$\mathrm{Cells/mL}$	Cells/mL Seeded	CS volume	Final Volume	Seeded Cells/mL
N2 RBL1	8.8×10^5 8.6×10^5	$3x10^5$. $3x10^5$.	$\begin{array}{c} 6.8~\mathrm{mL} \\ 11.6~\mathrm{mL} \end{array}$	$12~\mathrm{mL}$ $12~\mathrm{mL}$	$3x10^{5}$ $5x10^{5}$

Saturday 18-03-2023

N2/BLLW Growth Curve Day 3

• Counted by Chris

Sunday 18-03-2023

N2/BLLW Growth Curve Day 4

• Counted by Chris

Monday 20-03-2023

N2 Growth Curve Day 5

- BLLW were all dead, experiment stopped
- Cells gently mixed
- 30 μ L cell suspension taken and counted
- ullet Only counted 1 square and multiplied by 4

Cell Line	FDC Status	Count	Cell Number
N2	+ FDC	146	11680000
N2	+ FDC	112	8960000
N2	+ FDC	132	10560000
N2	- FDC	112	8960000
N2	- FDC	155	12400000
N2	- FDC	141	11280000

Split Cells

- Split cell lines N2/RBL1/JIRE
- Dirty split 1:2

Seed BLLW

- Given BLLW aliquot from Chris Steel
- Thawed and resuspended in 12 mL media
- Added to T75 pre-seeded with FDC cells (from Chris Steel)

Tuesday 21-03-2023

Large Infection in TC

- N2 Growth Curve stopped

Friday 24-03-2023

Seed Cells - N2/RBL1/JIRE

- Vials given by Chris Steel
- Thawed and resuspended in 12 mL media
- Added to T75 pre-seeded with FDC cells (from Chris Steel)

Monday 27-03-2023

Split Cells

- Split cell lines N2/RBL1/JIRE
- Dirty split 1:3

Wednesday 29-03-2023

Split Cells

• Split by Chris Steel.

Friday 31-03-2023

Split Cells

- Split cell lines N2/RBL1/JIRE
- Dirty split 1:4

Monday 03-04-2023

Large Infection in TC

Seed Cells - N2/RBL1/JIRE

- Vials given by Chris Steel
- Thawed and resuspended in 12 mL media
- Added to T75 pre-seeded with FDC cells (from Chris Steel)

Seed EC50

DNW - N2/JIRE/RBL1

- Vin/Dactolisib

Wednesday 05-04-2023

Seed Cells - N2/RBL1/JIRE

- Vials given by Chris Steel
- Thawed and resuspended in 12 mL media
- Added to T75 pre-seeded with FDC cells (from Chris Steel)

Thursday 06-04-2023

Collect EC50 Plate

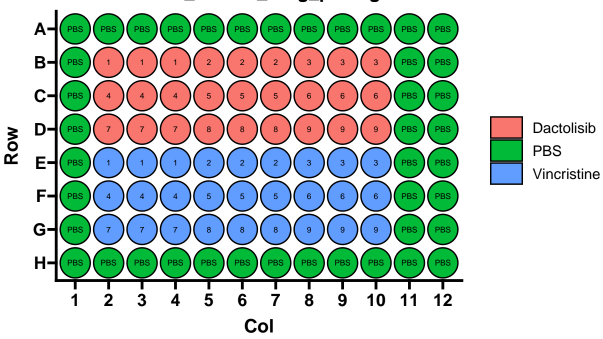
- Plate ID: EC50_230619_Plate1
- DNW

Tuesday 11-04-2023

Seed EC50_230411

96 Well Plate Map

EC50_230411_drug_plating



Drug Dilutions

Drug	Dilution	Concentration	Units	Starting_Concentration
Dactolisib	1	20	uM	20000
Dactolisib	2	2	uM	2000
Dactolisib	3	0.2	uM	200
Dactolisib	4	0.02	uM	20
Dactolisib	5	0.002	uM	2
Dactolisib	6	0.0002	uM	0.2
Dactolisib	7	2e-05	uM	0.02
Dactolisib	8	2e-06	uM	0.002
Dactolisib	9	2e-07	uM	0.0002

Drug	Dilution	Concentration	Units	Starting_Concentration
Vincristine	1	30	nM	30000
Vincristine	2	3	nM	3000
Vincristine	3	0.3	nM	300
Vincristine	4	0.03	nM	30
Vincristine	5	0.003	nM	3
Vincristine	6	0.0003	nM	0.3
Vincristine	7	3e-05	nM	0.03
Vincristine	8	3e-06	nM	0.003
Vincristine	9	3e-07	nM	0.0003

^{• 3} plates seeded

⁻ Plate 1: N2

- Plate 2: JIRE
- Plate 3: RBL1

Wednesday 12-04-2023

Cell Culture

• Thaw RBL1

Side Population - $N2_230412$

- Hoescht
 - -2.5 / 2.75 / 3
 - DNW

Tuesday 02-05-2023

Seeding Cells - N2/RBL1/RAMOS

1. Thawed N2, RBL1, and RAMOS into T75 flasks Note: N2 and RBL1 flaskes seeded with feeders

Wednesday 03-05-2023

Split Cells

Cell Line	$\mathrm{Cells/mL}$	Cells/mL Seeded	CS volume	Final Volume	Last day split	P number
N2	$8.93 \text{x} 10^5$	$3x10^{5}$.	$6.71~\mathrm{mL}$	12 mL	Tues 02-05-23	2
RBL1	$7.73 \mathrm{x} 10^5$	$3x10^{5}$.	$7.76~\mathrm{mL}$	$12 \mathrm{mL}$	Tues $02-05-23$	2

- 2. Expanded N2-Barcodes to T175
- 3. Seeded feeders into 96-well
 - a. Diluted feeders to 3.2×10^4 cells/mL
 - b. Add $100\mu L$ cell suspension to each well

Thursday 04-05-2023

Monday 19-06-2023

Split Cells

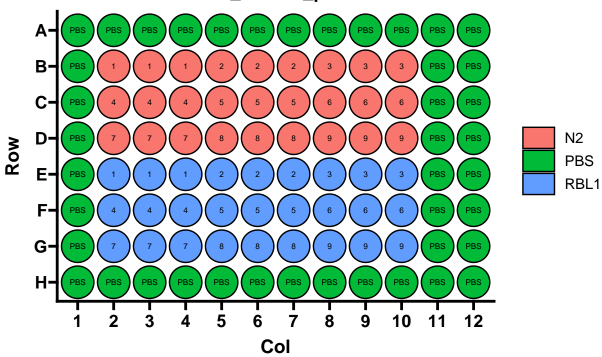
- Split N2/RBL1/JIRE
- $\bullet\,$ Added 10 mL media to BLLW
 - Lots of dead cells observed

Seed EC50

- Plate ID: EC50_230619_Plate1
- Cylophosphamide (Stock: 20mM) EC50 for N2 and RBL1

96 Well Plate Map

EC50_230619_plate1



Drug Dilutions

Drug	Dilution	Concentration	Units	Starting_Concentration
Cyclophosphamide	1	20	uM	20000
Cyclophosphamide	2	2	uM	2000
Cyclophosphamide	3	0.2	uM	200
Cyclophosphamide	4	0.02	uM	20
Cyclophosphamide	5	0.002	uM	2
Cyclophosphamide	6	0.0002	uM	0.2
Cyclophosphamide	7	2e-05	uM	0.02
Cyclophosphamide	8	2e-06	uM	0.002
Cyclophosphamide	9	2e-07	uM	0.0002

Saturday 20-05-2023

${\bf Tuesday~20\text{-}06\text{-}2023}$

RNA Extraction for SP Test 3 Sample

- Sample collected:
- Low number of cells so used Pico-Pure Kit
- 1 sample (n = 1)

Note before starting:

- Pre-heat block to 42C
- Pre-make DNAse treatment master mix (1.1n)

Component	Volume	MM volume
DNAse I Stock	$5 \mu L$	$5.5 \ \mu L$
DNA Disgestion Buffer	$35 \mu L$	$38.5 \ \mu L$

RNA Extraction Pico-Pure Protocol

- 1. Extract cells with 100 μ L of Extraction Buffer (XB)
 - Resuspend the cell pellet gently by pipetting
 - DO NOT VORTEX
- 2. Incubate at 42C for 30min
- 3. Centrifuge sample at 3000xg for two minutes
- 4. Collect supernatant
 - Can stop here and freeze RNA at 70C
- 5. Pre-condition the RNA Purification Column
 - Add 250 μ L Conditioning Buffer onto the purification column filter membrane
 - Incubate RNA Purification Column with CB for 5min at RT
 - Centrifuge purification column at 16000xg for 1 min
- 6. Pipette 100 μ L of 70% Ethanol (EtOH) into cell extract
 - Mix well by pipetting
 - DO NOT VORTEX
- 7. Add cell extract to column
- 8. Centrifuge column for 2 minutes at 100 x g and then 16000 x g for 30s
- 9. Pipette 100 µL Wash Buffer 1 (W1) into the purification column and centrifuge for 1min at 8000 x g
- 10. Add 40 μ L of DNAse treatment master mix to purification tube
- Add directly onto membrane
- 11. Incubate at RT for 15 min
- 12. Pipette 40 μ L Wash Buffer 1 (W1) onto purification column membrane and centrifuge at 8000 x g for 15 seconds
- 13. Pipette 100 μ L Wash Buffer 1 (W2) into purification column membrane and centriguge for one minute at 8000 x g
- 14. Pipette 100 μ L Wash Buffer 1 (W2) into purification column membrane and centriguge for two minutes at 16000 x g
 - Check if any wash buffer remains in column and centrifuge again at $16000 \times g$ for one minute to clear all the liquid

- 15. Transfer purification column to new 0.5mL tube
 - Insert purification column/0.5mL assembly into 1.5mL tube for centrifugation
- 16. Pipette EB directly onto membrane of the purification column
 - Gently touch tip to surface of membrane to ensure maximum absorption of EB into membrane
 - Use 11 μL to 30 μL EB
- 17. Incubate purification column for one minute at room temperature
- 18. Centrifuge column for one minute at 1000 x g to distribute EB in column
- 19. Centrifuge column for one minute at 16000 x g to elute RNA
- 20. Started PCR and completed Tapestation

Tapestation_230620

High-Sensitivity RNA ScreenTape Protocol

Results

- Inconclusive, proceeded with PCR

Reverse Transcription

Notes Before Starting:

- Label PCR strip tubes
- Thaw RT reagents
- Uses WILDseq specific RT primer
- Dilute RT primer to $2\mu M$
 - Do this in a PCR hood
 - IMPORTANT Do not mix primers and carefully note which RT primer is added to which sample
- Make RT and ExoI master mixes (MM) before starting
 - MM amounts are 1.1n

Protocol:

Note: Used "rt" protocol in callum folder on thermocycler for heating steps

- 1. In PCR strip tubes diluted 1-5ug RNA in a total volume of 10 μ L of RNAse/DNAse-free water
- 2. Added 1 ul of RT primer
- 3. Added 1 ul of dNTPs
- 4. Denatured at 65 C for 5 mins in PCR machine
- 5. Added straight to ice for 2 mins
- 6. Spun to get liquid to bottom of tube
- 7. Prepared master mix of RT enzyme and buffers (Made prior to starting)

Component	Volume	MM volume
5x SSIV Buffer	$4~\mu L$	$4.4~\mu L$
SSIV RT	$1~\mu L$	$1.1~\mu \mathrm{L}$
$100~\mathrm{mM}~\mathrm{DTT}$	$1~\mu { m L}$	$1.1~\mu\mathrm{L}$

Component	Volume	MM volume
RNAse Out	$1~\mu { m L}$	$1.1~\mu\mathrm{L}$

- 8. Added 7 μL of RT MM prepared above to each sample and mix
 - Spun briefly to get liquid to bottom of the tube
- 9. In PCR Machine: incubated at 53C for 10 mins followed by 80C for 10 mins
- 10. Added 3 μ L of ExoI MM to each sample

Component	Volume	MM volume
Thermolabile Exonuclease I NEBuffer r3.1*	$\begin{array}{ccc} 1 \ \mu L \\ 2 \ \mu L \end{array}$	$1.1 \ \mu L \\ 2.2 \ \mu L$

- Most PCR buffers are compatible
- 11. In PCR Machine: heat at $37\mathrm{C}$ for 4 mins followed by $80\mathrm{C}$ for 1 min
 - Spin briefly to get liquid to bottom of the tube
- 12. Add 1 μL of RNAse H and incubate at 37C for 20 mins
- 13. Dilute final cDNA 1 in 2 with DNAse/RNAse-free water

PCR Amplification of cDNA

1. Prepare mastermix of below reagents at 1.1x the number of samples

Component	Volume	MM volume
Q5 Master Mix	$25 \ \mu L$	
$P5_R1_fwd (100\mu M)$	$2.5~\mu\mathrm{L}$	
P7_rev	$2.5~\mu\mathrm{L}$	
DNAse/RNAse H20	$13.5~\mu\mathrm{L}$	
Q5 polymerase	$0.5~\mu\mathrm{L}$	
Total Master Mix volume	40 μ L	
cDNA from above	$10~\mu\mathrm{L}$	

2. Perform PCR using the following parameters:

Steps	Time
Step 1: 98C	30s
Step 2: 98C	10s
Step 3: 61C	30s
Step 4: 72C	30s
Step 5: 72C	$2 \min$
Step 6: 12C	Hold

20-25 cycles of steps 2-4

Ran overnight and left at 12C

Wednesday 21-06-2023

$Tape station_230619_SPTest3_cDNA$

High-Sensitivity DNA ScreenTape Protocol

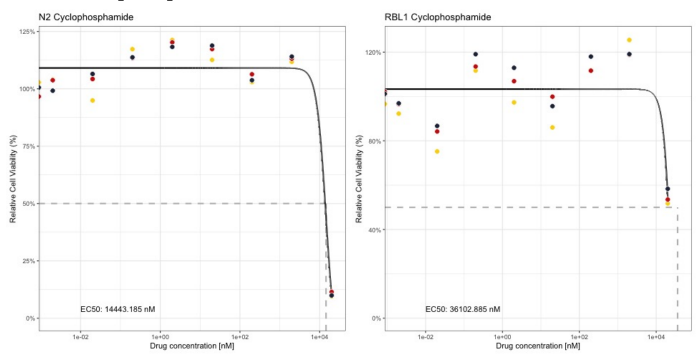
Results

- * Did not work, bands present but faint and not at 200bp
- * Potential solutions:
- Repeat PCR with increased cycles
- Repeat RNA extration with trizol protocol (Anna)

Thursday 22-06-2023

Collect EC50 Plate

- Plate ID: EC50_230619_Plate1



Friday 23-06-23

Split Cells

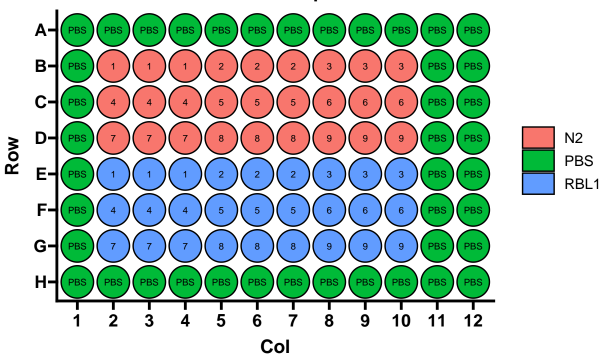
- 1. Threw out BLLW (all dead)
- 2. Split JIRE 1/2
- 3. Split N2/RBL1 1/4

Seed EC50

• EC50 Plate ID: EC50_230623_Plate1

96 Well Plate Map

EC50_230623_plate1



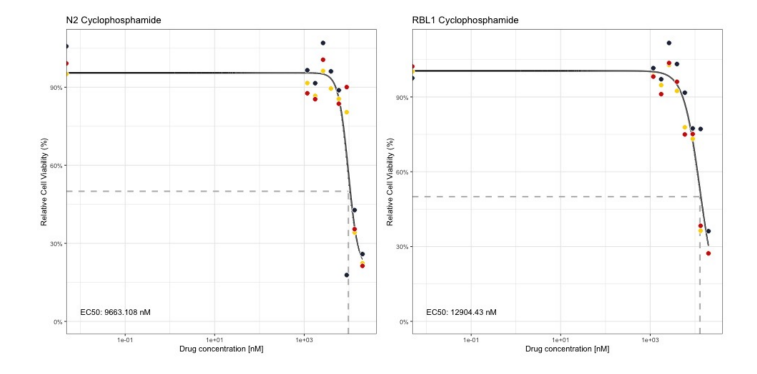
Drug Dilutions

Drug	Dilution	Concentration	Units	Starting_Concentration
Cyclophosphamide	1	20	uM	20000
Cyclophosphamide	2	13.3	uM	13300
Cyclophosphamide	3	8.89	uM	8890
Cyclophosphamide	4	5.93	uM	5930
Cyclophosphamide	5	3.95	uM	3950
Cyclophosphamide	6	2.63	uM	2630
Cyclophosphamide	7	1.76	uM	1760
Cyclophosphamide	8	1.17	uM	1170
Cyclophosphamide	9	0.78	uM	780

Monday 26-06-23

Collect EC50 Plate

- Plate ID: EC50_230623_Plate1



Break for writing MRes Report: 27/06/2023 - 24/07/2023

Monday 24-07-2023

Seeding Cells - RAMOS/FDC

Made New Media: AR-1

1. Made new media: AR-1

Solution	ID code	Volume	% Total volume
Advanced RMPI		500 mL	78%
Glutamax	—-	$6.5~\mathrm{mL}$	20%
Pen-Strep	—-	$6.5~\mathrm{mL}$	1%
FBS	—-	$128~\mathrm{mL}$	1%

2. Seeded Feeders

- 2 vial of FDC cells thawed
- Resuspended in 12 mL of media
- \bullet 6 mL of FDC cell suspension were added to 2 seperate T75 flasks (Seeding RBL1/BLLW on 25-07-2023)
- Remaining 12 mL of Media were added to T175 (Seeding N2 on 25-07-2023)
- 3. Seeded Ramos WILDseq Barcode Pool 2
- Thawed a vial of RAMOS pool 2 (previously barcorded DATE)
- Resuspended in 5 mL
- Spun down 90* for 5min
- Resuspended in 12 mL of AR-1 media
- Seeded into T75

Experimental Plans

- 1. Barcode BLLW/RBL1
- 2. Set up Drug pressure experiement for RAMOS barcoded cells
- EC50's for cyclophosphamide and methotrexate need to be completed
- Drugs to test (EC50):

Drug	Concentration	Replicates
DMSO	1%	3
Doxorubicin	$200\mathrm{nM}$	3
Methotrexate	$100\mathrm{nM}$	3
Vincristine	$1\mathrm{nM}$	3

- 3. Do Side Population Experiement for N2 Barcoded Cells
- Grow up sorted side pop to avoid issues with low cell numbers

Tuesday 25-07-2023

$Seed \ Cells - N2-BC4/RBL1/BLLW/RAMOS-BC1$

- Got vials from Chris
- Thawed, resuspended in 6 mL AR-1
- Spun down at 90* for 5 min
- Discarded supernatant and resuspended pellet in 12 mL of AR-1
- Added RBL1/BLLW to T75's pre-seeded with FDC (24-07-2023)
- Added RAMOS-BC1 to new T75
 - RAMOS-BC2 from 24-07-2023 was dead
 - Was taken from -80, New RAMOS vial from LN
- Added N2 to T125 seeded with FDC (24-07-2023)

Wednesday 26-07-2023

Cell Culture

- Checked cells, all flasks looked healthy
- Added media
 - N2 20ml of AR-1
 - RAMOS 10 mL of AR-1
 - RBL1 10ml of AR-1
 - BLLW 10ml of AR-1

Thursday 27-07-2023

Split Cells - RAMOS

- Split Ramos 1:2
- Added 10ml of AR-1 to N2
- Plan to split RBL1/BLLW tomorrow

Lucy Drug Randomizer

• Randomized drugs for Lucy Hare mouse experiment

Drug	Identification
Alectinib	A
Erda	В
Vehicle	$^{\mathrm{C}}$
${\bf Alectinib + Erda}$	D

Friday 28-07-2023

Drug	Dilution	Concentration	Units	Starting_Concentration
Cyclophosphamide	1	20	uM	20000
Cyclophosphamide	2	13.3	uM	13300
Cyclophosphamide	3	8.89	uM	8890
Cyclophosphamide	4	5.93	uM	5930
Cyclophosphamide	5	3.95	uM	3950
Cyclophosphamide	6	2.63	uM	2630
Cyclophosphamide	7	1.76	uM	1760
Cyclophosphamide	8	1.17	uM	1170
Cyclophosphamide	9	0.78	uM	780

Friday 06-10-2023

Cell Culture

Made New Media: AR-2

1. Made new media: AR-2

Solution	ID code	Volume	% Total volume
Advanced RMPI	—-	500 mL	78%
Glutamax	—-	$6.5~\mathrm{mL}$	20%
Pen-Strep	—-	$6.5~\mathrm{mL}$	1%
FBS	—-	$128~\mathrm{mL}$	1%

Monday 09-10-2023

Cell Culture

Seeded A20

 $\bullet~$ Got A20 from Swetha box: 2nd rack in shared -80

• AR-2

Seeded FDC

- Seeded feeders, taken from feeder box, irradiated by Chris
- Will seed N4 PDX tomorrow

Tuesday 10-10-2023

Cell Culture

Seeded N4

- Given by Chris -AR-2
- Seeded onto FDC cells from 09-10
- Future experiments:
 - Barcode PDX
 - EC50: Vincristine/Doxorubicin/Methotrexate/Cyclophosphamide (Active)

Seeded RAMOS-BC

- Seeded barcoded Ramos cells Pool#4
 - From Chris
 - LN tank 1 Rack 5 box 2nd from Bottom
- AR-2
- Future Experiments:
 - Combination Drug pressure: Vin/Dox/Meth

Checked A20

- Look healthy
- ~40% confluency